Drug-Induced Liver Injury in GI Practice

Naemat Sandhu and Victor Navarro

Although drug-induced liver injury (DILI) is a rare clinical event, it carries significant morbidity and mortality, leaving it as the leading cause of acute liver failure in the United States. It is one of the most challenging diagnoses encountered by gastroenterologists. The development of various drug injury networks has played a vital role in expanding our knowledge regarding drug-related and herbal and dietary supplement–related liver injury. In this review, we discuss what defines liver injury, epidemiology of DILI, its biochemical and pathologic patterns, and management. (Hepatology Communications 2020;4:631-645).

A

lthough drug-induced liver injury (DILI) is a rare clinical event, it carries significant morbidity and mortality, leaving it as the leading cause of acute liver failure (ALF) in the United States. It is one of the most challenging diagnoses encountered by gastroenterologists. DILI is also the most common single adverse event that has led to withdrawal of drugs from the marketplace, drug attrition, and failure of implicated drugs to obtain U.S. Food and Drug Administration (FDA) approval.

Defining DILI

Liver injury is recognized by abnormal liver biochemistries with or without associated clinical symptoms. Using liver biochemistry criteria, which will increase the specificity of hepatotoxicity causality assessment and eliminate false positives, is key. This aids in early detection, prediction, and risk stratification of suspected cases of DILI. The updated Roussel Uclaf Causality Assessment Method (RUCAM) uses an alanine aminotransferase (ALT) >5-times the upper limit of normal (ULN) and/or alkaline phosphatase (ALP) >2-times ULN to identify liver injury. Conventionally, liver biochemistry elevations to this degree, lesser elevations that are sustained over time, rapidly rising tests, or any elevation combined with signs of liver dysfunction, such as increase in the international normalized ratio or encephalopathy, are clinically significant and worthy of investigation.

Burden of DILI in the United States and Abroad

In western countries, acetaminophen (APAP)-related liver injury remains one of the leading causes of DILI. Given the challenges in detection and reporting, the incidence of DILI is difficult to ascertain. Annual incidence of DILI ranges from 2.3-13.9/100,000 inhabitants in population-based studies from Europe. The highest
incidence of non-APAP-related DILI was reported at 19.1/100,000 inhabitants/year with a steady increase in age-standardized incidence of DILI, in an Icelandic population-based study.\(^9\) Although most of the cases reported in the western countries are DILI secondary to prescription medications, traditional/complimentary and dietary supplements are the main causative agents of DILI in Asia.\(^{10,11}\) In the only U.S. population-based study, the yearly incidence of DILI was found to be approximately 3/100,000 residents.\(^{12}\)

**U.S. DILI Network**

The National Institute of Diabetes and Digestive and Kidney Diseases established the U.S. DILI Network (DILIN) in 2003 to identify, enroll, and characterize cases of non-APAP DILI and herbal and dietary supplements (HDSs).\(^{13}\) The U.S. DILIN has two registry studies at eight different academic centers across the United States. The network has expanded our understanding of DILI. Antimicrobials were noted to be the most common causative agents, accounting for 45% of cases in a 2004 study, followed by HDSs, cardiovascular drugs, central nervous system agents, antineoplastic agents, and analgesics.\(^{14}\) Of the antimicrobials, amoxicillin-clavulanate (22%), isoniazid (11.7%), and nitrofurantoin (10.2%) were the top three implicated agents.\(^{14}\) The network has also noted an increasing proportion of HDS-related liver injury, from 7% in 2004-2005 to 20% in 2013-2014.\(^{15,16}\)

**Patterns and Outcomes of DILI**

Recognizing the pattern of liver injury at the initial presentation is vital. It provides a useful foundation to establish a differential diagnosis and guides the diagnostic evaluation accordingly. Assessing the pattern of liver injury as hepatocellular, cholestatic, or mixed is based on which liver enzyme elevation predominates. For example, hepatocellular injury suggests that an elevation of ALT and/or AST is more prominent than ALP. Conversely, cholestatic injury suggests a predominant elevation of ALP. The R-ratio is a quantitative expression of the injury pattern; it is defined as the ratio of serum ALT to ALP values, both expressed as multiples of ULN, obtained at the onset of injury. An R-ratio of >5 indicates hepatocellular injury, <2 indicates cholestatic injury, and 2-5 indicates mixed injury.\(^{17,18}\) Table 1 lists the drugs and pattern of associated liver injury.

Hepatocellular injury is the most common pattern of DILI across various networks, with 52%-75% of cases reported from Spanish, Latin, and U.S. DILIN, and in the Swedish Adverse Drug Reaction Advisory Committee (SADRAC).\(^{14,19-21}\) Patients with hepatocellular injury tend to be younger, less likely to be clinically jaundiced, but have higher frequency of liver-related deaths. Hepatocellular injury is 2–3 times more likely to lead to liver transplantation. The clinical course tends to be more protracted in those with cholestatic injury.\(^{14}\)

Hepatocellular DILI with jaundice is an important pattern of biochemical injury to recognize, commonly referred to as “Hy’s Law.”\(^{22,23}\) It was first observed by Hyman Zimmerman in an analysis of 114 patients taking isoniazid, patients with an increased ALT >3-times ULN, and total bilirubin >2-times ULN without initial findings of cholestasis and after excluding other potential causes experienced a case fatality rate of 10%.\(^{24}\) This observation has been used by the FDA to identify drugs with the potential of severe liver injury. Similar fatality rates from DILI with hepatocellular jaundice have been seen in the Spanish DILIN (11.7%) and SADRAC (12.7%).\(^{13,25,26}\)
Adaptation is a transient and modest rise of liver tests due to a drug that persists without progression, or regresses back to normal despite continued use of a drug thought to be the cause. The common examples of liver adaptation are statins and isoniazid, although both agents have also been implicated in cases of idiosyncratic DILI. (27)

From a clinical standpoint, the development of low-level liver enzyme elevation (ALT < 3-times ULN or direct bilirubin <2-times ULN) in an otherwise asymptomatic patient presents an important challenge. The clinician must also weigh the risks and benefits of a drug for the patient, as the benefit of a medication may outweigh the risk of hepatic injury. In this context, if low-level enzymes are detected that, in the clinician’s assessment, are due to a drug, the patient has no liver-related symptoms, and the drug is of evidence-based clinical value to the patient, the enzymes should be followed monthly for at least 3 months to confirm their nonprogressive nature. A concurrent hepatic ultrasound to detect drug-related morphological alterations such as steatosis is also wise. If the enzymes’ direct bilirubin rises, or new steatosis is detected, an alternative agent should be sought, even if within the same class of drugs. If no better alternative exists, such as may be the case with some chemotherapies, dose reduction or drug continuation with close enzyme and imaging follow-up is recommended.

Most patients with DILI have both clinical and biochemical recovery. However, a small proportion of cases may develop chronic liver disease or chronic DILI, conventionally defined as persistence of liver-enzyme elevations for more than 6 months after withdrawal of an offending drug. The incidence of chronic DILI varies from 5%-20% in various DILI registries and population-based studies. (9,28,29)

### Types of DILI

Conventionally, DILI is classified as intrinsic and idiosyncratic. Intrinsic DILI is predictable, occurring in a dose-dependent manner. The typical example of an intrinsic DILI is APAP-related liver injury. The injury is usually within a short duration of time with a predictable latency period after drug exposure. APAP-related liver injury alone is responsible for about 46% of the cases of ALF in the United States. Pharmacodynamic studies have demonstrated a dose-response relationship between the extent of biochemical abnormalities and dose of APAP ingested. At higher doses of APAP, the conjugation pathways for its metabolism are saturated, thereby generating N-acetyl-p-benzoquinone-imine, a highly reactive toxic metabolite leading to hepatocellular necrosis. (30)

Idiosyncratic DILI (IDILI) is unpredictable and, arguably, may be the most common type of drug-associated hepatotoxicity. It has a variable latency period and is difficult to replicate in experimental models. Historically, IDILI has been considered to be dose-independent. In the last decade, however,
there have been studies showing an association of IDILI with drugs prescribed at a daily dose of more than 50 mg.\textsuperscript{(31,32)} Whether there is true association between dose and IDILI is not fully clear. The pathogenesis of idiosyncratic DILI can either be related to the pathway of metabolism of the drug and/or activation of the immune system, as will be discussed subsequently.

### Mechanism of Action

Direct hepatotoxicity occurs in the setting of a known hepatotoxic agent, which leads to cell death through necrosis or apoptosis. However, the mechanism of IDILI involves a complicated process involving the drug, its metabolites, and the host immune system. Most drugs implicated in DILI are lipophilic and metabolized by the liver. They undergo phase 1 reaction, which is usually mediated by the hepatic cytochrome p450 system. The generation of intermediate bioactive or reactive metabolites is an important step, leading to both intrinsic and idiosyncratic DILI.\textsuperscript{(33)}

These toxic intermediate products are usually inactivated by phase 2 reactions, such as glutathione or sulfate conjugation. If conjugation pathways are overwhelmed by the excess production of the toxic reactive metabolites or caused by depletion of the conjugating factors, covalent binding of the reactive metabolites to mitochondrial proteins results. This leads to production of reactive oxygen species and ATP depletion, causing cellular organelle dysfunction from activation of stress kinase signaling pathways and disruption of membrane permeability pores. The result is hepatocyte dysfunction, necrosis, and/or cell death.\textsuperscript{(33,34)}

Immune-mediated injury is an important mechanism for IDILI. Both the innate and the adaptive immune system play a vital role. Over the years, there have been various hypotheses for the mechanism of immune-mediated DILI. Under the hapten hypothesis, reactive metabolites irreversibly bind with cellular proteins to form neoantigens (also referred to as “haptens”), which then are presented to major histocompatibility complex molecules on antigen-presenting cells, triggering an immune response against the hepatocyte by recruitment of cytotoxic T lymphocytes, natural killer cells, and B cells. In some instances, the hapten can induce development of autoantibodies against cytochrome p450 enzymes, leading to cellular injury and death, as in the case of halothane-related liver injury.\textsuperscript{(34-36)} The pharmacological interaction concept proposes that a drug or its metabolite can bind directly to the human leukocyte antigen (HLA) molecule to trigger a T cell–mediated injury, especially in genetically susceptible individuals.\textsuperscript{(37)}

The liver has a remarkable capacity for immune tolerance. This is a necessary adaptation that serves as a barrier to hepatocyte dysfunction and injury from an inflammatory state, which develops in response to the constant exposure to ingested antigens. This is managed by induction of peripheral immune tolerance to incoming antigens.\textsuperscript{(38,39)} The adaptive immune response can be up-regulated in the setting of a sublethal stress from a hapten. HLA associations are indicative of the adaptive immune response in IDILI, and in genetically susceptible individuals, overt liver injury may occur as a result of “defective adaptation.”\textsuperscript{(40)} Conceivably, this complex interplay between the immune and nonimmune pathways is responsible for the unpredictability of idiosyncratic DILI. Figure 1 details the mechanism of direct and immune-mediated DILI.

### Risk Factors for DILI

#### PATIENT-RELATED FACTORS

Susceptibility to DILI is influenced by modifiable and nonmodifiable host demographic, clinical, and environmental factors.

#### Age

Various large prospective DILI registries differ on advancing age as a risk factor for DILI. A population-based Icelandic study showed increasing incidence of DILI with age: 5 times higher for patients over 70 years than those between 15 and 29 years. The former group was also noted to have a higher prescription rate of drugs,\textsuperscript{(9)} which could be a potential reason for increased DILI in this group as opposed to advanced age itself. The U.S. and Spanish networks did not identify advancing age as a risk factor for all-cause DILI.\textsuperscript{(14,33)} Increasing age does pose as a risk factor for liver injury from medications like isoniazid, amoxicillin/clavulanate, and nitrofurantoin.\textsuperscript{(41)} Cholestatic DILI is more common in the elderly as compared with hepatocellular injury in younger
individuals. Therefore, age may confer a susceptibility to DILI in a drug-specific manner.

**Gender**

Although epidemiological data from Spain, the United States, and Iceland do not suggest a heightened risk of DILI in women, others have found women to be more susceptible to DILI from some drugs. These agents include minocycline and nitrofurantoin, in which DILI is typically characterized by autoimmune features. Women with acute liver injury are more likely to progress to acute liver failure, as demonstrated in various registries. The reasons for worse liver-related outcomes and chronicity of DILI in African-Americans in unclear.

**Race**

Race and ethnicity are potential factors influencing DILI frequency, liver injury patterns, and outcomes. In a U.S. DILIN cohort, African-Americans with DILI tended to be younger with higher rates of chronic DILI. The most common agents in African-Americans were trimethoprim/sulfamethoxazole, methyldopa and phenytoin, as compared with amoxicillin/clavulanate in Caucasians. The pattern of liver injury as well as the time of recovery of DILI does not appear to be have significant differences when comparing African-Americans and Caucasians, the predominant pattern being that of a mixed injury. African-Americans were twice as likely to develop severe liver injury, either leading to death or liver transplantation, despite absence of disparity in attaining health care. The reasons for worse liver-related outcomes and chronicity of DILI in African-Americans is unclear. Agents associated with hypersensitivity reactions are known to be the common causes for DILI in African-Americans, which can be as a result of genetic factors. African-Americans have higher risk of severe
cutaneous reactions to allopurinol due to higher frequency of HLA-B*5801.\(^{(48)}\)

**Pregnancy**

Data on DILI during pregnancy is limited and limited to therapeutic agents used to treat gestational hypertension and hyperthyroidism. The most common drugs are methyldopa, hydralazine, propylthiouracil, and antimicrobial agents like tetracycline.\(^{(33,49)}\) It is imperative to differentiate DILI during pregnancy from other more common etiologies of abnormal liver tests including viral hepatitis, gallstone disease, or pregnancy-related complications such as intrahepatic cholestasis of pregnancy.

**Alcohol**

The pathophysiology of APAP-associated liver injury in the setting of alcohol use is complex and driven by the manner in which alcohol is consumed. Acute alcohol co-ingestion with APAP may be protective, as they compete with each other for use of CYP2E1 substrate, in turn reducing the byproduct N-acetyl-p-benzoquinone-imine. In contrast, chronic alcohol use augments APAP hepatotoxicity by acting as a CYP2E1 inducer. Chronic malnutrition, as may occur in patients with alcoholism, may further augment risk for DILI due to glutathione depletion in the malnourished state. In IDILI, there are no data to suggest increased risk of all-cause DILI in the setting of chronic alcohol use. Chronic alcohol consumption is a risk factor for DILI from isoniazid, methotrexate, and halothane.\(^{(50)}\)

**Chronic Liver Disease and Comorbid Conditions**

In a 2015 U.S. DILIN study, patients with pre-existing liver disease were noted to have higher rates of severe DILI and 3-times higher risk of mortality in comparison to those without prior liver disease.\(^{(14)}\) Nonalcoholic fatty liver disease (NAFLD) is also a potential factor to increased risk of DILI\(^{(34)}\); however, it should not preclude the use of statins in patients with NAFLD. In a 2010 *post hoc* analysis of the prospective Greek Atorvastatin and Coronary Heart Disease Evaluation study, those receiving statin therapy had significantly lower rates of cardiovascular events and significant improvement in their liver chemistries.\(^{(51)}\)

**Genetics**

Single nucleotide polymorphisms in numerous genes and HLA regions have been shown to have an increased risk for DILI. A U.K. genome-wide association (GWAS) identified HLA-B*5701 genotype as a major determinant of flucloxacillin DILI.\(^{(55)}\) Another European and U.S. GWAS showed evidence of HLA genotypes HLA DRB1*15:01 and DQB1*0602 playing a significant role in amoxicillin-clavulanate DILI susceptibility.\(^{(56)}\) HLA-B*35:02 has been associated with increased risk of minocycline DILI, with a 16% carrier frequency in DILI cases in comparison to 0.6% in the population control in a GWAS study.\(^{(57)}\) The rarity of occurrence of DILI in relation to a given drug confers a negative predictive value greater than 95% to these HLA alleles. HLA genotyping can potentially increase the accuracy and confidence in diagnosing DILI.\(^{(33)}\)

**DRUG-RELATED FACTORS**

**Dose**

There are data to suggest that drugs with daily oral dosing of ≥50 mg account for 70%-80% of DILI cases in multiple DILI registries and population-based studies.\(^{(9,32,58)}\)

**Drug Metabolism**

Ghabril in 2010 noted oral medications with more than 50% hepatic metabolism had significantly higher frequency of ALT >3-times ULN, liver failure, and fatal DILI. In the same study, drugs with biliary excretion had significantly higher frequency of jaundice. Medications with more than 50% hepatic metabolism and dose more than 50 mg/day were noted to have an additive effect, leading to higher risk of hepatotoxicity.\(^{(59)}\)

**Lipophilicity**

Oral medication with high lipophilicity measured as LogP is known to influence drug pharmacokinetics...
and toxicity. Drugs with higher lipophilicity have an increased volume of distribution, higher chances of off-target binding, and increased risk of toxicity with subsequent generation of increased reactive metabolites. A drug’s hepatotoxic potential has been conceptualized as the “rule of two,” characterized by high lipophilicity (LogP > 3) and daily dosing (>100 mg). These parameters have been associated with an increased risk of DILI.

Making a Diagnosis

DILI is a challenging and complex diagnosis. Given the overlap between presentation of DILI with both acute and chronic liver diseases, it remains a diagnosis of exclusion. Obtaining a detailed history and recognition of the pattern of liver injury is key. Using the R-ratio at presentation assists in a careful selection of diagnostic tests to evaluate for competing diagnoses: hepatic or systemic. In addition to a patient’s demographic variables, obtaining a full medication history is vital. Given the potential for prolonged latency periods, it is imperative to corroborate a patient’s medication list from their pharmacy and investigate over-the-counter medication/HDS use. This process can provide crucial information to establish a temporal relationship between drug exposure and development of signs/symptoms of liver disease. The spectrum of clinical presentation with DILI is broad, ranging from asymptomatic elevation in liver enzymes to nonspecific symptoms characterized by malaise, abdominal pain, and nausea to jaundice, pruritus, and encephalopathy. Establishing a timeline of onset of symptoms with respect to exposure of a culprit drug can be helpful, as the pattern of liver injury can change over the course of evolution of DILI. Figure 2 shows a stepwise approach to diagnosing DILI.

CAUSALITY ASSESSMENT

A physician’s awareness of a drug’s hepatotoxic potential and associated phenotypic pattern is useful when considering a diagnosis of DILI. LiverTox (https://www.ncbi.nlm.nih.gov/books/NBK547852/) provides a comprehensive characterization of drugs and HDSs, describing their typical patterns and presentations.

A number of DILI-specific causality assessment methods have been developed. These include general scales, algorithms, and expert opinions. The assessment provided by an algorithm or scale primarily depends on the weight given to each criterion. The validity of the method can vary as a result of differences in prioritizing the parameters. Table 2 details three liver-specific causality assessment scales. Causality scales are confounded by the absence of a true “gold standard” test for DILI. There is also varied interobserver reliability and reproducibility. Hence, these scales do not substitute a physician’s clinical judgement.

The DILIN uses a structured expert opinion process for categorization of probability of DILI. They describe the likelihood of DILI based on percent probability of diagnosis of DILI: definite (>95%), highly likely (75%-95%), probable (50%-74%), possible (25%-49%), or unlikely (<25%). An assessment for the first 300 patients enrolled to U.S. DILIN in 2003 determined the RUCAM approach was more conservative in assigning a high level of causality than the DILIN strategy. Although the two methods have been compared, the DILIN method has not been externally validated.

LIVER BIOPSY

Typically, a liver biopsy is not necessary to establish a diagnosis of DILI; however, it can prove useful in excluding other etiologies for liver injury and to assess the degree of inflammation and necrosis. The lack of resolution of liver injury serves as a strong rationale to obtain a liver biopsy.

Phenotypes

CLINICOPATHOLOGIC PHENOTYPES

Kleiner described 18 histopathologic patterns on liver biopsies of patients with suspected DILI. Most of the cases could be classified into one of five patterns: acute hepatitis, chronic hepatitis, acute cholestasis, chronic cholestasis, and cholestatic hepatitis.

Necroinflammatory

Inflammation, necrosis, and apoptosis are typical findings in hepatocellular DILI. The pattern of necrosis
can be zonal versus nonzonal, and acute versus chronic hepatitis-like.\(^{65}\) Acute hepatitis-like injury affects the hepatic parenchyma predominantly. Severe acute hepatitis is characterized by lobular disarray, from extensive sinusoidal architectural disruption, which correlates with a higher degree of hepatocyte apoptosis and confluent necrosis.\(^{64}\) The main differential for acute hepatitis-like DILI includes acute viral hepatitis and fulminant presentation of autoimmune hepatitis (AIH). Confluent necrosis with acute hepatitis has been seen in some cases of diclofenac related DILI.\(^{66}\) Chronic hepatitis-like DILI has a predominant portal infiltrate with interface hepatitis. Pure zonal necrosis, typically zone 3 necrosis, can mimic hypoxic-ischemic liver injury, typical of APAP hepatotoxicity. This usually signifies cytotoxicity secondary to toxic drug metabolites, as opposed to necrosis with immune cell infiltration suggestive of involvement of the adaptive immune system.\(^{65}\)

**Cholestatic**

Acute cholestasis from DILI can be categorized as bland cholestasis or cholestatic hepatitis. Bland cholestasis is classically associated with injury caused by anabolic steroids and estrogens.\(^ {67}\) Both forms of acute cholestasis share common features of zone 3 hepatocellular and canaliculic cholestasis. Concurrent granulomatous or eosinophilic inflammation in cholestatic hepatitis is suggestive of an immuno-allergic or hypersensitivity-type immune reaction.\(^ {68}\) Penicillins, especially with beta-lactamase inhibitors (amoxicillin/clavulanate), sulfonyleureas, methimazole and cephalosporins, have been implicated in causing cholestatic hepatitis.\(^ {69-72}\) Azathioprine and mercaptopurine can cause liver injury, resembling both forms of acute cholestasis.\(^ {73}\) Vanishing bile duct syndrome (VBDS) is an ominous finding in cholestatic DILI, commonly identified by a paucity of bile ducts in over 50% of

---

**FIG. 2.** Approach to diagnosis of DILI. Abbreviations: ANA, antinuclear antibody; ASMA, anti–smooth muscle actin; EtOH, ethanol; HBsAg, hepatitis B surface antigen; HCV, hepatitis C virus; HEV, hepatitis E virus; HSV, herpes simplex virus; IgG, immunoglobulin G; IgM, immunoglobulin M; LDH, lactate dehydrogenase.
A DILIN study confirmed the increased likelihood of development of chronic DILI in 94% of patients with VBDS in comparison to 47% in those without. (74) Amoxicillin/clavulanate, azithromycin, and fluoroquinolones were major causes of VBDS in this study cohort.

Steatohepatitis

Historically, the risk of drug-associated fatty liver disease related to tamoxifen or methotrexate use increased with the presence of obesity or diabetes. (33) Steatohepatitis has been associated with amiodarone, methotrexate, and tamoxifen. Microvesicular steatosis may indicate mitochondrial injury, which has a higher risk of severe liver injury, seen in DILI secondary to aspirin, valproate, amiodarone, and fialuridine. (75-80)

Vascular Injury

DILI from a vascular injury can present acutely with abdominal pain, hepatomegaly, abnormal liver enzymes, and acute onset portal hypertension. This usually occurs in the setting of Budd-Chiari syndrome and veno-occlusive disease/sinusoidal obstruction syndrome (VOD/SOS). VOD/SOS affects small veins and sinusoids, and typically presents as zone 3 necrosis. VOD/SOS is most commonly observed in the context of stem cell transplantation, exposure to toxins, pyrrolizidine alkaloids, onco-therapeutic agents, and purine analogues. Thrombosis of the major hepatic veins in Budd-Chiari can lead to massive congestion and ischemic necrosis. This is now an uncommon scenario, as it was associated with early formulations of contraceptive steroids. (65,81-83) Nodular regenerative hyperplasia, typically identified with a reticulin stain, has been seen in cases of exposure to azathioprine (as a result of active metabolite 6-thioguanine), oxaliplatin, and chlorambucil, in addition to various non–drug-related disease states. (33,65,81)

Immune Phenotypes

Drug-Induced AIH

In multiple case cohorts, 2%-9% of cases of AIH were considered to be induced by drugs, and drug-induced AIH is responsible for 9% of all cases of DILI. (33,84,85) Differentiation between drug-induced AIH and idiopathic AIH can be difficult for both the gastroenterologist and the pathologist, as patients often present with serologic and histologic markers of idiopathic AIH. As discussed previously, identification of DILI risk alleles can aide in the diagnosis of
drug-related AIH, such as HLA DRB1*15:01, which is positive in 57%-67% of DILI cases from amoxicillin/clavulanate, in comparison with 15%-20% cases in the general population. Other medications associated with drug-induced AIH include minocycline, nitrofurantoin, hydralazine, infliximab, and methyldopa.  

### Immune Therapy-Related Liver Injury

Immune checkpoint inhibitors (ICIs) increase T-cell responses and restore antitumor immune responses, which have been suppressed by cancer with the goal of inducing tumor rejection. The various targets for ICIs include cytotoxic T-lymphocyte antigen 4 (CTLA-4, the target for ipilimumab), programmed cell death 1 (PD-1, the target for pembrolizumab and nivolumab), and programmed cell death ligand 1 (the target for avelumab, durvalumab, and atezolizumab).  

However, the therapeutic reversal of immune tolerance following administration of these agents comes at the expense of immune-related adverse events (irAEs), including hepatotoxicity. A recent meta-analysis showed a higher rate of all-grade and high-grade hepatotoxicity with CTLA-4 inhibitors in comparison to PD-1 inhibitors. In other clinical trials with ipilimumab treatment, 11% of patients had early discontinuation of treatment due to hepatotoxicity, and this rose to 30% early discontinuation in combination therapy with ipilimumab and nivolumab.  

The mechanisms of the specific irAEs, which include rash, diarrhea, colitis, hepatitis and endocrinopathies, are not fully understood. It is postulated that hepatitis occurs from immune T-cell activation, leading to secretion of CD4 T-helper cell cytokines and cytolytic CD8 T-cell tissue infiltration.  

Hepatotoxicity from ICIs varies from an asymptomatic elevation of aminotransferases to acute hepatitis and fulminant liver failure. The pattern of liver injury can be nonspecific as well, ranging from hepatocellular to cholestatic. ICI-related hepatitis is typically a seronegative hepatitis in comparison to cases of idiopathic AIH. The 2018 American Society of Clinical Oncology Clinical Practice Guideline details the grading system for ICI-related hepatotoxicity and its management. Table 3 details the grades of toxicity from ICIs and proposed management.

### Treatment

The most important step in management of suspected DILI is the discontinuation of the implicated agent and avoiding re-exposure. In most cases (up to 90% or more), spontaneous recovery occurs as a result, without need for further treatment measures. Such improvement with discontinuation of a suspected offending agent is termed “dechallenge” and serves as good evidence of a causal association with injury.  

Short-term administration of a bile acid resin can be used in DILI by enhancing drug clearance by interrupting enterohepatic circulation of leflunomide and terbinafine. Carnitine is an antidote to valproate hepatotoxicity. It regulates mitochondrial acetyl-CoA levels, leading to enhanced fatty acid uptake and beta oxidation in the mitochondria.  

The role of N-acetylcysteine (NAC) in APAP-related liver injury has been well established. NAC should be considered in patients with ALF from IDILI. The U.S. ALF study group randomized 45 patients with ALF from non-APAP DILI to receive NAC versus placebo infusion. The NAC group had 2-times higher transplant-free survival rates. The use of corticosteroids should be limited to DILI in the setting of drug-induced AIH, immune therapy–related severe hepatitis, or in the presence of features of hypersensitivity. Corticosteroid treatment in one retrospective analysis was associated

### Table 3. ICI Hepatotoxicity Grading and Management

| Parameter                  | Grade 1 | Grade 2 | Grade 3 | Grade 4 |
|----------------------------|---------|---------|---------|---------|
| ALT (×ULN)                 | 1-3     | 4-5     | 6-20    | >20     |
| Bilirubin (×ULN)           | 1-1.5   | 1.6-3   | 4-10    | >10     |
| Liver disease/decompensation | Continue | Hold | Stop | Stop |
| ICI treatment              |         |         |         |         |
| Lab monitoring             | Twice weekly | Twice weekly | Daily | Daily |
| DILI treatment             | Prednisone | IV Solumedrol | IV Solumedrol | IV Solumedrol |

Abbreviation: IV, intravenous.
with lower survival in patients with more severe liver injury.\textsuperscript{(95)} When corticosteroids are used, close monitoring is essential, as a failure of response may either suggest no benefit or an alternative diagnosis, and they should be stopped. With improvement of liver injury, withdrawal of steroids should be performed gradually with close follow-up to detect resurgence of liver injury or uncover an AIH that warrants ongoing therapy.

Ursodiol can be used as pretreatment prophylaxis for patients undergoing myeloablative hematopoietic stem cell transplantation with good tolerability. Defibrotide is used as both a treatment for severe SOS and pretreatment prophylaxis in patients at high risk of developing SOS.\textsuperscript{(96)} Liver transplantation remains the primary rescue treatment for ALF from DILI. Timely identification of ALF should prompt a referral to a liver transplant center. Table 4 details the targeted therapies for DILI.

**HDS-Induced Liver injury**

More than half of U.S. households use some form of dietary supplement, most commonly multivitamins and minerals.\textsuperscript{(97-100)} The incidence of hepatotoxicity from HDSs is steadily increasing in the United States, up to 20\% in 2013-2014.\textsuperscript{(15,16)} The current regulatory framework established by the Dietary Supplement Health and Education Act of 1994 is suboptimal. The requirements for HDS products before being marketed are significantly less stringent than for pharmaceutical products; specifically, there is no requirement for assessments of product efficacy and safety. The FDA and other regulatory bodies take action only after a supplement has been shown to have potential toxicity.\textsuperscript{(101)}

Among HDSs, bodybuilding products are the most common cause of liver injury.\textsuperscript{(16)} The typical clinical scenario is that of prolonged jaundice in young men, with nonfatal outcomes from use of anabolic androgenic steroids.\textsuperscript{(102,103)} Non-bodybuilding HDS-related liver injury is commonly seen in women with a hepatocellular pattern of injury. This group of patients with HDS-related liver injury tend to have worse outcomes, with transplantation rates up to 11\%.\textsuperscript{(16)} Hillman found that in comparison to prescription medications, HDSs tend to have a higher transplantation rate and lower rate of ALF-specific transplant-free survival.\textsuperscript{(104)} Non-bodybuilding HDSs can be either single-ingredient or multi-ingredient products. One of the most common single-ingredient HDSs is green tea extract (GTE). Results from a randomized Minnesota study showed that the health of postmenopausal women without chronic liver disease were 7 times more likely to have ALT elevations with GTE use. GTE dechallenge leads to a downtrend in ALT levels, and ALT elevation recurrent following GTE rechallenge,\textsuperscript{(105)} indicating a causal relationship between GTE and hepatotoxicity.

Multi-ingredient products also pose a risk for HDS-related liver injury.\textsuperscript{(16,106)} Given their unclear chemical descriptors, identification of the culprit agent is difficult. In addition, there can be contamination of HDSs with microbials, heavy metals, and mycotoxins. Large batch-to-batch variation in HDS product content is common.\textsuperscript{(107-110)} The U.S. DILIN using toxicologic analysis confirmed the suspicion of supplement mislabeling in multi-ingredient products, with rates up to 50\%-80\%.\textsuperscript{(111)} The potential for more severe liver injury with these products remains largely unknown. However, the risk for toxicity may result from the complex interactions among individual ingredients, supplement overuse/misuse, or combination use with prescription medications.

**Summary**

DILI remains the leading cause for ALF in the U.S. adult population. It is a diagnosis of exclusion and requires a meticulous process of evaluation and a high index of clinical suspicion to attribute a potential agent as the cause of the liver injury. The accurate identification of clinical and biochemical patterns of liver injury at the onset enables prognostication for individual cases. HDSs are also a common cause for DILI, although the exact ingredient responsible for injury is difficult to confirm. The DILINs have played a vital role in expanding our understanding of liver injury from both drugs and HDSs. With continued

| Agent                        | Treatment                  
|------------------------------|----------------------------|
| APAP                         | NAC                        |
| Valproic acid                | Carnitine                  |
| Leflunomide                  | Bile salt resin            |
| Amanita (mushroom)           | Silimarin                  |
| ICI s                        | Steroids                   |
| Sinusoidal obstruction syndrome | Ursodiol, defibrotide     |
advancements and expansion of drug injury registries, our hope is that the understanding of DILI epidemiology, mechanism of injury, and establishment of causality will continue to improve.

REFERENCES

1) Ostapowicz G, Fontana RJ, Schiedt FV, Larson A, Davern TJ, Han SH, et al. Results of a prospective study of acute liver failure at 17 tertiary care centers in the United States. Ann Intern Med 2002;137:947-954.

2) Kaplowitz N. Drug-induced liver disorders: implications for drug development and regulation. Drug Saf 2001;24:483-490.

3) Aithal GP, Watkins PB, Andrade RJ, Larrey D, Molokhia M, Takikawa H, et al. Case definition and phenotype standardization in drug-induced liver injury. Clin Pharmacol Ther 2011;89:806-815.

4) Danan G, Teschke R. RUCAM in drug and herb induced liver injury: the update. Int J Mol Sci 2016;17:14.

5) Bunchornwutkul C, Reddy KR. Acetaminophen-related hepatotoxicity. Clin Liver Dis 2013;17:587-607.

6) Sgro C, Clinard F, Ouazir K, Chanay H, Allard C, Guilleminet C, et al. Incidence of drug-induced hepatic injuries: a French population-based study. Hepatology 2002;36:451-455.

7) Devalle MB, Av Klinteberg V, Alem N, Olsson R, Björnsson E. Drug-induced liver injury in a Swedish University hospital out-patient hepatology clinic. Aliment Pharmacol Ther 2006;24:1187-1195.

8) de Abajo FJ, Montero D, Madurga M, Rodriguez LAG. Acute and clinically relevant drug-induced liver injury: a population based case-control study. Br J Clin Pharmacol 2004;58:71-80.

9) Bjornsson ES, Bergmann OM, Björnsson HK, Kvaran RB, Olafsson S. Incidence, presentation, and outcomes in patients with drug-induced liver injury in the general population of Iceland. Gastroenterology 2013;144:1419-1425.e3.

10) Wai CT, Tan BH, Chan CL, Sutedja DS, Lee YM, Khor C, et al. Drug-induced liver injury at an Asian center: a prospective study. Liver Int 2007;27:465-474.

11) Shen T, Liu Y, Shang J, Xie Q, Li J, Yan M, et al. Incidence and etiology of drug-induced liver injury in mainland China. Gastroenterology 2019;156:1337-1345.e2.

12) Vega M, Verna M, Beswick D, Bey S, Hossack J, Merriman N, et al. The incidence of drug-and herbal and dietary supplement-induced liver injury: preliminary findings from gastroenterologist-based surveillance in the population of the state of Delaware. Drug Saf 2017;40:783-787.

13) Fontana RJ, Watkins PB, Bonkovsky HL, Chalasani N, Davern T, Serrano J, et al.; DILIN Study Group. Drug-induced liver injury network (DILIN) prospective study: rational, design and conduct. Drug Saf 2009;32:55-68.

14) Chalasani N, Bonkovsky HL, Fontana R, Lee W, Stolz A, Tawalkar J, et al.; United States Drug Induced Liver Injury Network. Features and outcomes of 899 patients with drug-induced liver injury: the DILIN prospective study. Gastroenterology 2015;148:1340-1352.

15) Navarro VJ, Khan I, Bjornsson E, Seeff LB, Serrano J, Hoofnagle JH. Liver injury from herbal and dietary supplements. Hepatology 2017;65:363-373.

16) Navarro VJ, Barnhart H, Bonkovsky HL, Davern T, Fontana RJ, Grant L, et al. Liver injury from herbs and dietary supplements in the U.S. drug induced liver injury network. Hepatology 2014;60:1399-1408.

17) Danan G, Benichou C. Causality assessment of adverse reactions to drugs. I: A novel method based on the conclusions of international consensus meeting. J Clin Epidemiol 1993;46:1323-1330.

18) Benichou C, Danan G, Flahault A. Causality assessment of adverse reactions to drugs. II: An original model for validation of drug causality assessment methods: case reports with positive rechallenge. J Clin Epidemiol 1993;46:1331-1336.

19) Saithanyamurthi H, Faust AJ. Drug-induced liver disease: clinical course. Clin Liver Dis 2017;21:21-34.

20) Bessone P, Hernandez N, Lucena M, Andrade R. The Latin American DILI registry experience: a successful ongoing collaborative strategic initiative. Int J Mol Sci 2016;17:313.

21) Björnsson E, Jerstad P, Bergqvist A, Olsson R. Fulminant drug-induced hepatic failure leading to death or liver transplantation in Sweden. Scand J Gastroenterol 2005;40:1095-1101.

22) Zimmerman HJ. Drug-induced liver disease. In: Zimmerman HJ, ed. Hepatoxicity: The Adverse Effects of Drugs and Other Chemicals on the Liver. 2nd Edition. Philadelphia, PA: Lippincott Williams & Wilkins; 1999:427-456.

23) Zimmerman HJ. The spectrum of hepatotoxicity. Perspect Biol Med 1968;12:135-161.

24) Black M, Mitchell JR, Zimmerman HJ, Ishak KG, Epler GR. Isoniazid-associated hepatitis in 114 patients. Gastroenterology 1975;69:289-302.

25) Rohles-Diaz M, Lucena MI, Kaplowitz N, Stephans C, Medina-Cálix I, González-Jimenez A, et al. Use of Hy’s law and a new composite algorithm to predict acute liver failure in patients with drug-induced liver injury. Gastroenterology 2014;147:109-118.

26) Björnsson ES, Björnsson HK. Mortality associated with drug-induced liver injury (DILI). Transl Gastroenterol Hepatol 2017;2:114.

27) Teschke R. Idiosyncratic DILI: analysis of 46,266 cases assessed for causality by RUCAM and published from 2014 to early 2019. Front Pharmacol 2019;23:730.

28) Andrade RJ, Lucena MI, Kaplowitz N, García-Munoz B, Borraz Y, Pachkoria K, et al. Outcome of acute idiosyncratic drug-induced liver injury: long-term follow up in a hepatotoxicity registry. Hepatology 2006;44:1581-1588.

29) Fontana RJ, Hayashi PH, Gu J, Reddy KR, Barnhart H, Watkins PB, et al. Idiosyncratic drug-induced liver injury is associated with substantial morbidity and mortality within 6 months from onset. Gastroenterology 2014;147:96-108.

30) Habib S, Shaikh OS. Liver transplantation for drug-induced acute liver failure. Eur Rev Med Pharmacol Sci 2017;21(Suppl. 1): 37-45.

31) Uetrecht J. Idiosyncratic drug reactions: current understanding. Annu Rev Pharmacol Toxicol 2007;47:513-539.

32) Lammert C, Einarsson S, Saha C, Niklasson A, Bjornsson E. Robles-Diaz M, Lucena MI, Kaplowitz N, García-Munoz B, Borraz Y, Pachkoria K, et al. Outcome of acute idiosyncratic drug-induced liver injury: long-term follow up in a hepatotoxicity registry. Hepatology 2006;44:1581-1588.

33) European Association for the Study of the Liver. EASL clinical practice guidelines: drug-induced liver injury. J Hepatol 2019;70:1222-1261.

34) European Association for the Study of the Liver. EASL clinical practice guidelines: drug-induced liver injury. J Hepatol 2019;70:1222-1261.

35) David S, Hamilton JP. Drug-induced liver injury. US Gastroenterol Hepatol Rev 2010;1:73-80.

36) Uetrecht J. Idiosyncratic drug reactions: past, present, and future. Chem Res Toxicol 2008;21:84-92.

37) Njoku DB, Greenberg RS, Bourdi M, Borkowf CB, Dake EM, Martin JL, et al. Autoantibodies associated with volatile anesthetic hepatitis found in the sera of a large cohort of pediatric anesthesiologists. Anesth Analg 2002;94:243-249.
37) Kullak-Ublick GA, Andrade RJ, Merz M, End P, Benesic A, Gerbes AL, et al. Drug-induced liver injury: recent advances in diagnosis and risk assessment. Gut 2017;66:1154-1164.

38) Knolle PA, Gerken G. Local control of the immune response in the liver. Immunol Rev 2000;174:21-34.

39) Cantor HM, Dumont AE. Hepatic suppression of sensitization to antigen absorbed into the portal system. Nature 1967;215:744-745.

40) Dara L, Liu Z-X, Kaplowitz N. Mechanisms of adaptation and progression in idiosyncratic drug induced liver injury, clinical implications. Liver Int 2016;36:158-165.

41) Vuppalanchi R, Chalasani N. Risk factors for drug-induced liver disease. In: Kaplowitz N, DeLeve OLD, eds. Drug-Induced Liver Disease, 3rd Edition. Waltham, MA: Academic Press; 2013: 265-274.

42) Guy J, Peters MG. Liver disease in women: the influence of gender on epidemiology, natural history, and patient outcomes. Gastroenterol Hepatol 2013;9:633-639.

43) deLemos A, Fouereau D, Jacobs C, Ahrens W, Russo M, Bonkovsky HL, Watkins PB, Davern T, Serrano J, et al. Causes, clinical features, and outcomes from a prospective study of drug-induced liver injury in the United States. Gastroenterology 2008;135:1924-1934.

44) Andersd R, Chalasani N, Fonatana RJ, Bonkovsky HL, Watkins PB, Davern T, Serrano J, et al. Causes, clinical features, and outcomes from a prospective study of drug-induced liver injury in the United States. Gastroenterology 2005;129:654-664.

45) Chalasani N, Fontana RJ, Bonkovsky HL, Watkins PB, Davern T, Serrano J, et al. Causes, clinical features, and outcomes from a prospective study of drug-induced liver injury in the United States. Gastroenterology 2005;129:512-521.

46) Chalasani N, Reddy RKK, Fonatana RJ, Barnhart H, Gu J, Hayashi PH, et al. Idiosyncratic drug induced liver injury in African-Americans is associated with greater morbidity and mortality compared to Caucasians. Am J Gastroenterol 2017;112:1382-1388.

47) Lu NA, Rai SK, Terkeltaub R, Kim SC, Menendez ME, Choi HK. Reddy RKK, Fonatana RJ, Barnhart H, Gu J, Hayashi PH, et al. Idiosyncratic drug induced liver injury in African-Americans is associated with greater morbidity and mortality compared to Caucasians. Am J Gastroenterol 2017;112:1382-1388.

48) Lu NA, Rai SK, Terkeltaub R, Kim SC, Menendez ME, Choi HK. Reddy RKK, Fonatana RJ, Barnhart H, Gu J, Hayashi PH, et al. Idiosyncratic drug induced liver injury in African-Americans is associated with greater morbidity and mortality compared to Caucasians. Am J Gastroenterol 2017;112:1382-1388.

49) Kunelis CT, Peters JL, Edmondson HA. Fatty liver of pregnancy. N Engl J Med 1965;273:603-609.

50) Zimmerman HJ. Effects of alcohol on other hepatotoxins. Alcohol Clin Exp Res 1986;10:3-15.

51) Athyros VG, Tzimos M, Gossios TG, Griva T, Anagnostis P, Kargiotsi K, et al. Safety and efficacy of long-term statin treatment for cardiovascular events in patients with coronary heart disease and abnormal liver tests in the Greek Aterovastatin and Coronary Heart Disease Evaluation (GREACE) Study: a post-hoc analysis. Lancet 2010;376:1916-1922.

52) Han KH, Rha SW, Kang HJ, Bae JW, Choi BJ, Choi SY, et al. Evaluation of short-term safety and efficacy of HMG-CoA reductase inhibitors in hypercholesterolemic patients with elevated serum alanine transaminase concentrations: PITCH study (PITavastatin versus atorvastatin to evaluate the effect on patients with hypercholesterolemia and mild to moderate hepatic damage). J Clin Lipidol 2012;6:340-351.

53) Gomez-Dominguez E, Gisbert JP, Moreno-Montenegro JA, Garcia-Buey L, Moreno-Otero R. A pilot study of atorvastatin treatment in dyslipidemia, non-alcoholic fatty liver patients. Aliment Pharmacol Ther 2006;23:1643-1647.

54) Antonopoulou S, Mikros S, Mylonopoulos M, Kokkori S, Giannoulis G. Rosuvastatin as a novel treatment of non-alcoholic fatty liver disease in hyperlipidemic patients. Atherosclerosis 2006;184:233-234.

55) Daly AK, Donaldson PT, Bhattachar P, Shen Y, Perez I, Floratos A, et al. HLA-B 5701 genotype is a major determinant of drug-induced liver injury due to fluvoxamine. Nat Genet 2009;41:816-819.

56) Lucena MI, Molokhia M, Shen Y, Urban TJ, Atital GP, Andrade RJ, et al. Susceptibility to amoxicillin-clavulanate induced liver injury is influenced by multiple HLA class I and II alleles. YGAST 2011:141:338-347.

57) Urban TJ, Nicoletti P, Chalasani N, Serrano J, Stolz A, Daly AK, et al. Minocycline hepatotoxicity: clinical characterization and identification of HLA-B*35:02 as a risk factor. J Hepatol 2017;67:137-144.

58) Lucena MI, Andrade RJ, Kaplowitz N, Garcia-Cortes M, Fernández MC, Romero-Gomez M, et al. Phenotypic characterization of idiosyncratic drug-induced liver injury: the influence of age and sex. Hepatology 2009;49:2001-2009.

59) Ghabril M, Chalasani N, Björnsson E. Drug-induced liver injury: a clinical update. Curr Opin Gastroenterol 2010;26:222-226.

60) Shen M, Borlak J, Tong W. High lipophilicity and high daily dose of oral medications are associated with significant risk for drug-induced liver injury. Hepatology 2013;58:388-396.

61) Maria VA, Victorino RM. Development and validation of a clinical scale for the diagnosis of drug-induced hepatitis. Hepatology 1997;26:664-669.

62) Takikawa H, Takamori Y, Kumagi T, Onji M, Watanabe M, Shibuya A, et al. Assessment of 287 Japanese cases of drug-induced liver injury by the diagnostic scale of the International Consensus Meeting. Hepatol Res 2003;27:192-195.

63) Rockey DC, Seef LB, Rochon J, Freston J, Chalasani N, Bonacini M, et al. Causality assessment in drug-induced liver injury using a structured expert opinion process: comparison to the Roussel-Uclaf causality assessment method. Hepatology 2010;51:2117-2126.

64) Kleiner DE, Chalasani NR, Lee WM, Fonatana RJ, Bonkovsky HL, Watkins PB, et al.; Drug-Induced Liver Injury Network. Hepatic histological findings in suspected drug-induced liver injury: systematic evaluation and clinical associations. Hepatology 2014;59:661-670.

65) Kleiner DE. Histopathological challenges in suspected drug-induced liver injury. Liver Int 2018;38:198-209.

66) Schmelzer PA, Kosinski AS, Kleiner DE, Hoofnagle JH, Stolz A, Fonatana RJ, et al. Liver injury from non-steroidal anti-inflammatory drugs in the United States. Liver Int 2016;36:603-609.

67) Bond P, Llewellyn W, Van Mol P. Anabolic androgenic steroid-induced hepatotoxicity. Med Hypotheses 2016;93:150-153.

68) Björnsson E, Kalaitzakis E, Olsson R. The impact of eosinophilia and its relationship to tetracycline therapy. Am J Med 1965;36:359-377.

69) Hautekeete ML. Hepatotoxicity of antibiotics. Acta Gastroenterol Belg 1995;58:290-296.

70) Chounta A, Zouridakis S, Ellinas C, Tsiodras S, Zoumpouli C, et al. Idiosyncratic drug-induced liver injury: clinical features and outcomes from a post-hoc analysis. Lancet 2010;376:1916-1922.

71) Han KH, Rha SW, Kang HJ, Bae JW, Choi BJ, Choi SY, et al. Evaluation of short-term safety and efficacy of HMG-CoA reductase inhibitors in hypercholesterolemic patients with elevated serum alanine transaminase concentrations: PITCH study (PITavastatin versus atorvastatin to evaluate the effect on patients with hypercholesterolemia and mild to moderate hepatic damage). J Clin Lipidol 2012;6:340-351.

72) Gomez-Dominguez E, Gisbert JP, Moreno-Montenegro JA, Garcia-Buey L, Moreno-Otero R. A pilot study of atorvastatin treatment in dyslipidemia, non-alcoholic fatty liver patients. Aliment Pharmacol Ther 2006;23:1643-1647.
74) Bondkovsky HL, Kleiner DE, Gu J, Odin JA, Russo MW, Navarro VM, et al. Clinical presentations and outcomes of bile duct loss caused by drugs and herbal and dietary supplements. Hepatology 2017;65:1267-1277.
75) Bregiche K, Massart J, Robin M-A, Borgne-Sanchez A, Fromenty B. Drug-induced toxicity on mitochondria and lipid metabolism: mechanistic diversity and deleterious consequences for the liver. J Hepatol 2011;54:773-794.
76) Trosi LC, Lemasters JJ. The mitochondrial permeability transition: a new pathophysiological mechanism for Roye's syndrome and toxic liver injury. J Pharmacol Exp Ther 1996;278:1000-1005.
77) Silva MF, Ruiter JF, Jhst L, Jakobs C, Duran M, de Almeida JT, et al. Differential effect of valproate and its Delta2- and Delta4-unsaturated metabolites, on the beta-oxidation rate of long-chain and medium-chain fatty acids. Chem Biol Interact 2001;137:203-212.
78) Card JW, Lalonde BR, Rafeiro E, Tam AS, Racz WJ, Brien JF, et al. Amiodarone-induced disruption of hamster liver and mitochondrial function: lack of association with thiobarbituric acid-reactive substance production. Toxicol Lett 1998;98:41-50.
79) Fromenty B, Fisch C, Berson A, Letteron P, Larrey D, Pessayre D. Dual effect of amiodarone on mitochondrial respiration. Initial protonophoric uncoupling effect followed by inhibition of the respiratory chain at the levels of complex I and complex II. J Pharmacol Exp Ther 1990;255:1377-1384.
80) Lewis W, Levine ES, Grinuvivi B, Tankersley KO, Colacino JM, Sommadossi JP, et al. Fluorouracil and its metabolites inhibit DNA polymerase gamma at sites of multiple adjacent analog incorporation, decrease mtDNA abundance, and cause mitochondrial structural defects in cultured hepatoblasts. Proc Natl Acad Sci U S A 1996;93:3592-3597.
81) Kleiner DE. Drug-induced liver injury: the hepatic pathologist's approach. Gastroenterol Clin North Am 2017;46:273-296.
82) Lewis JH, Tice HL, Zimmerman HJ. Chiari syndrome associated with oral contraceptive steroids. Review of treatment of 47 cases. Dig Dis Sci 1983;28:673-683.
83) Ulrickson M, Aldridge J, Kim HT, Hochberg EP, Svarcz AJ, et al. Carnitine in the treatment of valproic acid-induced toxicity. Clin Toxicol 2009;47:101-111.
84) Lee WM, Hyman LS, Rossaro L, Fontana RJ, Stravitz RT, Larson AM, et al. Intravenous N-acetylcysteine improves transplant-free survival in early stage nonacacetaminophen acute liver failure. Gastroenterology 2009;137:856-864.
85) Kantor ED, Rehm CD, Du M, White E, Giovannucci EL. Trends in dietary supplement use among US adults from 1999-2012. JAMA 2016;316:1464-1474.
86) Radimer K, Birnbaum Q, Hughes J, Ervin B, Swanson C, Picklino MF, Dietary supplement use by US adults: data from the National Health and Nutrition Examination Survey, 1999-2000. Am J Epidemiol 2004;160:339-349.
87) Food and Drug Administration. Dietary supplements. Updated August 16, 2019. http://www.fda.gov/Food/DietarySupplements/default.htm. Accessed January 2, 2020.
88) Krishnan PV, Feng ZZ, Gordon SC. Prolonged intrahepatic cholestasis and renal failure secondary to anabolic androgenic steroid-enriched dietary supplements. J Clin Gastroenterol 2009;43:672-675.
89) Kafrouni MI, Perkins KA, Verma S. Hepatotoxicity associated with dietary supplements containing anabolic steroids. Clin Gastroenterol Hepatol 2007;5:809-812.
90) Hillman L, Gottfried M, Whitsett M, Rakela J, Slivsky M, Lee WM, et al. Clinical features and outcomes of complementary and alternative medicine induced acute liver failure and injury. Am J Gastroenterol 2016;111:958-965.
91) Yu Z, Samavat H, Dostal AM, Wang R, Torkelson CJ, Yang CS, et al. Effect of green tea supplements on liver enzyme elevation: results from a randomized intervention study in the United States. Cancer Prev Res 2017;10:571-579.
92) Medina-Caliz I, Garcia-Cortes M, Gonzalez-Jimenez A, Caballo MR, Robles-Diaz M, Sanabria-Cabera J, et al. Herbal and dietary supplement-induced liver injuries in the Spanish DILLI registry. Clin Gastroenterol Hepatol 2018;16:1495-1502.
93) Liu J, Pei M, Zheng C, Li Y, Wang Y, Lu A, et al. A systems-pharmacology analysis of herbal medicines used in health improvement treatment: predicting potential new drugs
108) Stickel F, Droz S, Patsenker E, Bögli-Stuber K, Aebi B, Leib SL. Severe hepatotoxicity following ingestion of Herbalife contaminated with Bacillus subtilis. J Hepatol 2009;50:111-117.

109) Miller GM, Stripp R. A study of western pharmaceuticals contained within samples of Chinese herbal/patent medicines collected from New York City’s Chinatown. Leg Med (Tokyo) 2007; 9:258-264.

110) Gurley BJ, Gardner SF, Hubbard MA. Content versus label claims in ephedra-containing dietary supplements. Am J Health Syst Pharm 2000;57:963-969.

111) Navarro V The frequency of herbal and dietary supplement mislabeling: experience of the drug induced liver injury network. AASLD LiverLearning 2017;24:201449.

Author names in bold designate shared co-first authorship.