Mechanism of idiosyncratic drug induced liver injury (DILI): unresolved basic issues

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Abstract: Clinical features of idiosyncratic drug induced liver injury (DILI) are well described in cases that have been assessed for causality using the Roussel Uclaf Causality Assessment Method (RUCAM), but our understanding of the mechanistic steps leading to injury is fragmentary. The difficulties describing mechanistic events can be traced back to the lack of an animal model of experimental idiosyncratic DILI that can mimic the genetic requirements of human idiosyncratic DILI. However, immune tolerance plays a dominant role in the immune response of the liver, and impairment of immune tolerance with immune checkpoint inhibitors increases DILI in both humans and animals. This may provide one method to study the individual steps involved. In general, the human DILI liver is a secret keeper providing little insight into what occurs in the diseased organ. Sufficient evidence exists that most idiosyncratic cases are mediated by the adaptive immune system, which depends on stimulation of the innate immune system, but the triggering factors are unknown. It is attractive to hypothesize that the gut microbiome plays a role; however, it is very difficult to study. Similarly, exosomes are likely to play an important role in communication between hepatic cells and the immune system, but there is a lack of data on blood exosomes in affected patients. Reactive metabolites are likely to play an important role. This is supported by the current analysis, which revealed an association between metabolism by cytochrome P450 and drugs most commonly involved in causing idiosyncratic DILI with causality verified by RUCAM. Circumstantial evidence suggests that reactive oxygen species (ROS) generated by cytochrome P450 could be responsible for the initial steps of injury, but details are unknown. In conclusion, most of the mechanistic steps leading to idiosyncratic DILI remain unclear.

Keywords: Drug induced liver injury (DILI); idiosyncratic DILI; Roussel Uclaf Causality Assessment Method (RUCAM); innate immune system; lipopolysaccharides (LPS); gut microbiome; cytochrome P450; reactive oxygen species (ROS); oxidative stress; mechanistic steps

Introduction

A broad range of chemicals are injurious to the liver as evidenced by clinical and experimental studies, which commonly provide clues to mechanistic steps leading to injury. Chemicals of clinical interests include alcohol (1-5) and aliphatic halogenated hydrocarbons such as CCl₄ (carbon tetrachloride) (6-11). In addition, heavy metals are found in excess amounts in the liver of patients with genetic liver diseases such as primary hemochromatosis with excess...
of iron (12,13), or Wilson's disease with excess of copper (14,15). Clinically important are also toxic liver diseases caused by herbs leading to herb induced liver injury (HILI) (16-19), and drugs presenting as idiosyncratic drug induced liver injury (DILI) (17,20-24).

Idiosyncratic DILI is viewed as a complex and complicated human disease (17,20-30). The 10 drugs most commonly implicated in DILI worldwide starting with the most common are amoxicillin-clavulanate, flucloxacinil, atorvastatin, disulfiram, diclofenac, simvastatin, carbamazepine, ibuprofen, erythromycin, and anabolic steroids as body building agents (22). There is also a compilation of 46,266 worldwide DILI cases with established causality (23). However, data on the exact number of potentially hepatotoxic drugs are not available from these reports (22,23) or other publications (26). A recent review cited studies that estimated the number of drugs that cause DILI to range from 385 to 1,266 drugs (30), and other estimates from the United States came up with more than 1,000 drugs that cause DILI, although causality has not always been established clearly as explicitly mentioned (27). On the other hand, based on DILI cases contained in the LiverTox database, 353 drugs with the potential to injure the liver were recognized (25). The high variability of drug numbers is likely due to the lack of clear criteria of case inclusion or exclusion and confounding variables, poor case data quality, alternative causes, and incomplete causality assessment by a robust algorithm (22,23,25-31). It seems that perhaps a few hundred drugs are potentially hepatotoxic, but the exact number remains uncertain.

There is also much uncertainty on initial mechanistic steps leading to idiosyncratic DILI. Although much attention has been paid to this topic worldwide, it is a challenging issue due to lack of appropriate animal models leaving many open questions (24). These scientific conditions remind us of the musical “The unanswered question” by the US Charles Ives.

The aim of this article is to analyze unresolved basic issues of mechanistic steps in idiosyncratic DILI. Special attention is given to immune aspects in connection with the gut microbiome and the hepatic cytochrome P450 (CYP). It is hoped that a pragmatic and conceptual approach can at least partially close the gaps in our understanding of the mechanistic steps occurring initially and during the cascade of injurious events.

**Literature search and source**

The PubMed database was searched for articles on DILI by using the following terms: drug induced liver injury, DILI, idiosyncratic DILI, RUCAM, pathogenesis, mechanistic steps, adaptive immune system, innate immune system, HLA genes, hepatic cytochrome P450, reactive oxygen species (ROS), exosomes, and gut microbiome, whereby search terms were used alone or in combination. The electronic search was completed on 15 April 2020 and supplemented by a manual literature search, using also the private archives of the authors. Preference was given to articles in the English language. The final compilation consisted of original papers, consensus reports, and review articles with the most relevant publications included in the reference list.

**Definition**

Idiosyncratic DILI is a special type of human liver injury that is specific to an individual and does not occur in most patients treated with a drug. It is also idiosyncratic in animals, which poses a major challenge for developing animal models. Human idiosyncratic DILI reflects both the properties of the drug, with its special physicochemical features, and its interaction with a susceptible individual based on genetic and other unknown factors (23,24,32). In contrast, human intrinsic DILI is caused by specific toxic properties of the drug and occurs in virtually any human and most animals through mechanisms not requiring genetic predisposition (23,32). Differentiation between human idiosyncratic and intrinsic DILI is essential if clinical and mechanistic aspects are to be considered to avoid descriptions of idiosyncratic DILI characteristics based on mixed results of both DILI forms. The present article focuses on unresolved mechanistic steps of idiosyncratic DILI without extrapolation of mechanistic principles known for intrinsic DILI. Whereas mechanistic steps for drugs causing intrinsic DILI are commonly based on clear experimental or clinical evidence, this condition is not the case for idiosyncratic DILI, which relies mostly on circumstantial evidence.

**Unresolved key questions of basic mechanistic issues**

There are a number of interesting facts related to drugs and idiosyncratic DILI that warrant further consideration (Table 1). Although most of the clinical features are well established, a variety of mechanistic steps remain unresolved in this complex disease. They include drugs with
Table 1 | Fact sheet on drugs and DILI and unresolved mechanistic issues

| Facts of drugs and idiosyncratic DILI | Unresolved questions of basic mechanistic issues |
|--------------------------------------|-----------------------------------------------|
| Several hundred drugs are potentially hepatotoxic as compared to only a few drugs that appear never to cause idiosyncratic DILI | What are the differences between the two groups? Is it lipophilicity and active transport that concentrates the drug in the liver, reactive metabolite formation with the formation of neoantigens, interference with basic pathways such as BSEP inhibition leading to cell stress and the release of DAMPs, or a combination of such factors? |
| Although there are some common features, idiosyncratic DILI caused by different drugs is not uniform. In addition, the features of DILI overlap with liver injury caused by other agents such as viral hepatitis | What is the cause for the inhomogeneity of liver injuries? Can they be explained by differences in the mechanistic steps, or is it just the usual interindividual differences in the immune response to various agents? |
| The same drug may cause two or more different types of liver injury as assessed by liver histology and laboratory tests | Hepatocellular injury is the most common type of injury as compared with cholestatic injury and the rarer autoimmune DILI. Is an important factor that some drugs/metabolites are concentrated in bile? |
| Although the idiosyncratic DILI caused by some drugs is strongly associated with a specific HLA genotype, only a small fraction of patients with that genotype will experience DILI when exposed to the associated drug | What other risk factors in addition to HLA genotype are required? Is it the T cell receptor repertoire and/or other factors such as the gut microbiome? |
| Although in most cases it has never been tested, autoantibodies are found in the serum of patients with idiosyncratic DILI caused by some drugs | It is unclear whether such autoantibodies are pathogenetic or simply an indication of an immune response against the drug |
| There is strong evidence that the immune system is responsible for idiosyncratic DILI caused by many drugs | Are there other mechanisms of idiosyncratic DILI caused by some drugs that do not involve the immune system? |
| Adverse reactions mediated by the adaptive immune system require a prior activation of the innate immune system | What mediators are responsible for activation of the innate immune system? Are they produced solely by hepatocytes or are non-parenchymal cells also involved? |
| Some patients treated with potentially hepatotoxic drugs show clinical and laboratory signs of immunoallergy or even autoimmunity while others do not | Do differences in the chemical structure of drugs determine whether they can induce an immune response, and is prediction of liver injury possible in a setting of drug development? |
| Exosomes appear to be an essential mechanism by which organs such as the liver communicate with the immune system. For example, drug-modified proteins are present in the exosomes from drug-treated hepatocytes and are taken up by antigen presenting cells | Can exosomes from patients with idiosyncratic DILI confirmed by RUCAM be used to more accurately differentiate DILI from other types of liver injury. Can a study of exosomes provide a better fundamental understanding of the mechanisms of idiosyncratic DILI |
| Endotoxins such as lipopolysaccharides have been detected in patients with liver injury unrelated to the use of drugs, and it has been proposed that they are pathogenic. This has not been tested in patients with idiosyncratic DILI. Patients with inflammatory bowel disease do not appear to be at increased risk of idiosyncratic DILI | Do the presence of serum endotoxins indicate that they are important in the pathogenesis of liver injury, or are they simply an indication of decreased liver function with a failure of Kupffer cells to clear them? |
| There is a lack of an animal model that has the human HLA genotype required to mimic the full picture of human idiosyncratic DILI | Can the early innate immune response to drugs that cause idiosyncratic DILI be studied in humans, or even in animals, even though, without the required HLA/T cell receptor repertoire, it does not lead to significant liver injury. If so, this could provide a way to predict that a drug candidate would cause idiosyncratic DILI in some patients |

Table 1 (continued)
There is a strong correlation between reactive metabolite formation and the risk that a drug will cause idiosyncratic DILI. However, it is very difficult to prove that a specific reactive metabolite is responsible for idiosyncratic DILI. If reactive metabolites are responsible for most idiosyncratic DILI, what role do they play: neoantigen formation, production of ROS, other cellular damage leading to the release of DAMPs, etc.? Do enzymes other than CYP such as glucuronyltransferase also play an important role in metabolic activation leading to idiosyncratic DILI?

In addition to hepatocytes, various non-parenchymal cells of the liver and immune cells outside of the liver are considered to be involved in the pathogenesis of idiosyncratic DILI. How can data derived from experimental studies, not from the liver of patients with idiosyncratic DILI, be translated to human DILI in the absence of the required HLA molecules?

Multiple hepatic mediators released from hepatocytes and non-parenchymal cells have been implicated to contribute to idiosyncratic DILI development. The abundance of mediators and resulting hypotheses is challenging; how valid is it to translate results from mostly in vitro studies to human idiosyncratic DILI?

There are abundant publications that propose various mechanisms of idiosyncratic DILI. How can mechanistic hypotheses be rigorously tested?

There are multiple mechanistic studies, many in vitro and at high drug concentrations. Other studies involve drugs or chemicals that are intrinsically toxic. Many of the results from these studies are not reliable indicators of the mechanism of DILI in humans. Mechanistic hypotheses must be consistent with the characteristics of idiosyncratic DILI in humans, and whenever possible, mechanisms should be tested in humans.

BSEP, bile salt export pump; CYP, cytochrome P450; DAMPs, danger-associated molecular pattern molecules; DILI, drug-induced liver injury; HLA, human leucocyte antigen; ROS, reactive oxygen species; RUCAM, Roussel Uclaf Causality Assessment Method.

The human study model of idiosyncratic DILI

Idiosyncratic DILI mechanistic steps are best analyzed using patients with their unique genetic profile and other unknown risk factors instead of experimental models that lack these features. Inclusion in study cohorts will require patients with the diagnosis of real idiosyncratic DILI verified by a robust causality assessment method (CAM) such as the Roussel Uclaf Causality Assessment method (RUCAM), established in 1993 (33) and now the preferred version updated in 2016 (34). This will ensure cohort homogeneity through exclusion a priori cases with alternative causes unrelated to drug use. Indeed, it was early recognized that up to 47% patients with the initial diagnosis of idiosyncratic DILI had been wrongly diagnosed in face of overt alternative causes (35), findings in principle, confirmed in subsequent reports (28,29).

A prerequisite for evaluating mechanistic steps is the use of patients with high RUCAM based causality gradings of probable or highly probable. At the present time RUCAM cannot be replaced by any of the multiple diagnostic biomarkers that have been heavily promoted by interested parties, because virtually all biomarkers available on the market came under scientific fire recently due to misconducted studies carried out by groups outside the European Medicines Agency (EMA) (36,37). In more detail, EMA provided a statement on 17 April 2019 via internet that the earlier issued “Letter of Supprt for drug-induced liver injury (DILI) biomarker (EMA/4238702016) had been retracted from the EMA website on 15 April 2019. The decision was based on information received by the IMI TransBioLine consortium, which is a successor of the former SAFE-T consortium, which itself was the applicant of the “Letter of Support” issued in 2016. The consortium indicated that, due to a case of scientific misconduct at one of the collaboration partner centers of the consortium, the IMI TransBioLine consortium is no longer recommending the exploratory use of hyperacetylated HMBGB1 isoforms in clinical studies. The overall promising nature of other recommended biomarkers was considered to be highly dependent on the results for the incriminated biomarker HMGB1. The CHMP/EMA has therefore decided to retract this Letter of Support that affected many other biomarkers that had
been initially supported (37). A detailed discussion on this subject was published earlier (36).

**DILI liver as a secret keeper organ**

The liver of patients with idiosyncratic DILI is not readily accessible for investigations exploring the initial steps triggering the development of liver injury. In particular, invasive diagnostic procedures such as liver biopsies are not practicable and must be declined for ethical reasons considering the lack of any significant benefit to the patient. Liver histology is also not part of the diagnostic algorithms of RUCAM (34). Instead, we must rely upon biomarkers released into body fluids, usually blood.

**Serum enzyme activities and bile acids**

In patients with idiosyncratic DILI, a battery of laboratory parameters, including aminotransferases, merely signify the existence of a liver disease (37). More specifically, increased serum activities of glutamate dehydrogenase (GDH) would provide circumstantial evidence of an involvement of mitochondria in liver injury, whereas increased serum bile acids (BA) would reflect drug associated injury of the biliary system (37). However, these two parameters do not necessarily allow any conclusions on the underlying mechanistic steps leading to injury. These, and other laboratory parameters, indicate different types of liver injury: one is for hepatocellular injury and the other one for cholestatic injury (34). It is obvious that with at least 2 different liver injury types, a uniform hypothesis based on a single mechanistic step may not be feasible. Also in face of the variability of liver histology that includes, for instance, steatosis, granulomatous hepatitis, vanishing bile duct syndrome, and hepatic sinusoidal obstruction syndrome (HSOS) (38), one single mechanism may not be able to explain the liver histology of idiosyncratic DILI. However, if idiosyncratic DILI is mediated by the adaptive immune system, it is common for different individuals to have quite different responses. For example, the response to SARS-CoV-2 and the manifestations of the infection varies widely between different individuals. Some individuals become quite ill including a “cytokine storm”, while others have little or no symptoms; some individuals lose the sense of smell while others do not, some have gastrointestinal symptoms or strokes while others do not. Such variations in individual responses are common, especially when the immune system is involved.

**Blood exosomes**

Exosomes found in the blood of patients with various liver diseases have provided an important development in the study of liver injury. It has been proposed that they have the potential to serve as diagnostic markers for specific liver diseases (39,40). In addition, they likely play an important mechanistic role in communication between the liver and the immune system. They have been studied both in patients and in animal models of liver injury (39-50). Belonging to the group of extracellular vehicles comprising also microvesicles and apoptotic bodies (40), exosomes represent vesicles, which are commonly derived from cell membranes of various organs including the liver (39). Exosomes released from the liver may be traced back to hepatocytes, hepatic stellate cells, and immune cells in normal and pathological conditions (39,48). They carry lipids, proteins, coding and non-coding RNAs, and mitochondrial DNA (48). Most important for the mechanism of immune adaptation and immune idiosyncratic DILI, exosomes carry drug-modified proteins that are taken up by antigen presenting cells (48). In these studies, however, the exosomes did not lead to activation of the antigen presenting cells; therefore, in these cases immune tolerance may prevail rather than immune reactions associated with idiosyncratic DILI. In other studies, blood exosomes were detected in animals with liver injury caused by CCl₄ (50), but they have not yet been studied and confirmed using the recently published animal model of CCl₄ liver injury (7). In addition, they have not been examined in patients with liver injury following intoxication by CCl₄ (8,9). Exosomes were also found in the blood of patients with alcoholic liver disease (ALD) and in animals with experimental alcoholic liver injury (4,41,42). These blood exosomes contain CYP 2E1 originating from the injured liver where this CYP isoform is part of the hepatic microsomal ethanol oxidizing system (MEOS), which contributes to ethanol metabolism and the associated liver injury (1,2,4,51). Increased exosomal CYP 2E1 is considered as a possible diagnostic biomarker in the blood of humans with alcoholism and microsomal stress. These exosomes may help disclose mechanistic steps leading to alcoholic liver injury, and they could assign alcohol as a cause in a liver disease of unknown etiology (4,42). Exosomes are also released into the blood from the liver of mice treated with acetaminophen (chemically known as N-acetyl-p-aminophenol; APAP) leading to experimental intrinsic DILI (41,42,45). However, their diagnostic or mechanistic role in human intrinsic HILI by APAP remains...
to be established (42,44,45,49). Surprisingly, no data on blood exosomes in patients with idiosyncratic DILI have been published (38-40,42-50). Instead, experimental studies only showed that exosomes originating from hepatocytes are closely related to exosomes derived from cholangiocytes that can affect various signaling pathways (47). Whether these connections explain cholestatic injury in addition to hepatocellular injury in human idiosyncratic DILI remains to be established. Future studies may be useful for elucidating initial mechanistic steps of liver injury (39,40,42-50).

### Blood microRNA

MicroRNAs originating mainly from the liver are found in the blood of patients with liver diseases, either included in exosomes or freely floating (41,52). They were promoted not only as diagnostic biomarkers (27,52-60), but also as mechanistic biomarkers providing insights into to pathogenetic steps in liver injury (61,62). However, the enthusiasm for their use as diagnostic biomarkers was not shared by others because they lack superiority over existing parameters (63). In addition, there were methodology shortcomings (64). In particular, the combination of diagnostic biomarker qualities and idiosyncratic DILI is problematic because many biomarkers were not validated using cases of idiosyncratic DILI assessed for causality by RUCAM with causality gradings limited to highly probable or probable ones (36,64). In addition, most biomarkers have been described for human or animal intrinsic DILI caused by APAP overdoses, providing results that are not necessarily suitable for translation to human idiosyncratic DILI (27,36,52-61). Of note, blood microRNA was analyzed in capillary blood of RUCAM based cases with intrinsic DILI caused by APAP and in one single case associated with the use of clarithromycin causing idiosyncratic DILI (52), but no studies were carried out in patients with idiosyncratic DILI by drugs others than clarithromycin (61). Respective data were also not published for the 46,266 cases of RUCAM based idiosyncratic DILI (23). To date, it appears that microRNA analyses have failed as a diagnostic biomarker of idiosyncratic DILI.

With respect to microRNA, other problems emerged (23,36). Triggered by promotional statements issued by regulators and consortia, a scientific and clinical biomarker hype emerged in 2016 following EMAs online presentation of a Letter of Support to use several diagnostic biomarkers including microRNA in order to verify or exclude liver injury cases as summarized recently (36). However, on 15 April 2019, confusion emerged due to the EMA issuing a retraction note regarding microRNA and various other potential biomarkers due to external data manipulation. This led to a dramatic scientific and regulatory dilemma because previously published analyses and recommendations now require reconsideration. The website with the retraction notification remained accessible only for a short period before it was removed from the internet. The initial letter of support was also removed. As a consequence, related regulatory or consortia Letters of Recommendations previously provided by the FDA and SAFE-T (Safer and Faster Evidence-based Translation) Consortium also disappeared from their websites. The lack of accessibility of the retraction notices likely explains why some reports on microRNA, other biomarkers, and DILI published after April 2019 did not consider this new state of affairs. It is also obvious that data derived from blood microRNA cannot be used to identify the multiple mechanistic steps in idiosyncratic DILI.

### Gut microbiome and blood lipopolysaccharides (LPS)

There is a significant immunological interaction between the gut and the liver (65). For instance, evidence exists that the gut microbiome plays an important role in several types of liver injury. In particular, the presence of specific gut bacteria is associated with a decrease in the autoimmune reactions caused by immune checkpoint inhibitors used to treat cancer (66). There is also evidence that endotoxins, such as LPS derived from the gut microbiome, enter the hepatic blood vessels through a leaky gut, and if not cleared by the liver, reach the systemic circulation where they can be quantified (66-69). They are under consideration as toxins for a variety of human and experimental liver injuries (2,4,24,68,70-72) caused, for instance, by alcohol (2,4,72), CCl\(_4\) (68,70), or APAP overdoses leading to intrinsic DILI (71). No published data on blood LPS are available in patients with idiosyncratic DILI when reports up to 2014 were evaluated (56). Findings of missing LPS data were currently confirmed using the PubMed database for search of respective reports published until mid of April 2020. Seemingly a neglected topic, the lack of LPS data requires further investigations in patients with idiosyncratic DILI with high causality gradings based on the use of RUCAM. As it presently stands, LPS cannot assist in clarifying mechanistic steps in human idiosyncratic DILI. In particular, the
incidence of idiosyncratic DILI does not appear to be higher in patients with inflammatory bowel disease even though the liver in such patients is exposed to much higher levels of LPS and other inflammatory molecules.

**Blood monocytes**

Macrophages and monocytes play a critical role in the control of immune responses. Therefore, it seems plausible that they would play an important role in the mechanism of idiosyncratic DILI. Studies have shown that they even play a role in acetaminophen-induced DILI (73). In one study, peripheral blood monocytes from patients suspected to have diclofenac-induced DILI were cultured under conditions to convert them to “hepatocyte-like” (MH) cells (74). These cells increased the expression of Integrin beta 3 (ITGB3) when incubated with diclofenac. Cases were partially assessed for causality by the updated RUCAM (34), but some methodology uncertainties remained (23,36). Integrin beta 3 (ITGB3) derived from MH cells was also promoted as a biomarker candidate for idiosyncratic DILI (34), but requiring conformation by peers as outlined previously (23,36). Studies on MH cells and ITGB3 were published in 2018 based on Letters of Recommendations presented by EMA through IMI projects SAFE-T and MIP-DILI and supported by the US FDA (34). However, a few months later these parties revoked their recommendations in 2019 (36). In addition, the relationship between MH cells and hepatocytes as well as the steps leading to the production of MH cells are seemingly vague (74). At the present time, it is not possible to be confident of the relationship between ITGB3 produced by blood-derived MH cells and the mechanism of idiosyncratic DILI.

**Hepatic immune system**

Compelling evidence exists that for most idiosyncratic DILI cases the hepatic immune system plays a prominent pathogenetic role (24,56,74-77). A review of this evidence is beyond the scope of this paper, but is presented in a recent review (24). In short, a consensus exists that the hepatic immune system is involved in DILI caused by many drugs (24,56,74-77). Although there are multiple lines of evidence for an immune mechanism of most IDILI, and clinical evidence for alternative mechanisms is scant, given that biological systems are very complex, it is possible that some IDILI involves nonimmune mechanisms. In most cases the injury appears to be mediated by CD8 T cells of the adaptive immune system, which requires prior activation of the innate immune system. Early steps in this process likely involves activation of antigen presenting cells by molecules such as danger associated molecular pattern molecules (DAMPs) (24). Support for an involvement of the immune system in idiosyncratic DILI is provided by autoimmune parameters in the blood of patients (20) and clinical features, liver histology, and in some cases with human leucocyte antigen (HLA) genotypes (24).

An association of HLA genes as risk factor for idiosyncratic DILI caused by selected drugs has been described in a variety of reports (17,20,21,24), but their clinical utility for use as a pre-prescription screening tool with potential liver injury remains unclear (24). Pharmacogenetic HLA associations were found in a few, but not all, patients under a treatment with a few drugs including abacavir, amoxicillin clavulanate, fluvoxacinil, isoniazid and other antituberculosis drugs, lumiracoxib (21), and other drugs (20). In addition, for many published DILI cases, an association with HLA genes has been assumed, but it is unclear whether the liver injury was induced by drugs or due to alternative causes (17,21-23,25,26,28,29,35) not carefully excluded by the use of a strong causality assessment tool such as the updated RUCAM (34). Concern also exists that for most drugs, there are too few cases available for investigators to evaluate (24). Although it is likely that most idiosyncratic DILI is immune mediated, the details of how a drug induces an immune response leading to DILI are unknown and likely different for different drugs (24,78). Among the possible general mechanisms are that reactive drugs such as ß-lactams or reactive metabolites can act as hapten to form neoantigens, and they can also cause cell damage leading to the release of DAMPs and activate inflamasomes. There is a large amount of circumstantial evidence for the involvement of reactive metabolites in the mechanism of idiosyncratic DILI; however, it is very difficult to prove that a specific reactive metabolite is responsible for human idiosyncratic DILI caused by a specific drug. Drug metabolism occurs in the liver preferentially via CYP (79,80) and to a lesser extent through non CYP pathways (81). In particular, aryl glucuronides have been proposed to be responsible for DILI caused by carboxylic acids such as diclofenac; however, the evidence is far from compelling (82).

**Hepatic cytochrome P450 and oxidative stress**

CYP is primarily localized in the liver and degrades many
drugs to harmless metabolites; however, it also has the potential to produce toxic metabolites (79,80), which have the potential to initiate idiosyncratic DILI. To further evaluate this issue, top drugs causing idiosyncratic DILI as assessed in 48 cases worldwide with verified causality by RUCAM were analyzed for their possible metabolism via (83-100) CYP (101-118) (Table 2). In at least 28/48 drugs (58.3%), clinical or experimental evidence exists that metabolism proceeds via CYP, whereas for the remaining 20 drugs (41.7%) there were negative or missing results implicating CYP in the metabolism of these drugs (Table 2).

The CYP dependent metabolism of drugs and other

| Drug                     | RUCAM based DILI cases (n) | Substrates of CYP | References        |
|--------------------------|---------------------------|-------------------|-------------------|
| 1. Amoxicillin-clavulanate | 333                       | CYP –             | Hautekeete (97)   |
| 2. Flucloxacillin         | 130                       | CYP +             | Dekker (91)       |
| 3. Atorvastatin          | 50                        | CYP +             | Zanger (116)      |
| 4. Disulfiram             | 48                        | CYP +             | Hopley (99)       |
| 5. Diclofenac            | 46                        | CYP +             | Zanger (116)      |
| 6. Simvastatin           | 41                        | CYP +             | Fatunde (96)      |
| 7. Carbamazepine         | 38                        | CYP +             | Zanger (116)      |
| 8. Ibuprofen             | 37                        | CYP +             | Hopley (99)       |
| 9. Erythromycin          | 27                        | CYP +             | Hopley (99)       |
| 10. Anabolic steroids    | 26                        | CYP +             | Yamazaki (114)    |
| 11. Phenytoin            | 22                        | CYP +             | Hopley (99)       |
| 12. Sulfamethoxazole/trimethoprim | 21     | CYP +             | Hopley (99)       |
| 13. Isoniazid            | 19                        | CYP +             | Hopley (99)       |
| 14. Ticlopidine          | 19                        | CYP +             | Hopley (99)       |
| 15. Azathioprine/6-mercaptopurine | 17     | CYP –             | Johansson (100)   |
| 16. Contraceptives       | 17                        | CYP +             | Scott (108)       |
| 17. Flutamide            | 17                        | CYP +             | Zanger (116)      |
| 18. Halothane            | 15                        | CYP +             | Zanger (116)      |
| 19. Nimesulide           | 13                        | CYP +             | Yu (115)          |
| 20. Valproate            | 13                        | CYP +             | Kiang (101)       |
| 21. Chlorpromazine       | 11                        | CYP +             | Hopley (99)       |
| 22. Nitrofurantoin       | 11                        | CYP –             | Wang (113)        |
| 23. Methotrexate         | 8                         | CYP –             | Donehower (93)    |
| 24. Rifampicin           | 7                         | CYP –             | Acocella (83)     |
| 25. Sulfasalazine        | 7                         | CYP –             | Das (90)          |
| 26. Pyrazinamide         | 6                         | CYP –             | Shih (110)        |
| 27. Natriumaurothiolate  | 5                         | CYP –             | Björnsson (88)    |
| 28. Sulindac             | 5                         | CYP +             | Brunell (117)     |
| 29. Amiodarone           | 4                         | CYP +             | Hopley (99)       |

Table 2 (continued)
exogenous chemicals such as ethanol and CCl₄ proceeds in the endoplasmic reticulum of the hepatocyte, which corresponds to the microsomal fraction obtained from liver homogenate after subcellular fractionation using ultracentrifugation (51). CYP is not a single enzyme but consists of several highly polymorphic isoforms (100). The individual steps of the catalytic CYP cycle are complex (119), presented in Figure 1 as a simplified multistep process (4).

The reactions within the CYP cycle start with the binding of a drug as substrate to the ferric heme moiety of CYP-Fe³⁺ (Figure 1). This is followed by the uptake of an electron provided by the NADPH-cytochrome P450 reductase that converts the ferric state of CYP-Fe³⁺ to its ferrous state CYP-Fe²⁺, with subsequent inclusion of molecular oxygen and another electron provided again by the reductase (Figure 1) (4,119). These reactions are supported by

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**Table 2 (continued)**

| Drug                        | RUCAM based DILI cases (n) | Substrates of CYP | References                  |
|-----------------------------|----------------------------|-------------------|-----------------------------|
| 30. Interferon beta         | 3                          | CYP −             | Bertz (87)                  |
| 31. Propylthiouracil        | 2                          | CYP +             | Heidari (98)                |
| 32. Allopurinol             | 1                          | CYP −             | Turnheim (112)              |
| 33. Hydralazine             | 1                          | CYP −             | Talseth (111)               |
| 34. Infliximab              | 1                          | CYP −             | LiverTox (104)              |
| 35. Interferon alpha/ Peginterferon | 1                  | CYP −             | Okuno (107)                 |
| 36. Ketoconazole            | 1                          | CYP −             | Kim (102)                   |
| 37. Busulfan                | 0                          | CYP −             | Myers (105)                 |
| 38. Dantrolene              | 0                          | CYP −             | Amano (84)                  |
| 39. Didanosine              | 0                          | CYP −             | Andrade (85)                |
| 40. Efavirenz               | 0                          | CYP +             | Desta (92)                  |
| 41. Floxuridine             | 0                          | CYP −             | Landowski (103)             |
| 42. Methyldopa              | 0                          | CYP +             | Dybing (94)                 |
| 43. Minocycline             | 0                          | CYP −             | Nelis (106)                 |
| 44. Telithromycin           | 0                          | CYP +             | Shi (109)                   |
| 45. Nevirapine              | 0                          | CYP +             | Erickson (95)               |
| 46. Quinidine               | 0                          | CYP +             | Nielsen (118)               |
| 47. Sulfonamides            | 0                          | CYP −             | Back (86)                   |
| 48. Thioguanine             | 0                          | CYP −             | Choughule (89)              |

Listed are the top ranking 48 drugs worldwide causing idiosyncratic DILI with verified causality using RUCAM, with details presented in a previous publication (22). The references refer to the first author of the study that delineates whether the drug under consideration is a substrate of and metabolized by CYP (CYP +) or not (CYP −). CYP, cytochrome P450; DILI, drug induced liver injury; RUCAM, Roussel Uclaf Causality Assessment Method.

![Figure 1](image-url) **Figure 1** Metabolism of drugs and other substrates through the cytochrome P450 cycle. The figure is derived from a previous report (4).
phospholipids, which are essential components of the microsomal membrane (4), and could be involved as peroxidized phospholipids in the development of the complex consisting of the reduced CYP, an activated form of oxygen, and the substrate (Figure 1). Under conditions of incomplete oxygen splitting, radicals may be generated from oxygen, phospholipids, and drugs, but the individual mechanistic steps occurring in the human liver of patients with idiosyncratic DILI are difficult to assess and are currently unknown. Finally, and under physiological conditions of drug use, the drug is split off as oxidized substrate, whereas the CYP molecule returns from its ferrous state (2') to its ferric state (3') and is again available for binding to the next substrate molecule to keep the CYP cycle running (Figure 1). Clearly, CYP presents with 2 sides of a coin: on the one side, CYP metabolizes drugs and clears them from the body, but on the other side, it produces ROS that may attack membrane structures of cell organelles. The damage produced could lead to direct liver injury, although it is harder to explain why this would be idiosyncratic, and it could also cause the release of DAMPs that promote immune mediated injury in susceptible individuals (24,56,75-77,120,121). The reactive species could also act as haptens that provoke an immune response in susceptible individuals (120).

In addition to CYP in the endoplasmic reticulum, hepatic mitochondria also contain CYP, which possibly contributes to mitochondrial injury as suggested by experiments in alcohol-induced liver injury (4). Furthermore, hepatocytes release exosomes containing CYP as a result of toxicity associated with drug metabolism (122,123), which is analogous to exosomal CYP found in the blood of patients with ALD or animals with alcoholic liver injury (42). However, data from studies of exosomes in the blood of patients with idiosyncratic DILI containing CYP are not available that could be used to substantiate a mechanistic role of CYP in this disease (43). In idiosyncratic DILI caused by a few drugs, anti-CYP can be detected in the blood, implying a close association between the incriminated drug and CYP (124-126) in line with the proposed involvement of the adaptive immune system (24). Many questions relating to an association between CYP polymorphisms and the risk of idiosyncratic DILI remain unanswered (24,127).

Cellular oxidative stress in idiosyncratic DILI includes mitochondrial stress (128,129) and microsomal stress (129,130), and this may be related to ROS generated by CYP in these subcellular domains of the hepatocytes (128-130), but specific ROS types involved in human idiosyncratic DILI have not yet been identified. For alcoholic liver injury, which involves the MEOS and is dependent on CYP and NADPH-CYP reductase, the following radicals have been implicated: singlet radical \( \cdot O_2 \), superoxide radical \( \cdot O_2^- \), hydrogen peroxide \( H_2O_2 \), hydroxyl radical \( \cdot OH \), alkoxyl radical \( RO^- \), and peroxyl radical \( ROO^- \) (4). Involvement of these radical types in the mechanism of idiosyncratic DILI is possible; however, despite multiple efforts, a satisfactory unifying mechanistic concept for idiosyncratic DILI with the involvement of various ROS types is currently not available.

### Hepatic non CYPs

There are some drugs that cause idiosyncratic DILI validated by RUCAM, which do not appear to be metabolized by CYP (Table 2). In some cases, such as ß-lactams and busulfan it is likely because the drug is intrinsically reactive. In other cases such as methotrexate, the parent drug is toxic. In some cases, such as nitrofurantoin and dantrolene, reactive metabolites are formed by reduction rather than oxidation. In still other cases other enzymes may oxidize the drug, and it is unclear whether the generation of ROS may be involved (81,131). Among these enzymes are aldehyde oxidase present in the cytosol, carbboxylesterase, and UDP-glucuronosyltransferase (131). Despite FDA guidance documents that specify that acyl glucuronides are toxic (132), clear evidence that they are involved in the mechanism of idiosyncratic DILI is lacking. It will be very interesting to find out whether new chemical entities in the drug pipeline that are not metabolized by CYP are associated with a significant risk of idiosyncratic DILI. In particular, there are many new biological drugs that cause idiosyncratic DILI, and the mechanism presumably does not involve metabolism by CYP (133).

### Animal models

Given the unpredictable nature of idiosyncratic DILI, it is impossible to prospectively study patients who will be affected. Animal models are very important for most biomedical research; however, idiosyncratic DILI is also idiosyncratic in animals. Therefore, it is difficult to develop animal models in which the incidence is sufficient to be practical for research. Most animal models involve high doses of the drug and/or other manipulations that lead to direct and immediate liver injury, and in general the
histology is different (24,134,135). These characteristics are different from the characteristics of idiosyncratic DILI; therefore, the mechanisms are almost surely different. Animals do not have the same HLA genotypes as those associated with an increased risk of DILI associated with specific drugs in humans; therefore, it would be impossible to develop an animal model to drugs that require a specific HLA genotype. However, not all idiosyncratic DILI appears to be associated with specific HLA genotypes. This is likely because some drugs react with so many endogenous proteins that there is likely one modified protein, which produces peptides. They in turn may bind to one of the HLA molecules that any individual human or animal may express.

A major development in the treatment of cancer has been drugs, which target immune checkpoints that prevent the immune system from destroying cancers (136). Two important immune checkpoints are programmed cell death protein 1 (PD-1) and cytotoxic T-lymphocyte-associated protein 4 (CTLA-4). The dominant immune response in the liver is immune tolerance. One side effect of these immune checkpoint inhibitors is DILI, and they also increase the risk of idiosyncratic DILI of coadministered drugs (137). The immune response is always a balance between an active immune response and immune tolerance, and immune checkpoint inhibitors shift this balance. Treatment of PD-1−/− mice with a combination of anti-CTLA-4 and amodiaquine (a drug associated with a relatively high incidence of idiosyncratic DILI) leads to a delayed onset liver injury with histology indistinguishable from that of idiosyncratic DILI in humans (138). This injury is mediated by CD8 T cells (139). Although the injury is significant and leads to decreased liver function with an increase in bilirubin, it does not lead to overt liver failure. This model also unmasks the potential of other drugs to cause DILI; however, the liver injury is less than with amodiaquine (140). It is likely that the lack of liver failure is due to the lack of the optimal HLA and T cell receptors required to produce the maximal amount of liver injury. In fact, drugs that cause serious liver injury are always associated with a higher incidence of mild liver injury, which often resolves with continued drug treatment. It is likely that most humans and even many animals have an innate immune response to drugs that can cause idiosyncratic DILI, but without the required HLA and T cell receptors, no adaptive immune response leading to injury occurs. A study of the innate immune response to drugs could provide a method to test the early immune events in idiosyncratic DILI and even a mechanism to screen drug candidates for the risk that they will be associated with a significant risk of DILI (24).

Non parenchymal cells and mediators

Given that the major site of reactive metabolite formation is hepatocytes, they are an important target of idiosyncratic DILI. However, there is a large amount of communication between hepatocytes and other cells in the liver, including Kupffer cells, hepatic stellate cells, and liver sinusoidal endothelial cells. In addition, immune cells in the liver and outside of the liver presumably also play important roles in the mechanism of idiosyncratic DILI. Injury to the liver usually causes a decrease in the number of Kupffer cells, but an increase in infiltrating monocyte-derived macrophages, which display a large amount of heterogeneity and change as the injury evolves (78). The number of publications on various pathogenic aspects in idiosyncratic DILI is impressive (17,21,24,32,61,75–77,121,127,128,130). The number of various hypotheses is not surprising given the variability of clinical features and the difficulty in performing rigorous mechanistic studies in humans. Therefore, it requires extrapolation from other types of experiments such as in vitro studies, studies with intrinsically toxic agents, and experiments using extreme conditions. In line with these considerations and restrictions, the conclusion can be reached that a satisfactory evidence for a unifying mechanism for individual susceptibility, initiation, and progression of idiosyncratic DILI is not available.

Hepatocellular injury versus cholestatic injury

The majority of idiosyncratic DILI presents clinically as hepatocellular injury; cholestatic injury is less common (19). This is why most studies have focused on the hepatocellular type rather than the on the cholestatic injury (24). Circumstantial evidence suggests that the hepatocellular injury with its increased serum activities of ALT, AST, and GDH can be attributed to a variety of mechanistic processes that include immune and non immune systems, CYP and non CYP pathways, and mitochondria and non mitochondria targets of the hepatocytes. Conversely, pathogenetic steps leading to the impairment of the bile salt export pump (BSEP) of the hepatocyte as a cause of the cholestatic injury is evidenced by increased serum alkaline phosphatase (ALP) activities. Increased serum bile acid levels is less studied, but some details warrant mentioning (24). In particular, the composition of bile acids is different in rodents than
in humans with a higher percentage of the more polar and
less toxic taurine conjugates in mice and more of the toxic
glycine conjugates in humans (3,24). This complicates
the development of a valid rodent model to study BSEP
inhibition or other bile acid transporters, and it makes it
more difficult to test the hypothesis that BSEP inhibition
leads to cholestatic idiosyncratic DILI (24). Thus it remains
to be established which mechanism triggers the cholestatic
injury. However, it is likely that transport and concentration
of a drug or its metabolites in the biliary system plays an
important role.

Conclusions

Idiosyncratic DILI is a multifaceted disease that provides
a large opportunity to resolve basic mechanistic issues. Clinical features are well described in cases assessed for
causality using RUCAM, but a large portion of mechanistic
steps triggering the liver injury remain unclear in face of both the behavior of the human DILI liver as a secret
keeper organ and the lack of an appropriate animal model
with genetic specificities of susceptible patients with
idiosyncratic DILI. Sufficient evidence exist that the hepatic
adaptive immune system mediates most of the liver injury
cases, a process requiring stimulation of the innate immune
system and transition to an adaptive response, but the events
facilitating the stimulation remain unclear. Additional
gaps of understanding relate to the failure to detect in the
blood of patients with idiosyncratic DILI specific exosomes
released from the injured liver that may carry information
on coding and non-coding RNAs, mitochondrial DNA, and
CYP isoforms that might help clarify initial mechanistic
steps of injury. Data on patients’ exosomes have not been published, either because they were not analyzed, or the
levels were too low to provide a clear picture. Similarly, patients’ blood microRNA has not been helpful to clarify
existing mechanistic issues, and at present they represent
a now outdated biomarker after losing regulatory support
by EMA and the FDA following detected misconduct of
an expert team. The current analysis also revealed that
most of the drugs causing liver injury are metabolized
by cytochrome P450 and NADPH cytochrome P450
reductase. These microsomal constituents may generate
toxic ROS as by products with the potential to initiate liver
injury, but evidence is only circumstantial. Given that drugs
can target all non parenchymal cells such as Kupffer cells,
hepatic stellate cells, liver sinusoidal endothelial cells, as well
as intrahepatic granulocytes, lymphocytes, and monocytes,
this makes it more difficult to define their role and the impact
of active mediators for signaling and mechanistic pathways
in human idiosyncratic DILI. In essence, from a scientific
point of view, it would be desirable to elucidate the cascade
of events with identification of each mechanistic step. A
better understanding of the mechanisms of idiosyncratic
DILI could lead to ways to prevent or treat it. However, at
the present time and for the sake of patients, it is important
to recognize early symptoms of idiosyncratic DILI and
to quickly establish the diagnosis using a robust method
for causality assessment such as the updated RUCAM
assessment with the aim to discontinue the use of the
incriminated drug.

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