Review

Role of Biotransformation in Drug-Induced Toxicity:
Influence of Intra- and Inter-Species Differences in Drug Metabolism

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Summary: It is now widely appreciated that drug metabolites, in addition to the parent drugs themselves, can mediate the serious adverse effects exhibited by some new therapeutic agents, and as a result, there has been heightened interest in the field of drug metabolism from researchers in academia, the pharmaceutical industry, and regulatory agencies. Much progress has been made in recent years in understanding mechanisms of toxicities caused by drug metabolites, and in understanding the numerous factors that influence individual exposure to products of drug biotransformation. This review addresses some of these factors, including the role of drug-drug interactions, reactive metabolite formation, individual susceptibility, and species differences in drug disposition caused by genetic polymorphisms in drug-metabolizing enzymes. Examples are provided of adverse reactions that are linked to drug metabolism, and the mechanisms underlying variability in toxic response are discussed. Finally, some future directions for research in this field are highlighted in the context of the discovery and development of new therapeutic agents.

Keywords: drug toxicity; reactive metabolites; drug interactions; species differences; pharmacoge-nomics; safety evaluation

Introduction

Studies on the metabolic fate of new drugs have been an integral component of the drug discovery and development process for many years, largely in recognition of the potential of circulating metabolites to serve as mediators of both the efficacy and toxicity of their respective parent compounds. While initial interest in drug metabolites focused primarily on their contribution to the pharmacology of the parent (and led to the successful development of several metabolites as drugs in their own right), the role of metabolites in adverse drug reactions has become a topic of considerable interest in recent years. Given the remarkable advances over the past two decades in analytical technologies (notably liquid chromatography-tandem mass spectrometry, LC-MS/MS) for the detection, identification, and quantitative analysis of drug metabolites in biological fluids, it has now become relatively straightforward to characterize the metabolic profile of a new chemical entity in animals and human subjects. However, the translation of chemical structure to toxicological potential remains challenging, and the effectiveness of preclinical safety assessment programs in predicting drug toxicity in man continues to be debated. Inter-species differences in drug metabolism are the rule, rather than the exception, and frequently complicate extrapolation of animal safety data to the human situation. In addition, genetic polymorphisms in drug-metabolizing enzymes and transporters, both in animals and human subjects, can result in pronounced intra-species differences in drug disposition, further compounding the problem. Nevertheless, much progress has been made in gaining a scientific understanding of these issues, which has led to a deeper appreciation of the role of biotransformation in drug-induced toxicities. This review attempts to summarize current knowledge on the topic, and draws upon selected examples from the literature where metabolism of therapeutic agents is believed to have played a central role in drug-induced toxicity.

Classification of Drug-Induced Toxicities

Based on an analysis of the primary causes of withdrawal of approved drugs and the attrition of new chemical entities during the drug development process, Park and co-workers proposed a useful classification scheme that describes broad
mechanisms of drug-induced toxicities. More recently, Smith and Obach added to this classification, such that four fundamental mechanisms of toxicity (designated A through D) may be discerned, as shown in Table 1. Type A toxicities result from untoward pharmacology (either on- or off-target), while Types B through D are believed to be triggered by a discrete chemical insult which results in a cascade of events that ultimately manifest as one of a wide variety of drug-mediated toxic effects. It is noteworthy that products of drug biotransformation are highlighted as playing a causative role in mechanisms B, C, and D, underscoring the important role that drug metabolism may play in mediating the adverse effects of therapeutic agents. It also should be noted that some Type A toxicities result indirectly from metabolic events, particularly in cases where pharmacokinetic drug-drug interactions elevate circulating levels of one interactant to the point that an exaggerated pharmacological response, either at the intended or an unrelated biological target, leads to the expression of clinical toxicity. Some examples of these mechanisms are discussed in the following sections.

Role of Drug Metabolism in Drug-Mediated Organ Damage

From an anatomical and functional perspective, the liver plays a critical role in the disposition of therapeutic agents administered by the oral route, serving as a portal to the tissues, and represents the major site of drug metabolism. As pointed out above, products of drug metabolism have been implicated as causative agents in several types of toxicity, and it is not surprising, therefore, that the liver represents a major site of drug-induced toxicity. Indeed, a wide range of therapeutic agents have been withdrawn from clinical use in the U.S. and elsewhere due to an unacceptably high incidence of hepatotoxicity, including aclofenac, alpidem, amodiaquine, benoxaprofen, bromfenac, ibufenac, iproniazid, nefazodone, sudoxicam, tienilic acid, tolxerat, troglita-

Table 1. Mechanisms of drug-induced toxicities (adapted from Park et al., 1998, and Smith & Obach, 2009)

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type A</td>
<td>Normally reversible, involving a defined target leading to a predictable pharmacodynamic outcome. Two sub-types are distinguished:</td>
</tr>
<tr>
<td>A1</td>
<td>Desired target pharmacology (exaggerated response or wrong tissue, e.g. myopathy with statins)</td>
</tr>
<tr>
<td>A2</td>
<td>“Off-target” pharmacology (lack of selectivity, e.g. binding to K+ channel)</td>
</tr>
<tr>
<td>Type B</td>
<td>“Idiosyncratic” drug toxicities (not predictable):</td>
</tr>
<tr>
<td></td>
<td>Rare, do not exhibit classic dose-response relationship</td>
</tr>
<tr>
<td></td>
<td>Manifest after repeat administrations (e.g. halothane-induced hepatitis)</td>
</tr>
<tr>
<td></td>
<td>Triggered by reactive drug metabolites that form adducts with proteins</td>
</tr>
<tr>
<td>Type C</td>
<td>May occur after a single high dose</td>
</tr>
<tr>
<td></td>
<td>Exhibit classic dose-response relationship (above dose “threshold,” e.g. APAP)</td>
</tr>
<tr>
<td></td>
<td>Triggered by reactive drug metabolites that form adducts to proteins and/or cause oxidative stress</td>
</tr>
<tr>
<td>Type D</td>
<td>Occur only after prolonged dosing (carcinogenicity, teratology)</td>
</tr>
<tr>
<td></td>
<td>Associated with chemotherapeutic alkylating agents and reactive drug metabolites that form adducts to DNA and/or cause oxidative stress</td>
</tr>
</tbody>
</table>

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Nrf2 cell defense system, and also serves as an intracellular oxidizing agent. A number of hypotheses have been advanced to account for the hepatotoxic properties of APAP, and while there remains lack of clarity on the detailed molecular events, it appears that the metabolic formation of NAPQI “upstream” induces cellular stress and triggers a complex series of immune-mediated responses “downstream.” These changes, in turn, perturb the balance of pro- and anti-inflammatory cytokines, ultimately bringing about the centrilobular hepatic necrosis that is characteristic of APAP overdose. Although liver injury has been recognized as a serious consequence of APAP overdose, it is remarkable that APAP-mediated hepatotoxicity is claimed to be the most common cause of acute liver failure in the United States today.

Gaining an understanding of the role of drug metabolism in the liver injury caused by APAP has been important to the field of drug-induced hepatotoxicity from several perspectives. First, it led to the development of intravenous N-acetylcysteine as an antidote for APAP poisoning, which was believed initially to function as a thiol-containing surrogate for GSH, but which was shown subsequently to act as a precursor for GSH biosynthesis and thereby to replenish GSH pools used for detoxification of NAPQI. Second, the p-aminophenol sub-structure contained in the APAP molecule is imbedded in many other therapeutic agents, or can be introduced or unmasked via metabolic reactions. Awareness of the potential for further metabolic activation of this element to yield an electrophilic quinone imine species has been invoked retrospectively to account for the hepatotoxic properties of drugs such as diclofenac, nefazodone, trazodone, tacrine, amodiaquine, and lapatinib (Fig. 2), and also may be employed in a prospective sense in screening new chemical entities for possible bioactivation liabilities based on the detection of GSH adducts in vitro or in vivo. Indeed, it is now appreciated that a wide variety of compounds with heteroatom-substituted benzene rings can undergo metabolic activation (normally catalyzed by P450 enzymes) to generate electrophilic “quinoid” products (quinones, quinone imines, quinone methides, etc.) that bind covalently to cellular
macromolecules and, in some cases, cause oxidative stress via reactive oxygen species; both of these mechanisms can lead to liver toxicity (Fig. 3).

Structure-metabolism relationships such as those discussed above have led to the development of "structural alerts," or "toxicophores," i.e., lists of functional groups that may undergo metabolism to generate chemically reactive, potentially toxic species. Examples include thiophenes and other sulfur-containing heterocycles (via S-oxidation), furans (via epoxidation), anilines (via N- or C-oxidation), nitrobenzenes (via nitro reduction), hydrazines (via oxidation to free radical species), and some carboxylic acid derivatives (via acyl glucuronide or acyl-coenzyme A thioester formation). It should be emphasized, however, that in the context of drug discovery and lead optimization efforts, it is imperative to determine experimentally whether one of these functional groups, if present in a candidate of interest, actually is subject to metabolic activation. In many cases, a thiophene or furan substituent in a particular molecule may not be accessible to drug-metabolizing enzymes, and therefore be "silent" with respect to bioactivation. In addition, acyl glucuronide conjugates differ widely in terms of their intrinsic reactivity and propensity to rearrange through acyl migration, and the mere observation of such a conjugate being formed as a metabolite of a carboxylic acid-containing drug should not necessarily be taken as a portent of liver toxicity.25

The foregoing discussion has focused on the role of reactive, electrophilic metabolites in drug-induced liver injury where the underlying mechanism is believed to involve cellular damage through either covalent modification of critical cellular macromolecules, oxidative stress, or both. However, an additional mechanism of hepatotoxicity that has attracted considerable interest in the area of idiosyncratic drug reactions involves activation of the immune system through haptenization of proteins by reactive drug metabolites.26 Classic examples of this phenomenon are the inhalation anesthetic agent halothane, which undergoes P450-mediated oxidation to yield trifluoroacetyl chloride, and the anti-inflammatory drug tienilic acid, which is metabolized to a thiophene remote interwaters. These reactive metabolites covalently modify hepatocellular proteins which, in turn, are recognized as foreign by the immune system and induce an antibody response. Trifluoroacetyl chloride selectively acylates free amino groups on proteins, and antibodies have been detected in the sera of patients exposed to halothane that recognize liver neoantigens containing the trifluoroacetyl moiety.27 In the case of tienilic acid, the reactive metabolite covalently modifies CYP2C9, the P450 isofrom responsible for S-oxidation of the drug, resulting in the appearance of "anti-LKM1," autoantibodies in patients who developed hepatitis from tienilic acid therapy.28 Drug-induced toxicities that are mediated by the immune system are especially difficult to predict from preclinical toxicology studies due to the lack of generally applicable animal models for the human immune system. While the protein targets of some reactive metabolites are now known, much remains to be learned on this subject before a knowledge of the chemistry of metabolic activation can be linked to the toxicological outcome. Consequently, it would seem prudent, during the lead optimization phase of early drug development, to evaluate candidate molecules of interest for their propensity to generate chemically reactive metabolites, and to attempt (through appropriate structural modification) to minimize this potential liability in order to reduce the risk of downstream metabolism-dependent toxicities.29

Whereas drug-induced liver injury has been, and remains, an area of significant concern during both drug development and post-marketing surveillance, other metabolism-based iatrogenic organ toxicities can also occur. Indeed, extrahepatic tissue-selective toxicities caused by a variety of environmental chemicals are well documented. 3-Methylindole and 4-ipomeanol are two of the most intensively studied lung-selective toxicants, both of which require P450-mediated bioactivation to elicit their toxicity.30,31 The expression of lung-selective CYP2F and CYP4B enzymes has been associated with the metabolic activation of 3-methylindole to 3-methylenedione and 4-ipomeanol to its reactive ene-dial,32 respectively. These two agents are well recognized pulmonary toxins in several experimental animal species. However, extrapolation to humans of the lung-specific toxicities displayed by such model compounds is complicated by the much lower concentrations of P450 isoforms that are present in human lung compared to
rodents and other susceptible species.\textsuperscript{34} The organ-specific nephrotoxic potential of halogenated alkenes, such as trichloroethylene, has been long recognized.\textsuperscript{35} Bioactivation of haloalkenes generally involves the GSH pathway, with a major mechanism for toxicity resulting from the further processing of cysteine conjugates to reactive intermediates by renal cysteine conjugate \(\beta\)-lyase.\textsuperscript{36} \(P450\)-dependent metabolism also is implicated in the renal damage observed with numerous environmental chemicals.\textsuperscript{37} For example, chloroform can be bioactivated to the reactive metabolite phosgene by CYP2E1. Good evidence exists that the kidney toxicity of chloroform in mice is attributable to renal Cyp2e1 because Cyp2e1-null mice were resistant to chloroform-induced kidney toxicity,\textsuperscript{38} whereas mice harboring a liver-specific \(P450\) reductase knockout still displayed renal damage.\textsuperscript{39} Kidney damage can also result from the physical deposition of poorly soluble metabolites, such as those derived from ethylene glycol. Case examples illustrating species differences in metabolism-based renal toxicity are provided in a later section.

**Metabolic Drug-Drug Interactions and Product Withdrawals**

In addition to the drug metabolism-dependent organ toxicities described above, where cellular injury is caused by one or more metabolites of a single parent compound, drug-drug interactions in polytherapy may lead to a serious adverse event. In most cases, the underlying mechanism involves inhibition of \(P450\) activity by one of the interactants, resulting in decreased clearance of the second agent and expression of a toxicity that normally is observed only in overdose situations. Although a number of organ systems may be affected by this mechanism, much attention in recent years has focused on the cardiovascular system.

Cardiovascular adverse events most frequently have been a consequence of delayed ventricular polarization (QTc interval prolongation) resulting from drug binding to (and inhibition of) hERG, the \(\alpha\)-subunit of the cardiac \(I_{\text{K}}\) channel. Numerous approved drugs, including terodiline, sertindole, terfenadine, astemizole, grepafloxacin, cisapride, droperidol, levacetylmethadol, thioridazine, and doxetilide, were withdrawn from the market as a result of adverse QTc effects, in many cases precipitated by drug-drug interactions. Perhaps the best documented example of such an interaction was with terfenadine (Seldane), a potent histamine H1 antagonist (\(IC_{50} = 1\) nM) introduced in the U.S. in 1985 for the treatment of seasonal allergies. While terfenadine also inhibits hERG (\(IC_{50} = 56\) nM), circulating levels of the drug in humans following a therapeutic dose rarely approach toxic concentrations due to extensive first-pass metabolism to yield the corresponding active carboxylic acid metabolite, fexofenadine (\(IC_{50} at H1 = 15\) nM), and so QTc prolongation with terfenadine was believed only to be an issue in cases of terfenadine overdose. However, the biotransformation of terfenadine to fexofenadine (which binds weakly to hERG) is catalyzed primarily by hepatic and intestinal CYP3A4, and can be blocked by potent inhibitors of this enzyme, \textit{e.g.}, erythromycin and ketoconazole (Fig. 4). Under such conditions, blood levels of terfenadine rise to values that may exceed its \(IC_{50}\) for hERG, with resulting prolongation of the QTc interval and, in severe cases, precipitation of life-threatening cardio toxicity in the form of Torsades de Pointes.\textsuperscript{40} Recognition of the mechanism of this toxic drug-drug interaction led to the effective replacement of terfenadine by its active metabolite, fexofenadine (Allegra) in 1996, and terfenadine was withdrawn from the U.S. market the following year. In light of this experience, screening of new chemical entities for hERG binding has been implemented throughout the pharmaceutical indus-

![Fig. 4. Oxidation of terfenadine to its active carboxylic acid metabolite fexofenadine, and inhibition of this pathway by inhibitors of CYP3A4](image-url)
try, as have assays for P450 inhibition that are designed to eliminate potent P450 inhibitors as development candidates, thereby minimizing the risk of failure due to an underlying drug interaction liability. It should be pointed out that one potent mechanism-based P450 inhibitor, ritonavir, has found utility in anti-HIV therapy where it is used primarily to “boost” the systemic exposure of other antiretroviral drugs. However, the general utility of such “therapeutic enhancers” likely will be limited to life-threatening diseases due to the incidence of unintended interactions with co-administered drugs.

**Inter-Species and Intra-Species Differences in Drug Metabolism**

Animal studies during preclinical drug development are used to evaluate the absorption, distribution, metabolism, and excretion (ADME) profile and safety of new candidate drugs. Ideally, data obtained from such animal studies can be used to screen out severe toxicities and to estimate a safe, maximum starting dose in human Phase I clinical trials. However, this begs the important question, “To what extent can animal studies predict the ADME properties of new chemical entities and (potentially related) human toxicities?” In terms of predicting human toxicities from animal data, a great deal of this information is, of course, not publicly available. Nonetheless, attempts have been made to address this question through compilation of available toxicity data in non-human primates, dogs, rats, and mice. While this analysis has revealed that human cardiovascular, gastrointestinal, and blood cell toxicities are predicted successfully from animal data in more than 80% of cases, overall, about a third of global human toxicities are not predicted in ANY of the standard animal species employed for drug safety assessment. (Dogs and rats are the commonest non-rodent and rodent species, respectively, that are used for pre-clinical toxicology testing). Further analysis demonstrated that dogs are much more useful than rodents in predicting up to 150 different human toxicities. However, the dog was still not predictive for about one-third of drug toxicities reported ultimately in humans. Perhaps surprisingly, non-human primates performed no better than dogs as predictive species for drug-related toxicities in humans.

Differences in drug response, including toxicity, across species (and individuals) will reflect both pharmacokinetic and pharmacodynamic variables. Drug pharmacodynamics often are highly individualized according to the pharmacological target involved, whereas pharmacokinetics are dictated by relatively conserved ADME processes. However, cross-species ADME comparisons also are complicated by a variety of factors, cogently summarized by Lin. For example, when considering the oral absorption of a drug, it is well recognized that dogs are poor acid secretors, whereas humans and rats are good acid secretors. Therefore, when the solubility of a drug is pH-dependent, species differences in drug absorption can be expected. In a similar vein, species differences exist in biliary excretion. Rodents and dogs are considered good biliary excretors compared to humans, and this can be reflected in species differences in the extent of fecal excretion and enterohepatic recycling. However, of more importance than biliary excretion to the overall elimination of drugs are renal and metabolic clearance. Predicting renal excretion across species is one area where allometric scaling of relative glomerular filtration rates works quite well, at least for high extraction ratio drugs. On the other hand, scaling metabolism across species is much more problematic. This is particularly unfortunate because it is estimated that some 70% of drugs are cleared primarily by metabolism. Moreover, metabolism often is an important determinant of toxicity through the formation of reactive (electrophilic) intermediates that may react with tissue nucleophiles and DNA (vide infra).

As noted above, metabolism dominates clearance routes, with about two-thirds of all drugs cleared primarily by Phase I and Phase II enzymes. P450s and UGTs dominate Phase I and Phase II processes, respectively, and in humans CYP3A4 and UGT2B7 are the specific enzymes that respectively oxidize and conjugate the majority of drugs. In terms of animal-to-human predictions, a sensible question would be, “Are all these same drug-metabolizing enzymes present and active in all animal species?” The short answer is NO – and indeed there are many examples of very large differences in metabolizing enzyme capacity across species. For example, species sensitivity to organophosphates (OPs) has been linked to the widely differing plasma levels of paraoxonase (PON1), which hydrolyzes many OPs. Organophosphates elicit neurotoxicity by irreversibly inactivating acetylcholinesterase in the brain, so PON1 serves a protective function against exposure to these toxins. The particularly high sensitivity of birds to OP poisoning has been attributed to their extremely low levels of plasma PON1. On a more domestic level, inadvertent facile poisoning of cats with APAP is due to a lack of the feline NAT enzyme that conjugates and eliminates the drug, thereby permitting its build-up to toxic levels. Of high relevance to drug disposition studies in industry is the knowledge that dogs (and other canids) completely lack N-acetyltransferase (NAT) genes. Therefore, these species have a greatly reduced capacity to metabolize primary aromatic amines by the N-acetylation pathway.

Of substantial relevance to the drug discovery/development process is the extent to which P450 activities are conserved across species, especially those species used in safety assessment, such as the rat and dog. Table 2 lists all the important human liver drug-metabolizing P450s and identifies homologs in other species. The term “homolog” in biology refers to a related gene, which can be identified on the basis of sequence similarities, whereas “ortholog” is generally reserved for a related gene that maintains functional similarities. For example, rats possess at least six forms of CYP2D enzymes. However, none of these rat forms have
the same substrate preferences as human CYP2D6, and should be described as homologs. Only CYP2E1 (and to a lesser extent the CYP1A enzymes) maintain functional similarities across species.50

Another way to highlight inter-species differences in P450 metabolism is to compare the activity of a substrate probe for a given human enzyme across different species. Pelkonen and coworkers compared the rates of metabolism of eleven such P450 probes in microsomes from mouse, rat, rabbit, dog, mini-pig, monkey, and human livers,51 such P450 probes in microsomes from mouse, rat, rabbit, and human livers.52

Table 2. Homologs of the major human liver P450s present in experimental animals

<table>
<thead>
<tr>
<th>Subfamily</th>
<th>Human</th>
<th>Rat</th>
<th>Mouse</th>
<th>Dog</th>
<th>Monkey</th>
</tr>
</thead>
<tbody>
<tr>
<td>CYP1A</td>
<td>1A1, 1A2</td>
<td>1A1, 1A2</td>
<td>1A1, 1A2</td>
<td>1A1, 1A2</td>
<td>1A1, 1A2</td>
</tr>
<tr>
<td>CYP2A</td>
<td>2A6, 2A7, 2A13</td>
<td>2A1, 2A2, 2A3</td>
<td>2A4, 2A5, 2A12, 2A22</td>
<td>2A13, 2A25</td>
<td>2A23, 2A24</td>
</tr>
<tr>
<td>CYP2B</td>
<td>2B6</td>
<td>2B1, 2B2, 2B3</td>
<td>2B9, 2B10</td>
<td>2B11</td>
<td>2B6, 2B17</td>
</tr>
<tr>
<td>CYP2C</td>
<td>2C8, 2C9, 2C19</td>
<td>2C6, 2C7, 2C11, 2C12 +</td>
<td>2C29, 2C37, 2C38, 2C19 +</td>
<td>2C21, 2C41</td>
<td>2C20, 2C43</td>
</tr>
<tr>
<td>CYP2D</td>
<td>2D6</td>
<td>2D1, 2D2, 2D3, 2D4, 2D5, 2D18</td>
<td>2D9, 2D10, 2D11, 2D12 +</td>
<td>2D15</td>
<td>2D17, 2D19, 2D29, 2D30</td>
</tr>
<tr>
<td>CYP2E</td>
<td>2E1</td>
<td>2E1</td>
<td>2E1</td>
<td>2E1</td>
<td>2E1</td>
</tr>
<tr>
<td>CYP2F</td>
<td>3A4, 3A5, (3A7)</td>
<td>3A1/23, 3A2, 3A9, 3A18 +</td>
<td>3A11, 3A13, 3A16 +</td>
<td>3A12, 3A26</td>
<td>3A8</td>
</tr>
</tbody>
</table>

For non-human primates, it might be anticipated that liver P450s in humans and monkeys would be very closely related—and this does, in fact, turn out to be the case. Whereas sequence similarities between human and dog P450 “orthologs” are only about 70%, monkey orthologs of human drug-metabolizing P450s in the CYP1-3 families with sequence homologies as high as 95%, are readily identified.53 There are two points of caution, however. First, human CYP2C9 and CYP2C19 share >90% sequence identity, yet they catalyze quite different metabolic reactions. Second, a new monkey P450, CYP2C76, has been identified recently, but this enzyme has no ortholog in humans.54 These considerations provide a molecular basis for the species differences in P450-dependent drug metabolism that can arise between humans and non-human primates. To summarize, highly variable microsomal P450 probe metabolism across species could be a result of: (a) differences in the concentration and/or intrinsic activity of the orthologous enzyme(s) that catalyze the probe reaction, or (b) a lack of functionally similar enzymes, and metabolism by either a homologous enzyme or a completely different form of P450 in that species.

Species differences in drug metabolism take on an added significance when they are responsible for species differences in toxicity. The following three examples will illustrate this point for different Phase I and Phase II enzymes. The first example is 4-ipomeanol, a furan-containing compound that was first found to cause lung toxicity after farm animals ingested it unintentionally in their feed.56 4-Ipomeanol became a model for xenobiotic-induced lung injury and its metabolism has been studied extensively.57 It was established more than 25 years ago that lung CYP4B1 catalyzed the bioactivation of 4-ipomeanol in animals,57 and it has been shown recently that CYP4B1 can convert ipomeanol in vitro to a reactive ene-dial (Fig. 5), presumably following initial epoxidation on the furan ring.53 In the late 1980s, 4-ipomeanol was evaluated in humans by the National Cancer Institute (NCI) as a possible treatment for non-small cell lung carcinoma,58 but the compound was found to be ineffective, probably because lung CYP4B1 in humans has little or no
catalytic activity. In other human studies, a dose-limiting liver toxicity was observed. What, then, is the basis for the species difference in organ toxicity? At least two possibilities can be considered: (a) alternative bioactivating liver P450 enzyme(s) in humans, or (b) a lack of detoxifying or conjugating enzyme(s) in human liver. Further studies are needed to discriminate between these possibilities.

The second example is renal toxicity of the non-nucleoside reverse transcriptase inhibitor efavirenz. During development, efavirenz was found to cause renal tubule epithelial cell toxicity in rats, but not in monkeys or humans. The drug was metabolized extensively in all three species, but only the cysteinylglycine conjugate M10 (Fig. 6) and its precursor GSH conjugate, M9, were formed exclusively in the rat. This observation was critical because the kidney is the major site for processing GSH conjugates through the mercapturic acid pathway, a process that is initiated here by the conversion of M9 to M10 by the enzyme γ-glutamyl-transpeptidase (GGT). It was proposed that metabolite M10 would be subject to further metabolism under the action of a dipeptidase to generate the corresponding cysteine conjugate, which, in turn, could be activated, either by S-oxidation or via the cysteine conjugate β-lyase pathway, to yield the ultimate toxic species (Fig. 6). Evidence in support of this hypothesis was obtained from studies in rats pretreated with the glutathione-S-transferase (GST) inhibitor acivicin, which protected animals from the renal toxicity of efavirenz. It was concluded, therefore, that the basis for the species difference in efavirenz renal toxicity likely is related to species difference in GST activity between rats, monkeys, and humans.

Finally, a very recent example of renal toxicity in humans arose with the experimental kinase inhibitor SGX523. When the drug’s metabolic profile was elucidated, only humans formed substantial quantities of a lactam metabolite, M11, which was determined to be formed by aldehyde oxidase (AO). The neutral, AO-derived lactam metabolite was very poorly soluble in patient urine, and so lactam...
precipitation was suggested to be the cause of the renal crystal deposits. Notably, in vitro studies with liver microsomes would not have revealed this metabolite because AO is a cytosolic enzyme, and in vivo studies in dogs and rats failed to identify M11 as a possible metabolite because AO activity is low or absent in these preclinical animal species. In summary, predicting drug metabolism and drug toxicity in humans from animal data often is problematic. Regarding metabolism, a central issue is that homologous Phase I and Phase II enzymes across species often:

- exhibit different substrate specificities,
- catalyze the same reaction at very different rates, or
- are present in liver at vastly different concentrations.

Genetic polymorphisms in drug-metabolizing enzymes add a further complication in extrapolating metabolic data obtained from a sub-set of human subjects (or even animals) across the larger population. In terms of animal experimentation, this is usually a minor issue because most preclinical animal studies are conducted on inbred strains, which generally results in only an approximately two- to three-fold inter-animal variation in the metabolic data. Notable exceptions occur in the beagle, which exhibits a pronounced polymorphism in CYP1A2, and in Gunn rats, which are deficient in UGT1 activity.

In contrast to the typical situation in experimental animals, in humans there can be huge (100-fold) variations in a drug’s metabolic clearance when the pathway involved is under genetic control. Genetic polymorphisms in several Phase I and Phase II enzymes, notably the CYP2D6, CYP2C19, CYP2C9, and UGT1A1 enzymes, grossly affect human drug metabolism. The resulting phenotype in carriers of the minor allele is usually one of reduced metabolic function, although rarer polymorphisms in both the CYP2D6 and CYP2C19 genes are known that confer increased metabolic function. The consequences to the patient will also depend on whether metabolism of the parent drug terminates the desired therapeutic activity or, in the case of prodrugs, confers the pharmacological effect. In this section, genetic polymorphisms in human drug-metabolizing enzymes will be discussed as a complicating factor in assessing drug safety, both from the perspective of drug-induced liver injury and drug-drug interactions—two important reasons for drug failures in the clinical setting, as noted earlier.

Several drug-metabolizing enzyme (and transporter) variants are now recognized as risk factors for drug-induced liver injury. These effects presumably are manifest via an initiating toxic event at the level of reactive metabolite interaction with cellular constituents. A major difficulty in
this area has been the accumulation of sufficient samples of confirmed pathology for analysis, because the incidence of idiosyncratic hepatotoxicity is very low. However, recently established consortia that address this problem have provided new momentum to this field, along with the use of new analytical tools such as genome-wide association studies (GWAS). This latter approach recently enabled identification of the HLA-B*5701 allele as a major risk factor in the development of fluvoxacin-induced liver injury.67

In terms of drug-metabolizing enzymes, GWAS has yet to be exploited fully, but several candidate gene studies involving isoniazid and diclofenac have appeared. For isoniazid-induced liver injury, the GSTM1 null, NAT2 slow acetylator, and CYP2E1 wild-type alleles were found to be significantly associated with the adverse outcome.68,69 These findings are particularly interesting because each of the gene effects may be rationalized in terms of either increasing reactive metabolite formation or decreasing their detoxification. For diclofenac, several possible reactive intermediates have been postulated, including the 2,5- and 2,4′-quinone imines and both the parent and 4'-hydroxy-diclofenac acyl glucuronides (Fig. 7). These metabolites may result from combined metabolism involving CYP2C8 and UGT2B7, and, in fact, the CYP2C8*4 and UGT2B7*2 alleles were found to be associated with diclofenac-induced liver toxicity.70 Again, the pathway analysis used here is compelling because the effect of both variants can be rationalized in terms of increasing toxic metabolite formation. However, these two alleles are at best low-risk genetic variants, and it is clear that interactions between multiple risk factors for idiosyncratic hepatotoxicity are needed to explain the low incidence of this undesirable iatrogenic complication to therapy.66

The P450 genotype is now a well-recognized modifying influence on metabolic drug interactions involving CYP2D6 or CYP2C19 poor metabolizers (PMs). As reviewed by Lee et al.,71 the CYP2D6 inhibitors quinidine and fluoxetine had virtually no effect on the pharmacokinetics of the CYP2D6 substrates venlafaxine and risperidone in PMs, but increased the area under the plasma concentration-time curve (AUC) of these two drugs from 4- to 12-fold in extensive metabolizers (EMs). Additionally, the CYP2C19 inhibitors omeprazole and fluvoxamine increased the AUC of moclobemide and lansoprazole by 2.2- and 3.8-fold, respectively, in CYP2C19 EMs, but had little or no effect in PMs.72,73 When there is a clear functional P450 deficit, inductive drug interactions also appear to be abrogated in PMs. An early study by the Vanderbilt group demonstrated with mephenytoin that CYP2C19 EMs exhibited a three- to eight-fold increase in the ratio of urinary ratio of R:S mephenytoin and a 40–180% increase in the 0–8 hour urinary excretion of the 4′-hydroxy-mephenytoin metabolite. In stark contrast, CYP2C19 PMs did not respond to either parameter following induction with a course of rifampin.74 More recently, Tracy and coworkers have shown that changes in the oral clearance of flurbiprofen (a probe for CYP2C9) after fluconazole co-administration decreased in the order: CYP2C9*1*1 > *1*3 > *3*3 genotypes.75

Collectively, the foregoing in vivo examples provide the guiding principle: P450 genotype modifies the magnitude of a metabolic drug interaction as a function of the extent to which the polymorphic P450 catalyzes the target reaction. Consequently, one would anticipate that PM status for CYP2D6, CYP2C19, and CYP2C9 would afford protection against certain drug-drug interactions. This would affect approximately 7%, 2%, and <1%, respectively, of Caucasian populations, with some pronounced ethnic differences apparent, especially for CYP2C19 PMs, which are much more common (15–20%) in Asian populations.76 By extension, graded drug interaction responses would be expected for heterozygotes, or intermediate metabolizer phenotypes, and possibly magnified responses in ultra-rapid CYP2D6 metabolizers, although this remains to be investigated. In contrast to these CYP2 family enzymes, CYP3A4 is not subject to polymorphic variability to the extent that ultrarapid or poor metabolizer phenotype/genotypes are apparent in the general population. Therefore, genotype-dependent modulation of the serious drug-drug interactions that limited the clinical use of terfenadine (and other CYP3A4 substrates) would not be expected.

**Regulatory Guidances on Metabolites in Safety Testing**

The growing awareness of the potential role of drug metabolites in the toxicity of their respective parent compounds has led, in recent years, to increased scrutiny from regulatory agencies on the adequacy of safety assessment programs with regard to the toxicological properties of circulating drug metabolites. Following the publication of a “best practices” position paper on this issue in 2002 by the U.S. Pharmaceutical Research and Manufacturers of America (PhRMA),77 which stimulated much discussion at both scientific meetings and in the peer-reviewed literature, the U.S. Food & Drug Administration (FDA) and the International Conference on Harmonisation (ICH) each published regulatory guidance documents on the safety testing of drug metabolites.78,79 Although the two documents differ in some respects (notably on the definition of what constitutes the exposure threshold above which a metabolite is deemed of interest), the FDA announced recently that they now accept the criteria set out in the ICH document.80 The emphasis of both guidance documents is on biotransformation products that circulate and are anticipated to circulate in human plasma; the overall goal of this guidance is to provide assurance that preclinical safety findings are relevant to the human situation in terms of exposure to both the parent drug and its circulating metabolites. At the heart of the matter is the question of predicting drug toxicity in humans from results obtained in animals, where the types of
species differences in drug metabolism discussed above may confound the interpretation of animal toxicology data in terms of human risk assessment.81

Key components of the FDA guidance may be summarized as follows: The document applies only to small-molecule non-biologic products, and excludes anti-cancer agents, drug conjugates (other than acyl glucuronides), and chemically reactive metabolites. The primary focus is on stable drug metabolites circulating in human plasma, notably those that are “unique” or “disproportionate” in humans and which may require separate testing. Such evaluation (of the preformed metabolite) could include general toxicology studies (3 months duration), assessment of genotoxicity and embryo-fetal developmental toxicity, and, in some instances, carcinogenicity studies. An important element of both the FDA and ICH documents relates to the definition of a metabolite of interest. In this regard, the wording of the ICH guidance (which has now been endorsed by the FDA) states: “Nonclinical characterization of a human metabolite(s) is only warranted when that metabolite(s) is observed at exposures greater than 10 percent of total drug-related exposure and at significantly greater levels in humans than the maximum exposure seen in the toxicity studies.” The document goes on to state, “For drugs for which the daily administered dose is <10 mg, greater fractions of the drug related material might be more appropriate triggers for testing” and “The nonclinical characterization of metabolites with an identified cause for concern (e.g., a unique human metabolite) should be considered on a case-by-case basis.”

These two pieces of regulatory guidance, which address an important topic in the safety assessment of new therapeutic agents, raise a number of practical considerations for the pharmaceutical industry. For example, how does one best establish the identity and quantitative importance of individual drug metabolites in human plasma at a sufficiently early stage of clinical development that appropriate toxicological studies, where necessary, can be completed prior to the initiation of large-scale (Phase III) clinical trials? Related to this question, it should be noted that the traditional radiolabeled human ADME study typically is not conducted until after pharmacological proof-of-concept has been obtained (in Phase IIA), which leaves insufficient time to perform toxicology studies on previously unrecognized circulating metabolites with higher exposures in man relative to animals (“disproportionate metabolites”). Also, in the absence of data from a radiolabeled human ADME study, how does one assess whether a given circulating metabolite exceeds 10% of “total drug-related exposure”? One “cold” (i.e., non-radioactive) approach to the problem calls for the use of LC-MS/MS, under conditions of high mass resolution, to screen plasma samples from early (Phase I) clinical trials for components of plasma that exhibit accurate mass values consistent with being structurally related to the parent drug.82 This technique, known as “mass defect filtering,”83 represents a highly sensitive, selective, and relatively unbiased approach to the detection and identification of drug metabolites in biological fluids and, when mass spectrometric responses are calibrated by an external method (e.g., radioactivity measurements from preclinical samples), also can provide quantitative data on metabolite exposure.

Once a circulating human metabolite has been deemed to be “disproportionate” (i.e., exhibits an animal:human exposure margin less than unity), careful consideration needs to be given to the design of appropriate studies to assess its safety profile. While it is sometimes possible to escalate the dose of the parent compound in animal toxicology studies to establish required safety margins for a metabolite, or to employ a different animal species where the metabolite in question is better represented, these approaches are not always successful. In such situations, the FDA guidance recommends performing toxicological studies with the preformed metabolite, but it is known that the dispositional characteristics of a metabolite that is administered exogeneously often differ from those when the metabolite is generated in vivo from the parent drug.84 As such, dosing animals with synthetic metabolites, where feasible, runs the risk of generating data that are not relevant to the safety of the parent drug in man. Clearly, these are complex issues, and it is important that the approach that is adopted for a given development candidate is designed to generate scientifically meaningful data on the safety of circulating human metabolites, while simultaneously taking into consideration the constraints of available resources. In that context, a case-by-case approach to the issue would seem most logical.

Conclusions and Future Prospects

The 1971 Gordon Conference on Drug Metabolism opened with plenary lectures from two of the field’s founders, R. ’Tec’ Williams and Bernard R. Brodie, who spoke on the topics: “Factors Involved in Species Differences in Drug Metabolism” and “Biochemical Mechanisms of Drug-Induced Lesions,” respectively (Obach, personal communication). Forty years later, these two subjects continue to attract much attention, not least from scientists in the pharmaceutical industry charged with the development of new therapeutic agents. While the factors involved in species differences in drug metabolism are now reasonably well understood (vide infra), and considerable advances have occurred in our knowledge of both chemical and biochemical mechanisms of iatrogenic organ injury, prediction of the metabolic disposition and safety of new chemical entities in humans from animal studies remains a considerable challenge, although new biochemical and analytical tools continue to evolve to help mitigate this deficit. For example, in attempts to overcome the lack of conservation of drug-metabolizing functions for the major human P450s across different animal species, transgenic (or “humanized”) mice have been created, wherein the gene for a particular human P450 (e.g., CYP2D6) is introduced into their genome.
CYP2D is a human liver P450 that preferentially metabolizes ‘basic’ compounds and, consequently, this enzyme is responsible for the metabolism of many cardiovascular and CNS-acting drugs. A common probe reaction for CYP2D6 is 4-hydroxylation of debrisoquine, a relatively old antihypertensive drug. Opposed to one used extensively in drug development, created, but currently remain largely a research tool as with respect to several other human P450s have been increased greatly, thereby demonstrating the in vivo function of the transgene. Mice that have been humanized with respect to several other human P450s have been created, but currently remain largely a research tool as opposed to one used extensively in drug development.

Of the various categories of drug-induced toxicity classified in Table 1, the idiosyncratic, Type B, drug toxicities which have an immune basis continue to be the most difficult to rationalize and, therefore, to predict. Drug-induced immunogenicity is an especially serious concern during the development of peptide and protein drugs, and in silico and experimental approaches to preclinical screening for T-cell epitopes derived from normal cellular processing of biotherapeutics have been discussed. However, despite recent advances in our knowledge of the role of HLA variants as risk factors for drug-induced hypersensitivity reactions, including those that likely have a bioactivation component, devising a rational approach to preclinical screening of small molecules likely to possess this liability remains a daunting task. More promising, perhaps, is the development of lipopolysaccharide-based animal models for studying idiosyncratic toxicities with a strong inflammatory stress component, although, again, much remains to be learned about the extrapolation of data of this type to humans.

Finally, the technique of “microdosing,” in which a trace amount (typically 100 µg or less) of a 14C-labeled drug (usually <1 µCi) is administered to a human subject and radioactivity in plasma and excreta is followed by accelerator mass spectrometry, has been proposed as a highly sensitive and effective approach to assessing human drug metabolism during the lead optimization stage of drug development. Indeed, by means of this technique, a number of drug candidates can be assessed in humans in terms of their pharmacokinetic characteristics and metabolic profiles before one agent is selected for development. Importantly, this approach allows for metabolic profiles to be compared across species, such that differences in metabolism between humans and the animals employed in safety assessment studies can be taken into consideration in choosing the lead compound. The inclusion of patient cohorts genotyped for the common P450 polymorphisms in tracer dosing studies of this type could extend the utility of this approach to an early assessment of potential pharmacogenetic variability in candidate drug clearance, while minimizing drug exposure.

These and other technological developments promise to significantly advance our understanding of the molecular and biochemical basis of species differences in drug metabolism and toxicity, and thereby contribute to the development of safer, more effective therapies for unmet medical needs.

References


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