Pathological aspects of fatty liver disease

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Abstract
In a normal human liver 5% of its mass consists of lipids. When deposition of fat increases, the terms most often used are fatty liver or steatosis. This includes non-alcoholic fatty liver disease whose acronym is NAFLD and alcoholic liver disease (ALD). A liver biopsy is still considered to be the gold standard for determining the severity of liver damage in either of these entities.

Keywords
Steatosis, steatohepatitis, non-alcoholic steatosis, non-alcoholic steatohepatitis, NASH, alcoholic steatohepatitis, ALD, liver biopsy.

INTRODUCTION
Although numerous twentieth century studies showed a well-documented association between steatosis, cirrhosis, obesity, diabetes and alcohol, it was not until 1980 that Ludwig et al. at the Mayo Clinic published a study where the nature of nonalcoholic hepatic changes in steatosis and steatohepatitis were shown. In that study they coined the term NASH (Non-alcoholic steatohepatitis).

Since those first publications, fatty liver has become an increasingly common disease especially because of the increased prevalence of metabolic syndrome (obesity, hypertension, dyslipidemia, microalbuminuria and diabetes mellitus type II) in the general population. Currently, NAFLD is one of the most prevalent diseases worldwide.

These entities are accompanied by a spectrum of pathological changes ranging from simple steatosis to steatosis accompanied by inflammation with or without significant fibrosis to steatohepatitis. This can evolve into advanced chronic liver disease that progresses to cirrhosis, liver failure and increased risk of hepatocellular carcinoma (HCC). Approximately 30% of patients who have the sole initial finding of steatosis progress to cirrhosis. Of these 10% will develop HCC, and - over a period of 10 years - 30% to 40% will develop liver failure or die from their underlying disease (3).

Steatosis has been observed in 50% to 70% of patients with risk factors for NAFLD, especially obesity and type 2 diabetes. Of these 20% to 30% progress to steatohepatitis, and 2% to 3% progress to cirrhosis. The occurrence of cirrhosis has also been reported in between 5% and 15% of (missing word in original), with an overall mortality rate that ranges from 12% to 36% and a rate of mortality related to liver disease of between 2% and 7% which is significantly higher than in the general population. Morphological changes can be identical in both alcoholic and non-alcoholic etiologies (2, 4, 5).

There are still no clinical, diagnostic, laboratory or imaging methods that can establish with certainty the difference between simple steatosis and steatohepatitis or which can determine the degree of inflammatory activity and fibrosis and evaluate the existence of other coexisting disease processes. This is where the role of liver biopsy becomes importance for this condition.
DEFINITION

Fatty liver disease has traditionally been divided into alcoholic and non-alcoholic. Although their macro and microscopic appearances are very similar, their clinical signs and pathophysiology are different.

NAFLD is defined as the accumulation of liver fat that exceeds 5% to 10% of the liver’s weight. It is indispensable to discard substantial alcohol consumption (less than 20 g/day for women and less than 40 g/day for men). This directly correlates with the presence of fatty vacuoles in the cytoplasm of hepatocytes.

The condition known as nonalcoholic fatty liver disease (NAFLD) represents a spectrum of changes starting from a simple finding of steatosis alone which may be very mild and even have no inflammation. NAFLD’s evolution is benign, and its risk of chronicity with progression to cirrhosis is less than 5%. Steatohepatitis (NASH) steatosis and inflammation of variable intensity is always accompanied by hepatocellular lesions (ballooning and/or pericellular fibrosis) with progression to chronic disease and cirrhosis in 21% to 28% of cases. Progression to liver failure and hepatocellular carcinoma with its associated mortality occurs in 12% of cases (6, 7).

More than half of patients with cryptogenic cirrhosis have previously had a diagnosis of NASH or histological changes that suggest NAFLD. Up to 13% of them are associated with hepatocellular carcinoma (3).

RISK FACTORS

Many conditions are associated with the histological phenotype of NAFLD, but the vast majority of patients seen in clinical practice have metabolic syndrome with insulin resistance being the most important characteristic. This is the most common link of all etiologies associated with NAFLD. Metabolic syndrome, present in 22% to 30% of the population, increases the risk of cardiovascular morbidity and mortality and the risk of progression to fibrosis in nonalcoholic steatohepatitis (NASH). 90% of patients with NASH have metabolic syndrome. Between 70% and 80% of people with BMI >30 kg/m² have hepatic steatosis (8, 9, 10).

There are many other causes of fatty liver disease that are considered to be secondary. The size of the vacuole can help us in the search for an etiologic factor. For instance, a large vacuole is seen in rapid weight loss and malnutrition and is associated with Hepatitis C; genetic metabolic disorders such as Wilson’s disease, tyrosinemia and abetalipoproteinemia; other conditions such as congenital lipoatrophy and lipodystrophy acquired through HIV infection; large intestinal resections; gastroplasties and jejunoileal bypasses. Small vacuole steatosis is found in Reye syndrome, acute fatty liver during pregnancy, Hellp syndrome, metabolic errors such as acyl transferase lecithin deficiency, Wolman’s disease and deposition of cholesterol esters (10, 11).

Hepatic steatosis and steatohepatitis have also been linked with drugs. For example 1% to 3% of patients who use Amioradone for prolonged periods develop this condition. Perhexilene maleate (Pexid) and diethyl aminoethyl hexestrol hydrochloride (Coralgil) which are widely used in Europe and Japan can lead to the development of these conditions, as can calcium channel blockers such as nifedipine. Tamoxifen, isoniazid, estrogens, glucocorticoids, steroids and diethylstilbestrol have questionable etiologic associations, but their use has been proven to exacerbate NASH especially in patients with other risk factors. Methotrexate is also well known to cause steatosis, fibrosis and eventually cirrhosis. It has been suggested that its use increases chances of steatosis (12, 13, 14). Later we will expand on the findings observed in the parenchyma of the liver and those secondary to drug toxicity.

Predictors of severity and a greater probability of progression include ages over 45, AST/ALT greater than 1.0, obesity, type 2 diabetes mellitis, hypoalbuminemia, hypertension and the presence of liver fibrosis (15).

WHAT IS THE ROLE OF A LIVER BIOPSY?

Fatty liver disease has been identified as the most frequent cause of unexplained elevation of aminotransferases, but since a liver biopsy may also reveal other accompanying conditions in about 30% of cases, it remains the gold standard for confirmation of a diagnosis of NAFLD. A liver biopsy will show the clinical and pathological spectrum, and in many cases it will allow the clinician to establish the appropriate treatment. Repeated biopsies are used for evaluating treatment and for monitoring response to treatment especially in the context of research or use of new therapeutic methods.

When should a liver biopsy be performed? This remains a question that involves a lot of controversy. It is seen as a costly, invasive procedure with potential morbidity and vital risks. Therefore this decision should be made on an entirely individual basis that takes into account diagnostic images and especially the risk factors demonstrating progression of the condition. The main objectives of the biopsy should be to diagnose the type of steatohepatitis (NASH versus NAFL), to determine the stage of inflammatory activity or fibrosis, and to assess comorbid conditions and potential dual etiologies which are present in up to 20% of cases of ALD (16, 17).

HISTOPATHOLOGICAL FINDINGS

We will focus on the morphological changes necessary to establish a diagnosis and remember that in both alcoholic
and non-alcoholic etiologies the histological criteria of liver compromise may be indistinguishable. An NAFLD diagnosis is based on two major criteria. The first is determining the presence of fatty liver or steatohepatitis, the second is determining the nonalcoholic nature of the disease.

The morphological spectrum, and the severity and extent of each of the criteria evaluated varies from case to case. In early stages the changes are concentrated in zone 3 or the centrilobular zone. Then it progresses to the acinus, destroys the normal architecture, and becomes cirrhosis.

The main findings are given by steatosis, inflammation, hepatocellular lesions and fibrosis (17, 18).

**Steatosis**

Steatosis is present in 90% to 100% of cases. It is predominantly either macrovesicular or a combination of microvesicular and macrovesicular. It is localized primarily in zone 3 (Figures 1 and 2). A limit of more than 5% is used as a cutoff point to define significant steatosis, lesser amounts are considered to be within normal histological limits. Sometimes fat is minimal or practically disappears. This occurs in cases with significant fibrosis, some stages of cirrhosis, and when an alcoholic patient has stopped drinking (12). Steatosis alone is not a specific pattern and may have some mild inflammation.

**Inflammation**

A polymorphonuclear neutrophil (PMN) is a cell that is typically present in small clumps or sinusoids. The finding of PMNs surrounding ballooned hepatocytes is known as satellitosis (Figure 3). Lobular or acinar inflammation, usually mild with mixed infiltrates (lymphocytes and polymorphonuclear neutrophils), is considered to be one of the diagnostic features of steatohepatitis (18, 19). There are usually small amounts of acinar infiltrates, portal inflammation of lymphocytes and histiocytes, but these increase when they are accompanied by necroinflammatory activity superimposed on conditions such as viral hepatitis C, or cholestasis in acute alcoholic hepatitis (Figure 4) (20, 21).

**Hepatocellular lesions**

Hepatocellular lesions are essential for diagnosis of steatohepatitis. Steatosis alone, with or without inflammation, is not a sufficient criterion for diagnosis NAFLD: it must be accompanied by hepatocellular lesions. Lesion occurs in two forms: hepatocellular ballooning and/or subsinusoidal or pericellular fibrosis.
Ballooning is characterized by edema and cytoplasmic rarefaction resulting in enlarged cells with clear appearances with intracytoplasmic residual granular material located in Zone 3. In advanced stages the disease is adjacent to fibrous septa (Figure 5). As in other forms of hepatitis, isolated hepatocellular necrosis, acidophilic bodies and apoptotic bodies may be observed (18, 19).

**Fibrosis**

The distinctive fibrosis of steatohepatitis is pericellular and subsinusoidal. It is called chicken wire fibrosis due to the pattern caused by the collagen deposits in the space of Disse. It starts in zone 3 and can spread to the portal space. It occurs in the early stages of liver compromise (Figure 6).

In addition, fibrosis also develops around the terminal hepatic venules. This perivenular fibrosis or phlebosclerosis eventually completely obstructs the vein. In more advanced stages the combination of these progresses to replace scarred areas resulting in zone 3 sclerosing hyaline necrosis.

In endophlebitis there is inflammatory cells permeation areas next to terminal, sublobular or intercalated hepatic venules. Then it obliterates the lumen with subintimal proliferation and formation of veno-occlusive lesions. Portal and periportal fibrosis is unusual in early stages, although there may be evidence of veno-occlusive disease in portal veins.

In advanced stages of the disease bridging fibrosis extends from the perivenular areas to the portal tracts.
Characteristically cirrhosis is micronodular, but a mixed pattern of macronodular and micronodular can also be observed especially in those cases in which the initial stimulus is removed e.g. massive alcohol intake (22, 23).

Other findings

Megamitochondria can be observed in both NASH and ALH as small eosinophilic bodies between 3 and 10 microns. When they are negative with PAS staining it is indicative of high oxidative stress of the hepatocytes.

Glycogenated nuclei are observed in diabetic patients and in children with NASH.

Mallory bodies, also called Mallory-Denk bodies, appear in ballooned cells and are more abundant in zone 3. They are not well developed and are inconspicuous in NASH, but are prominent and well developed in AML (Figure 7).

Unusual findings that are more frequently found as part of the morphological chart of other conditions include microvesicular steatosis, less than 30% non-zonal macrovesicular steatosis, prominent portal or acinar inflammation, presence of an inflammatory infiltrate rich in plasma cells or eosinophils, epithelioid granulomas, portal or periportal fibrosis in the absence of pericellular fibrosis, chronic cholestasis with ductular proliferation and copper deposits and acute cholestasis with bile plug formation. Findings such as perivenular fibrosis, veno occlusive vascular lesions, phlebosclerosis and hyaline sclerosis. Intrahepatic iron deposits gradient from zone 1 to zone 3 (22, 23, 24, 25).

SPECIAL CONDITIONS

Steatohepatitis in children

For some authors the morphological differences observed in children suggest that this could be a different disease. Type 1 steatohepatitis occurs in approximately 17% of pediatric cases, and is most common among white girls. Changes are very similar to those observed in the adult population. Type 2 has slightly different features. 51% of cases occur in older children or adolescents, especially Hispanic, Asian and Native American males. It is more closely related to obesity in children. Morphological changes occur in zone 1 with increased inflammation and lymphocyte portal fibrosis with no ballooning (Figure 8). In the remaining 32% of cases we can observe a mixed pattern with overlapping of the two patterns described above (25, 26).

Alcoholic etiology

As mentioned, histopathological histories may be indistinguishable, but there is usually increased inflammatory acti-
vity. We will list some of the findings that may help suggest this etiology.

The presence of sclerosing hyaline necrosis is a combination of large quantities of well-formed Mallory-Denk bodies, perivenular fibrosis with type III collagen deposits, hepatocyte necrosis in zone 3 and veno-occlusive disease. This is the so-called alcoholic foamy degeneration corresponding to microvascular steatosis in zone 3 and cholestasis.

Noninvasive models have been proposed for differentiating between NASH and ALD. They use an ANI index that assesses the mean corpuscular volume, AST/ALT ratio, body mass index and gender. When this index is greater than 0 it supports an alcoholic etiology. However, these models have not been validated yet, and therefore liver biopsy is still used for diagnosis (16, 27).

**GRADING AND STAGING**

As with most liver diseases determining the stage of inflammatory activity is important for defining the prognosis and determining the appropriate therapy. The grade is defined by the extent of inflammation and hepatocellular lesions, and its stage is defined as the length and extent of fibrosis and architectural changes. Several groups have worked to create grading and staging schemes. The most frequently used is the Brunt’s semi-quantitative modified classification. No matter what system is used, it is advisable to always perform the most objective and proper possible classification of the morphological findings observed (18, 22, 23, 25, 28, 29). Tables 2 and 3 summarize the criteria to consider.

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<tr>
<th>Table 2. Steatohepatitis staging (Adapted from 19).</th>
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<tr>
<td><strong>Stage 1</strong> Low</td>
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<tr>
<td>Variable Steatosis of any stage, even greater than 66%, macrovesicular Ballooning: occasionally in zone 3, mixed lobular inflammation, absent or mild portal inflammation</td>
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<tr>
<td><strong>Stage 2</strong> Moderate</td>
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<tr>
<td>Steatosis of any stage, macrovesicular or mixed Ballooning: zone 3 PMN lobular inflammation, +/- chronic inflammation Absent, low or mild portal inflammation</td>
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<tr>
<td><strong>Grade 3</strong> Severe</td>
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<tr>
<td>Steatosis greater than 66%, mixed Lobular inflammation, mixed PMN infiltration in zone 3 Mild to moderate portal inflammation</td>
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<tr>
<th>Table 3. Steatohepatitis stages (Adapted from 19).</th>
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<tr>
<td><strong>Stage 1</strong> Perivenular fibrosis or in zone 3, pericellular or subsinusoidal 1a: If it is focalized 1b: If it is extended</td>
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<tr>
<td><strong>Stage 2</strong> Perivenular fibrosis or in zone 3, pericellular or subsinusoidal Portal fibrosis which may be focalized or extended</td>
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<tr>
<td><strong>Stage 3</strong> Portal fibrosis with bridge formation, which may be focalized or extended</td>
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<td><strong>Stage 4</strong> Established cirrhosis There may be residual pericellular fibrosis in the parenchyma of cirrhotic nodules</td>
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**REFERENCES**