Imaging patterns of fatty liver in pediatric patients

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ABSTRACT
We review imaging patterns of steatosis and new quantitative methods such as proton density fat fraction and magnetic resonance elastography for diagnosis of nonalcoholic fatty liver disease in children. Fatty liver can present as focal, diffuse, heterogeneous, and multinodular forms. Being familiar with various patterns of steatosis can enable correct diagnosis. In patients with equivocal findings on ultrasonography, magnetic resonance imaging can be used as a problem-solving tool. New techniques are promising for diagnosis and follow-up.

Nonalcoholic fatty liver disease (NAFLD) is as widely encountered in children as in adults, with an estimated prevalence of 9.6% (1). It occurs due to accumulation of triglyceride in hepatocytes without alcohol ingestion. Nonalcoholic steatohepatitis (NASH) was first defined in children in 1983 (2). NAFLD includes a broad range of clinicopathological features ranging from simple steatosis (fat with inflammation and/or fibrosis), steatohepatitis/NASH to cirrhosis. Some other diseases of liver can also cause hepatic steatosis including hepatitis B and C, Wilson’s disease, α-1-antitrypsin deficiency, autoimmune hepatitis, drug-induced liver injury (valproate, methotrexate, tetracycline, amiodarone, and prednisone), and total parenteral nutrition (3). Furthermore, fatty liver is a risk factor for cirrhosis, diabetes, and cardiovascular disease.

In clinical practice, the diagnosis of NAFLD is made by increased serum ALT and/or presence of enlarged echogenic liver in ultrasonography. Being overweight or obese, and/or insulin resistance are highly indicative but not absolutely necessary for diagnosing NAFLD (4). The gold standard for diagnosis is liver biopsy, which additionally provides semi-quantitative analysis of NASH damage in children (5). It is an expensive, invasive procedure with a risk of morbidity (0.06%–0.35%) and mortality (0.01%–0.1%) (6).

The evaluation of liver fat in children via noninvasive imaging modalities is needed to avoid complications of biopsy and for follow-up. Main imaging modalities for the assessment of pediatric NAFLD are ultrasonography (US) and magnetic resonance imaging (MRI). Computed tomography is the other imaging method for liver fat assessment, but ionizing radiation is a major drawback in children (7). Assessment of fat accumulation may cause diagnostic dilemmas and confusion due to manifestations with unusual structural patterns and imaging appearance of the liver. This article reviews the histopathology of pediatric NAFLD, radiologic evaluation and different structural patterns of childhood NAFLD/NASH on US and MRI. We also discuss diagnostic pitfalls and briefly review new imaging techniques.

Histopathology of pediatric fatty liver
Pediatric studies of NAFLD/NASH have described significantly different histopathological findings from the adult disease (8). The general findings in adulthood, which are designated in children as “type 1” includes macrovesicular steatosis, lobular inflammation, and ballooning degeneration, usually with poorly formed Mallory hyaline with or without perisinusoidal fibrosis with a zone 3 (perivenular) distribution. Pediatric “type 2” NASH is characterized by macrovesicular steatosis with...
portal inflammation and/or fibrosis. Perisinusoidal fibrosis or evidence of ballooning degeneration is not present (Fig. 1). Schwimmer et al. (5) characterized the liver biopsy results of pediatric patients with NAFLD and formed histologic definition of pediatric NASH. In contrast with adult patients, NASH in pediatric patients occurred with more portal inflammation (70%) and more portal fibrosis (60%) in biopsies. Type 2 NASH was detected in 51% and type 1 NASH in 17% of the study population. In type 2, children were younger, severity of obesity was greater, and advanced fibrosis was more common compared with type 1 NASH. Boys were significantly more likely to have type 2 NASH than girls.

Radiologic evaluation of pediatric fatty liver

Ultrasonography

US is the most preferred imaging technique for the assessment of hepatic steatosis due to low cost, easy accessibility, and outstanding safety. Normal liver echogenicity is similar to renal cortex or spleen echogenicity. Normal liver has intrahepatic vessels that are plainly demarcated and posterior parts are clearly illustrated. On the other hand, fat accumulation in the liver increases echogenicity and this can lead to indistinction of vessels and bile ducts and blurring of the diaphragm due to depth-dependent signal reduction (9). Despite mentioned advantages of US, operator and machine dependency, lack of objective quantification and significant decrease in sensitivity and specificity in morbid obesity are the major limitations for evaluation of hepatic steatosis (10). Furthermore, when biopsy-proven steatosis ratio is less than 30%, the sensitivity of US diminishes remarkably (7). Pathologic conditions such as fibrosis and inflammation may intensify liver echogenicity and can result in misinterpretations (11, 12). Given all these limitations, US evaluation of hepatic steatosis should be performed with attention and interpreted cautiously, and should not be recommended routinely as a sole diagnostic or monitoring tool for NAFLD in children.

Magnetic resonance imaging

Different methods are applied for the evaluation of liver fat by MRI, but the most widely preferred method is 2-point Dixon technique, also known as chemical shift imaging or dual-echo method (13, 14). In this technique, two sets of gradient-echo images of the liver are obtained and echo-time-dependent signal interference between fat and water is considered. On in-phase echo-time, water and fat signals add up and therefore, the total signal intensity is higher. On out-of-phase echo-time, water and fat signals cancel each other and consequently the total signal intensity diminishes. Healthy liver has no difference in signal intensities between the in-phase and out-of-phase images, however, in case of fat accumulation, the liver signal intensity decreases on the out-of-phase image. This imaging method is reliable in the absence of magnetic field inhomogeneity and iron deposition. The main drawback is that the quantity of water and fat affects the signals from fat and water; this can be managed by acquiring new images with variable T1-weighting by applying two flip angles (15). High-flip-angle imaging is desirable for uncovering small amounts of fat in tissues that include mainly water; low-flip-angle imaging should be applied for revealing small amounts of water in fat-rich tissues (14).

Magnetic resonance spectroscopy and proton density fat fraction

Magnetic resonance spectroscopy (MRS) is established as the most reliable
technique for the evaluation of hepatic steatosis, as it can detect the amount of water and fat in the liver. The prevailing frequency of hydrogen protons in water molecules in the MRI spectrum is at 4.7 ppm, while hydrogen protons in triglyceride molecules have lower values, 1.2 ppm. However, hydrogen protons in other triglyceride moieties have extra peaks at other frequencies (16). Accordingly, water has a single large peak in contrast to triglyceride, which has multiple frequency peaks. In normal cases, the dominant water peak is clear, and no triglyceride peaks are present. Existence of liver fat permits evaluation of a hepatic fat fraction via the area-under-water peak versus the area-under-fat peaks (17).

Proton density fat fraction (PDFF) measurement by MRI appears to be the most objective test for fat tissue quantification, as it removes T2* effect, T1 bias, spectral complexity of fat, eddy currents, noise bias, and magnetic inhomogeneity from the image and measures six peaks of fat. PDFF is the ratio of density of mobile fat protons (primarily triglycerides) and the total density of protons from mobile fat and mobile water. Moreover, it allows objective fat quantification and grading similar to MRS in a breath-hold time in patients with NAFLD (18). The practical strength of PDFF is recording of PDFF values from PDFF maps (Fig. 2). MRI-based PDFF maps or MRS-based PDFF values can be rapidly produced within seconds. Above all, it is very easy and practical for fat fraction measurement. In pediatric patients comparison of biopsy and PDFF showed promising results (19). PDFF is an unconfounded and fundamental property of tissue, and it is insensitive to changes in acquisition parameters which makes it a robust quantitative biomarker (20). PDFF is stable across scanner platform, scanner manufacturer, imaging center, and even field strength and can standardize MRI-based fat quantification (19). Moreover, simultaneous calculation of R2* and T2* maps from the same MRI sequence enables estimation of hepatic iron content, which can be important for diagnosis of coexisting fatty liver and hepatic iron overload (Fig. 3) (21, 22).

**Magnetic resonance elastography**

Nonalcoholic fatty liver disease has the potential to progress into hepatic fibrosis. Early diagnosis of fibrosis is important because, if treated, the degree of fibrosis can be minimized or reversed. Magnetic resonance elastography (MRE) is suggested to evaluate the stiffness of liver parenchyma (23). It allows detection of fibrosis and differentiation of low-grade fibrosis from high-grade and also it may be feasible to distinguish steatosis from steatohepatitis (24, 25). The MRE pulse sequences developed for adults have been modified for pediatric patients. The absorption rate and field of view are adopted for pediatric age group; additionally, the power of driver is decreased by about 20% for patients between five and 18 years of age and by 40%–50% for patients younger than two years, compared with the settings used in adult patients (26). The reason to decrease power is to reduce the abdominal pain. Normal values of stiffness for pediatric age group have not been suggested for MRE, because liver stiffness was found to be not related with age by the ultrasound-based transient elastography. The success rate and accuracy of MRE is higher than ultrasound-based transient elastography (27). Furthermore, more of the liver can be evaluated with deep penetration, and ascites and obesity are not obstacles for examination with MRE. The drawbacks are higher cost and presence of iron load that makes detection of shear waves very difficult.
**Patterns of fat deposition**

*Diffuse homogeneous and heterogeneous deposition*

Diffuse fat deposition is the most frequently encountered pattern in the liver, and the distribution of fat is generally homogeneous (Fig. 4). Multifocal sparing can be seen in patients with diffuse steatosis (Fig. 5). Sometimes diffuse heterogeneous steatosis can be seen on US as scattered echogenic areas, which can be mild, moderate, or severe. MRI may be required to diagnose this condition (Figs. 6 and 7). Also areas of fat sparing in diffuse steatosis can mimic lesions, and MRI can allow diagnosis in patients with equivocal findings on US (Fig. 4b).

**Focal fat deposition**

Focal fat deposition is the second most common involvement. Focal fat accumulation generally exists adjacent to the falciform ligament, in the porta hepatis, and in the gallbladder fossa. This distribution has been linked with variant venous circulation, such as anomalous gastric venous and pancreaticoduodenal venous drainage (28, 29). Focal fat depositions have been noticed emerging next to insulinoma metastases and they may cause diagnostic confusion by mimicking mass lesions. This appearance was hypothesized to be related to local insulin activity on hepatocyte triglyceride synthesis (30, 31). Specific location and chemical shift MRI are key findings for overcoming diagnostic dilemmas and hurdles.

**Multifocal fat deposition**

Multifocal fat deposition is an unusual distribution in childhood (Fig. 8). Multiple foci of fat accumulation are scattered in atypical locations all around the liver (32, 33). The fat centers mimic true nodules, which can be the source of diagnostic confusion especially in preexisting malignancy. Under such conditions, in-phase and out-of-phase MRI is superior to computed tomography and US. Absence of a mass effect, lack of hypointensity on in-phase image, absence of restriction on diffusion-weighted images and stability

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**Figure 5.** a, b. A 15-year-old girl with diffuse heterogeneous liver and subtle hypoechogenic areas on US. In-phase (a) and out-of-phase (b) images show diffuse heterogeneous liver steatosis. Diffuse heterogeneous signal drop is seen on out-of-phase image consistent with steatosis. Note the submillimeter focal sparing areas within the heterogeneous steatosis consistent with multifocal fat sparing.

**Figure 6.** a–c. US image (a) shows focal hyperechogenic areas in segments 7-8 of a 10-year-old boy (arrows). In-phase (b) and out-of-phase (c) images show mild heterogeneous liver steatosis. Out-of-phase image shows subtle signal drop compared with in-phase image confirming mild heterogeneous steatosis (arrows in c).

**Figure 7.** a–c. US image (a) shows multiple hyperechoic lesions in the right liver lobe of a six-year-old boy. In-phase (b) and out-of-phase (c) images show severe heterogeneous liver steatosis. Out-of-phase image shows multiple lesions with signal drop compared with in-phase image consistent with severe heterogeneous steatosis (arrows in c).
in size during follow-up are distinguishing features of multifocal fat areas.

**Pitfalls: fat-containing hepatic tumors**

Hepatic adenomas, hepatocellular carcinomas, and regenerative nodules may have intracellular lipid (Fig. 9) (34, 35). However, postcontrast imaging features of these lesions allow differentiation from areas of focal steatosis.

**Conclusion**

Fatty liver is a common clinical problem for children and adolescents. NAFLD constitutes a risk for diabetes, cardiovascular disease, cirrhosis, and end-stage liver disease. Currently, liver biopsy is the clinical standard for diagnosing and grading NAFLD. Noninvasive imaging modalities to assess liver fat in children have gained importance for avoiding invasive procedures and were proven to be reliable. The available evidence does not suggest the use of US as a sole method for the diagnosis or grading of fatty liver in children.

Chemical shift GRE MRI is more trustworthy than US for evaluation and diagnosis of steatosis. The most common imaging pattern of steatosis is diffuse homogeneous fat deposition. Less common patterns include focal deposition, diffuse heterogeneous deposition, and multifocal deposition. These patterns may mimic neoplasms, leading to confusion and unnecessary diagnostic invasive procedures. Assessment of the fat content of the lesion, location, morphologic features, contrast enhancement, and mass effect usually permits a correct diagnosis.

**Conflict of interest disclosure**

The authors declared no conflicts of interest.

**References**

7. Saadeh S, Younossi ZM, Remer EM, et al. The utility of radiological imaging in nonalcoholic fatty liver disease. Gastroenterology 2002; 123:745–750. [CrossRef]

**Fatty liver imaging in pediatric patients**

**Figure 8.** a–c. US image (a) shows multiple hyperechoic lesions in the liver of a nine-year-old boy. In-phase (b) and out-of-phase (c) images show multifocal liver steatosis. Out-of-phase image shows multiple lesions with signal drop compared with in-phase image consistent with multifocal steatosis.

**Figure 9.** a, b. A one-year-old girl with tyrosinemia type 1. In-phase (a) and out-of-phase (b) images show multifocal fat containing lesions. Out-of-phase image shows multiple lesions with signal drop compared with in-phase image consistent with multifocal fat containing lesions diagnosed as multiple regenerative nodules, which were stable in size during the follow-up.

7. Saadeh S, Younossi ZM, Remer EM, et al. The utility of radiological imaging in nonalcoholic fatty liver disease. Gastroenterology 2002; 123:745–750. [CrossRef]
31. Sohn J, Siegelman E, Osiason A. Unusual patterns of hepatic steatosis caused by the local effect of insulin revealed on chemical shift MR imaging. AJR Am J Roentgenol 2001; 176:471–474. [CrossRef]