Liver Fibrosis Helps to Distinguish Autoimmune Hepatitis from DILI with Autoimmune Features: A Review of Twenty Cases

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Abstract

Background and Aims: Drug-induced liver injury with autoimmune features (AI-DILI) mimics the clinical presentation, and laboratory and pathologic features of idiopathic autoimmune hepatitis (AIH). We aimed to identify histopathologic hallmarks to differentiate these entities. Methods: All liver biopsies archived for the past 10 years were reviewed retrospectively to identify cases of recently detected liver injury associated with predominantly lymphoplasmacytic interphase hepatitis, positive markers for liver autoimmunity, and negative tests for viral hepatitis. Twenty cases were divided into AIH (n = 12) or AI-DILI (n = 8) groups. Blind qualitative evaluation of necroinflammatory changes and liver fibrosis were performed according to the Scheuer scoring system. Cellular densities were determined using ImageJ (V.1.51t, National Institutes of Health, Bethesda, MD, USA). Fibrosis was assessed on Masson trichrome-stained slides, and collagen deposition was estimated following a protocol of color deconvolution. Results: Necroinflammatory changes as well as densities (portal and lobular) of neutrophils and eosinophils, intracellular cholestasis, and regenerative changes did not differ between the two groups (P ≥ 0.05). Neutrophil densities but not eosinophils showed a positive correlation with the severity of hepatocellular damage (r = 0.6 and 0.58, vs. alanine aminotransferase, P < 0.05). Ceroid-laden macrophages but not histiocytic aggregates appeared to be more common in AI-DILI (P < 0.05). AIH patients presented more often with evidence of chronic damage, including higher scores of fibrosis and collagen deposition, in comparison to AI-DILI (P < 0.05). Conclusions: Although there is no histologic feature pathognomonic for AI-DILI or AIH, advanced stages of liver fibrosis can be used to support the diagnosis of AIH in some cases. Definitive diagnosis of AI-DILI requires follow-up and demonstration of complete remission after drug withdrawal with no need for immunosuppression.

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Introduction

Autoimmune hepatitis consists of immune-mediated damage of hepatocytes associated with the development of autoantibodies.1 The diagnoses of idiopathic autoimmune hepatitis (AIH) and drug-induced liver injury (DILI) are challenging because both conditions show overlapping manifestations. Recently, DILI with autoimmune features (AI-DILI) has been recognized as a discrete entity.2 AI-DILI is characterized by liver injury due to the ingestion of medications or herbal products with simultaneous positivity for markers of liver autoimmunity (high IgG immunoglobulin levels, anti-nuclear antibodies, anti-smooth muscle antibodies, anti-liver-kidney microsomal antibodies, and rarely anti-mitochondrial antibodies). The liver damage becomes clinically evident within 3 months following the drug exposure, but it can appear after a longer latency interval.2,3 Among the medications causing this syndrome are statins, diclofenac, hydralazine, methyldopa, minocycline, nitrofurantoin, and procainamide.3

By definition, in AI-DILI, the liver injury must completely resolve after drug withdrawal with no recurrence. Recovery is reached either spontaneously or with immunosuppressive therapy. Steroids and azathioprine are among the medications applied to treat AI-DILI.2,3 Time to resolution varies from days to weeks, lasting months for a complete normalization of liver enzymes in a few cases. Those cases that relapse after drug withdrawal have been described as AIH triggered by drugs. Furthermore, relapse after immunosuppressive therapy withdrawal does not occur in AI-DILI, and its absence distinguishes AI-DILI from classical AIH.4,5 Timely diagnosis is critical for proper management in both conditions. Early immunosuppression can lead to remission in AIH. Likewise, prompt identification and discontinuation of toxic drugs halt liver injury in AI-DILI. Failure to properly treat AIH or AI-DILI could result in adverse clinical outcomes.3 AIH can present with typical histopathological findings, but DILI may mimic any non-DILI pattern of liver injury, including AIH. Critical characteristics of AIH include a mixed inflammatory infiltrate composed of lymphocytes and plasma cells that is most marked around portal areas and is referred to as interface hepatitis. The presence of a predominance of plasma cells within the infiltrate is highly suggestive of the diagnosis. Necrosis of periportal hepatocytes, scattered acidophil or apoptotic bodies, and cytoplasmic swelling are
manifestations of hepatocellular damage. AIH may also cause the full temporal spectrum of liver injury, from mild fibrosis to established cirrhosis.1 Diagnosing AIH also requires ruling out infectious causes of liver injury, including viral hepatitides.6 AI-DILI mimics the morphological pattern of AIH, including the prominent lymphoplasmacytic infiltrates in portal spaces and interface hepatitis.2,7 In this study, we performed a histopathological evaluation of liver biopsies to further identify potential hallmarks for differentiating both entities.

Methods

Case selection

Liver biopsies stored in the tissue archives of Mount Sinai Medical Center Department of Pathology for the past 10 years were reviewed. Cases were included if there was no history of prior liver disease and when the primary clinical and pathological differential diagnosis was AIH, according to recommendations of the American Association for the Study of Liver Diseases.4 The diagnosis of AIH was based on the presence of the autoantibodies anti-smooth muscle antibodies or anti-liver-kidney microsomal antibodies, or high IgG levels with compatible histology and exclusion of infectious etiologies. All cases had to show predominant lymphoplasmacytic infiltrates with piecemeal necrosis (Fig. 1). Sorting into the AIH or AI-DILI groups was done retrospectively according to pertinent clinical history, follow-up notes, and laboratory tests supporting liver autoimmunity and ruling out viral hepatitis. Liver biopsies included in the analysis were those performed at the time of presentation, when no diagnosis was known. For AI-DILI, the liver injury had to be associated with a drug exposure and completely resolve after drug withdrawal and with no need for immunosuppression in subsequent follow-ups. If a patient developed a relapse or persistent liver damage after recovery from a drug exposure, the case was included in the AIH group. Charts were reviewed to rule out re-challenge to the hepatotoxic drug. Relapsing cases of AIH triggered by drugs must not have been exposed again to the suspected drug in order to be considered part of this group. The Mount Sinai Medical Center Institutional Review Board approved the study in conformance to the ethical guidelines of the 1975 Declaration of Helsinki.

Histological evaluation

Blind qualitative evaluation of necroinflammatory changes and fibrosis were graded according to the Scheuer scoring system.8 Presence or absence of intracellular cholestasis, ceroid-laden macrophages, and histiocytic aggregates were also recorded (Fig. 1). Regenerative changes and hepatocyte necrosis were evaluated in reticulin-stained slides. Cellular densities were determined on captured images using ImageJ (V1.51t, National Institutes of Health, Bethesda, MD, USA). Liver fibrosis was assessed on Masson trichrome-stained slides, and collagen deposition was estimated on captured images following a protocol of color deconvolution.9,10 The amount of collagen was recorded as a fraction area of the adjusted threshold value of green color within the region of interest (Fig. 2a).
Statistical analysis

Scheuer scores and other histologic features were compared using Wilcoxon rank-sum and Fisher’s exact tests. Cellular densities and fraction areas of collagen deposition were compared using Pearson or Spearman tests depending on the type of variable. All P values presented were two-sided and considered statistically significant when less than 0.05. The statistical analysis was performed in Microsoft Excel® (V15; Microsoft Corporation, Redmond, WA, USA) and GraphPad Prism® (V6; GraphPad Software Inc, La Jolla, CA, USA).

Results

Patient features

The total number of cases was 20. Detailed patient’s characteristics including history of autoimmune diseases, evidence of liver autoimmunity at presentation, latency period, and recovery time are shown in Table 1. There were no significant differences between the AIH (n = 12) and AI-DILI (n = 8) groups regarding age (49 [24–66] vs. 54 [17–71], median [range], respectively). The male gender was relatively more frequent in AIH than in AI-DILI (33% vs. 13%, respectively). The mean±SEM of laboratory liver parameters at presentation were not significantly different between the AIH and AI-DILI groups: 938.9±281.9 versus 1339±395.7 U/L for alanine aminotransferase (ALT) levels; 1084±337.2 versus 836.7±257.7 U/L for aspartate aminotransferase levels; 187±26.0 versus 299.8±114.8 U/L for alkaline phosphatase levels; and 7.41±3.001 versus 4.58±1.54 mg/dL for total bilirubin levels (P = 0.05, unpaired t-test). Exposure to a drug known to cause AI-DILI was documented in 33% (4/12) of AIH cases and 100% (8/8) of AI-DILI. Drugs of the AIH group included clavulanic acid, levofloxacin, ramipril, and simvastatin. Doxycycline, atorvastatin, simvastatin, ciprofloxacin, lisinopril, isoniazid, OxyELITE, and ustekinumab were identified in the AI-DILI group. The mean follow-up period for both groups was 57 months (12–108, range).

Patients with AIH presented with a higher degree of liver fibrosis than those with AI-DILI

According to the Scheuer scoring system (Fig. 2b), 8 out of 20 patients presented with no liver fibrosis (score: 0), 5 out of 20 patients with enlarged fibrotic portal tracts (score: 1), 7 out of 20 patients with higher grades of liver fibrosis (scores: 2, 3, and 4). The amount of collagen deposition shows a strong positive correlation with Scheuer stages of liver fibrosis (r = 0.9201, P < 0.0001, Spearman correlation; Fig. 2b). The mean Scheuer score for liver fibrosis in AIH was 1.58±1.16 versus 0.37±0.74 in AI-DILI (mean±SD, P < 0.01, Wilcoxon rank-sum test; Fig. 2c). Advanced stages of liver fibrosis (3 and 4) were seen in only two patients with AIH. Likewise, the amount of collagen deposition was more prominent in AIH than AI-DILI (14.83±3.775 vs. 4.20±1.3565, mean±SEM, P < 0.05, unpaired t-test; Fig. 2d).

Fig. 2. Evaluation of liver fibrosis in AIH versus AI-DILI. Digital imaging protocol using the ImageJ software (5) was applied to quantify collagen deposition in liver biopsies (a). Color deconvolution of captured trichrome Masson-stained images allows separation of collagen (green component) from the background. The threshold was manually adjusted until the entire green area was highlighted in red, then converted in gray/black color. The amount of collagen was measured as fraction area-based quantification in the regions of interest (ROI). Collagen deposition determined by this method shows a strong correlation with the qualitative Scheuer scoring system for liver fibrosis (b) (mean, r = 0.9201, Spearman correlation). Patients with AIH were more likely to present with higher degrees of liver fibrosis than those with AI-DILI, as shown by collagen deposition (c, mean±SD, unpaired t-test) and Scheuer scores (d, mean±SD, Wilcoxon rank-sum test). Fibrosis in AIH is a consequence of chronicity of liver damage, a condition that is diagnosed during flares of disease, while AI-DILI more likely presents as (sub)acute onset cases with minimal or no fibrosis.
The degree of necroinflammatory changes correlates with the severity of hepatocellular damage but not with the etiology, in both groups

The severity of the inflammatory response, determined by Scheuer score (0–4), did not differ between patients with AIH versus AI-DILI in either portal spaces (2.083±0.996 vs. 2.125±0.64) or hepatic lobules (1.917±1.084 vs. 1.875 ±0.991) (mean±SD, P > 0.05, Wilcoxon rank-sum test). The cellular density of neutrophils and eosinophils in hotspots of portal spaces and hepatic lobules was also similar in both groups (mean±SD, P > 0.05, multiple t-test using the Sidak-Bonferroni method; Fig. 3a). Moreover, the cellular density of neutrophils in portal spaces and lobules, but not eosinophils, showed a positive correlation with the severity of hepatocellular damage in both groups, as measured by ALT levels (r = 0.608 vs. neutrophil density in portal spaces, P < 0.05; r = 0.586 vs. neutrophil density in hepatic lobules, P < 0.05; r = 0.209 vs. eosinophil density in portal spaces, P = 0.05; r = 0.338 vs. eosinophil density in hepatic lobules, P ≥ 0.05, Pearson correlation; Fig. 3b). The density of neutrophils in lobules but not portal spaces showed a positive correlation with necroinflammatory scores by the Scheuer method (r = 0.5766, P < 0.01, in lobules and r = 0.3236, P ≥ 0.05, in portal spaces, Spearman correlation).

The presence of ceroid-laden macrophages appears to be more common in AI-DILI and histiocytic aggregates in AIH

The presence or absence of additional histologic features of liver damage was also assessed and showed that ceroid-laden macrophages were more common in AI-DILI (87.5% vs. 33.3% in AIH, P < 0.05, Fisher’s exact test). Histiocytic
aggregates were more common in AIH (75% vs. 37.5% in AI-DILI, \( P < 0.05 \), Fisher’s exact test). The frequency of intracellular cholestasis (62.5% in AI-DILI vs. 33.3% in AIH) and regenerative changes (12.5% in AI-DILI vs. 50% in AIH) were not statistically different between groups (\( P \approx 0.05 \), Fisher’s exact test).

**Discussion**

The etiology of autoimmune liver disease is unknown, for the most part. The occurrence of liver autoantibodies can even be detected in infectious hepatitides, which suggest that they are rather nonspecific and should be used with caution when diagnosing AIH.\(^1\) There are reports of drug-induced hepatotoxicity accompanied by an autoimmune response. For that reason, AI-DILI is a differential diagnosis of AIH in daily practice.\(^2\) Histopathology of liver biopsies has been useful in diagnosing DILI and AIH. However, a significant problem for hepatologists and pathologists comes when separating AIH from AI-DILI as both entities share clinical, biochemical, and histopathologic features.\(^1\)–\(^3\) Patients with new-onset AIH frequently report recent use of medications that are also associated with DILI, such as antibiotics, statins or antihypertensive drugs.\(^1\)–\(^3\),\(^12\) We aimed in this study to identify histopathological features that can help separate AIH from AI-DILI.

Microscopic evaluation showed comparable severity in interface hepatitis, hepatocyte necrosis, portal and lobular inflammation, and infiltration of eosinophils in both conditions. In a previous study, the presence of cholestasis and portal neutrophils favored AI-DILI over AIH.\(^7\) Our results suggest that infiltration of neutrophils positively correlates with the severity of hepatocellular damage but not with the etiology. Intracellular cholestasis seemed to be more frequent in AI-DILI and regenerative changes in AIH, but they were not statistically significant. Cereoid-laden macrophages were more common in AI-DILI, and prominent histiocytic infiltrates were mainly seen in AIH; however, none of those features were skewed enough to be useful as a marker to differentiate AI-DILI from AIH in this small series. The full spectrum of noninflammatory changes that can develop in the context of liver autoimmunity was also observed in AI-DILI.

Historically, liver infiltration of eosinophils has been used to identify cases of drug hypersensitivity, allergic diseases, malignancies, hyper-eosinophilic syndrome, collagen vascular diseases, and, most commonly, parasitic infections. However, we found that eosinophils are observed in liver biopsies in patients with AIH. In terms of quantity, some forms of DILI (particularly the immunoallergic-type) are accompanied by copious amounts of eosinophils undoubtedly greater than what is seen in a typical case of AIH.\(^13\) However, AI-DILI is a distinct and rare entity among DILI reactions that appears to be mediated by immune mechanisms very similar to AIH. Thus, the density of eosinophils is not expected to be quite different between AI-DILI and AIH. We excluded other forms of DILI in this study, which may also explain that result. The inclusion criteria was very strict, and evidence of liver autoimmunity must have been documented in the chart for all cases. Liver infiltration of eosinophils has diagnostic value for some liver diseases but does not appear to help in the distinction of AI-DILI versus AIH.

Interestingly, owing to the chronic nature of AIH, persistent loss of hepatocytes leads to progressive fibrosis and eventually cirrhosis. In this way, the presence of advanced stages of liver fibrosis could be used clinically to favor the diagnosis of AIH over AI-DILI in patients with no history of a previously diagnosed chronic liver disease. This proposal was demonstrated by the presence of higher Scheuer scores of liver fibrosis and abundant collagen deposition in patients with AIH. The Scheuer scoring system has been commonly used in grading viral hepatitis-associated chronic damage and has shown good reproducibility.\(^8\) The protocol of color deconvolution applied to quantify collagen had not been validated for staging liver fibrosis; however, we found an excellent correlation between Scheuer scores and collagen deposition measurements. Fibrosis in the AIH group represents a marker of chronicity of disease. By the time a patient with AIH develops significant symptoms, it may be the expression of acute-onset AIH or a flare of previously unrecognized AIH. Therefore, significant liver fibrosis detected on liver biopsies can be used to favor a diagnosis of AIH, regardless of whether or not it is considered acute-onset AIH.\(^12\)
Early stages of liver fibrosis can be seen in AI-DILI. It is unclear whether mild fibrosis in AI-DILI is a consequence of prolonged drug toxicity or a different unrecognized liver insult. Liver fibrosis in many diseases, including AIH, is a progressive phenomenon and takes years to develop. Furthermore, drug toxicity in AI-DILI has been documented within 3 months before clinically detectable liver damage, which makes any degree of fibrosis unlikely to be secondary to drug toxicity. In a prior study, cirrhosis was only observed in AIH cases, whereas no cirrhosis was present among AI-DILI cases. Although not all AI-DILI cases are acute or chronic, AI-DILI usually causes liver damage severe enough to produce significant symptoms, leading to an early diagnosis. In many patients with AIH, the liver injury tends to be subclinical with waxing and waning episodes. AIH is detected during work-ups of long-standing unexplained increased liver enzymes or severe acute episodes of liver injury. Cases of late-onset AI-DILI, presenting at 1–2 years after drug exposure, are uncommon but possible, being more frequent with minocycline and nitrofurantoin. Advanced liver fibrosis and cirrhosis is absent in those cases of late-onset AI-DILI cases. In a series of 24 patients with AI-DILI, none of them presented with cirrhosis or significant fibrosis at baseline. Our findings are consistent with the fact that advanced fibrosis (i.e. marked bridging fibrosis) was observed mainly among AIH cases, but not AI-DILI.

Our study has limitations. First, our sample size is small, and its statistical power was not sufficient for the analysis of some histopathological features that are likely to be diagnostically useful in conjunction by multivariate analysis. The follow-up in some patients was short, and progression to a fully developed AIH phenotype cannot be completely ruled out. We limited the assessment of histological features to variables with high reproducibility among pathologists. Features such as the so-called hepatocyte rosette and emperipolesis are difficult to evaluate and lack diagnostic value for AIH. For that reason, they were not included in this study.

Conclusions

In summary, AI-DILI exhibits the clinical and pathological features of AIH. Necroinflammatory changes, and infiltration of neutrophils or eosinophils are manifestations of hepatocellular damage, and are nonspecific findings. There is no individual histopathologic feature decisive for diagnosing AI-DILI over AIH. AIH can cause the full spectrum of liver injury, from mild fibrosis to established cirrhosis. Advanced stages of liver fibrosis can be used to favor the diagnosis of AIH over AI-DILI in patients with no history of liver disease. Definitive diagnosis of cases presenting with autoimmune liver injury and mild liver fibrosis must be made by follow-up. AI-DILI requires demonstration of complete remission after drug withdrawal and no need for continuous immunosuppression.

Conflict of interest

The authors have no conflict of interests related to this publication.

Author contributions

Designed the study (CAF, RJP), wrote the manuscript, designed the figures and table, reviewed the medical records and performed the statistical analyses (CAF), performed the pathology evaluation of liver biopsies (CAF, SA, KK, RJP). All authors drafted and approved the final version of this manuscript.

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