Non-invasive scoring systems for predicting NASH in Japan: evidences from Japan Study Group of NAFLD

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Abstract
A disease spectrum of nonalcoholic fatty liver disease (NAFLD) ranges from nonalcoholic fatty liver (NAFL), a benign condition to nonalcoholic steatohepatitis (NASH) which can lead to liver-related mortality such as hepatocellular carcinoma, hepatic failure, and portal hypertension. Although liver biopsy is the gold standard for the diagnosis of NASH, there have been several drawbacks such as complication, cost, and sampling errors. A reliable and inexpensive noninvasive parameter for predicting NASH or severe hepatic fibrosis is required in patients with NAFLD. In this article, we reviewed several scoring systems which have been constructed by the Japan Study Group of NAFLD (JSG-NAFLD), a multi-center study group for the diagnosis and management of NAFLD in Japan. First of all, the NAFLD activity score calculated from the ferritin, insulin, and type 4 collagen 7S levels was established to diagnose NASH. We recently suggest that modified NAFLD score may be better than conventional NAFIC score, although external validation studies using a large population of NAFLD are essential. In order to exclude severe fibrosis (stage 3-4), FIB4 index and NAFLD fibrosis score (NFS) are the most simple and reliable scoring systems in consistent with the Western studies. The combination of platelet count and AST/ALT ratio (PAAR index) is useful and easily determined for excluding severe fibrosis. To detect cirrhosis (stage 4), we formed an easily calculated composite score, called the PLALA (platelet, albumin, AST/ALT ratio) score. Introduction of these scores in clinical practice may reduce the proportion of patients that require liver biopsy to diagnose mild disease.

Introduction
Nonalcoholic fatty liver disease (NAFLD) is an important cause of chronic liver injury in many countries including Japan. NAFLD represents a wide spectrum of conditions that are characterized histologically by macrovesicular hepatic steatosis (>5% of hepatocytes affected), and the diagnosis is made after excluding a history of consumption of alcohol in amounts sufficient to be considered harmful to the liver. NAFLD ranges over a wide spectrum, extending from nonalcoholic fatty liver (NAFL) which is generally benign, through to nonalcoholic steatohepatitis (NASH) to liver cirrhosis, end-stage liver disease, and even hepatocellular carcinoma (HCC) despite the absence of significant alcohol consumption [1]. The probability of developing advanced fibrosis and HCC is significantly greater in individuals with NASH than in those with NAFL. Liver biopsy as the gold standard tool of NASH can reveal the histologic steatosis with hepatocyte ballooning. However, it is difficult to perform liver biopsy for every patient with NAFLD to ascertain the presence of NASH and determine the stage and grade of the disease. In 2010, we established the Japan Study Group of NAFLD (JSG-NAFLD), represented by the following 10 hepatology centers: Yokohama City University, Asahikawa Medical College, Kurume University, Nara City Hospital, Hiroshima University, Saga Medical School, Osaka City University, Kyoto Prefectural University of Medicine, Kochi Medical School, and Saiseikai Suita Hospital. In 2014, JSG-NAFLD includes eleven hepatology centers through all over Japan. The aim of our study group is 1) to clarify prevalence, natural history, and prognosis in Japanese NAFLD, 2) to establish and validate parameters or scoring systems for the noninvasive diagnosis of NASH or severe fibrosis, and 3) to develop the pharmacological treatments for NASH.

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Scoring systems for predicting NASH in NAFLD

Several lines of evidences have confirmed that NASH cannot be differentiated from NAFL by routine clinical parameters. Serum transaminase activities are known to be usually elevated in patients with NAFLD, but those levels do not always reflect severity of NAFLD. Two representative cases were shown in Figure 2. A 27 year-old male patient with NAFLD had elevated transaminase activities (AST/ALT 74/233 IU/L). His histological examination revealed hepatic steatosis (>66% of hepatocytes affected). He was diagnosed NAFL (Matteoni type 2, NAS score: 3+1+0=4) (Figure 2a). On the other hand, a 57 year-old female patient with NAFLD almost mildly elevated levels of transaminase activities (AST/ALT 38/40 IU/L). Her liver histology revealed moderate steatosis with hepatocyte ballooning and extensive fibrosis. She was diagnosed NASH stage 3-4 (Matteoni type 4, NAS score: 2+1+2+5) (Figure 2b). A variety of parameters has been suggested to detect NASH [2], including oxidative stress markers (thioredoxin, lipidperoxides), iron-related parameters (ferritin), hepatic fibrosis markers (type1 collagen 7S, procollagen III peptide), and insulin resistance (HOMA-IR), adipokines (adiponectin, leptin), and apoptosis markers (soluble Fas, cytokeratin 18 (CK-18)). Among those, circulating levels of CK18 fragments have been investigated extensively as novel biomarkers for the presence of steatohepatitis in patients with NAFLD [3]. Although these are very encouraging results, currently this assay is not commercially available. Furthermore, as each study utilized a study-specific cut-off value, there is not an established cut-off value for identifying steatohepatitis. According to the guidelines of the American Association for the Study of Liver Diseases (AASLD) [1], CK18 is not recommended in routine clinical practice. The first aim of JSG-NAFLD is to construct a simple index for detecting NASH among a large population of NAFLD in Japan. NAFIC score assigns one point for 200 (female) or 300 (male) ng/ml or higher ferritin, one point for 10 μU/ml or higher fasting insulin, and two points for 5.0 ng/ml or higher type 4 collagen 7S. The total of these points is regarded as the NAFIC score (Table 1), and the possibility of NASH is high when the score is two or higher [4]. NAFIC score was 1 in the NASH case (Figure 2a), while that was 4 in the NASH case (Figure 2b). A limitation of NAFIC score is low sensitivity and negative predictive value (NPV). At a cutoff value of NAFIC score 2, sensitivity, specificity, positive predictive value (PPV), and NPV were 60, 87, 85, and 64%, respectively. We reported the usefulness of the modified NAFIC scoring system, modified by changing the weightage assigned to the serum insulin levels, to obtain a reduced number of false-negative cases. Modified NAFIC is calculated as follows: serum ferritin ≥ 200 ng/ml (female) or ≥ 300 ng/ml (male), 1 point; serum type IV collagen 7S ≥ 5.0 ng/ml, 2 points; serum fasting insulin 10-15 μU/ml, 1 point and ≥ 15 μU/ml, 2 points. The modified NAFIC score showed improved sensitivity and NPV for the diagnosis of NASH [5]. Three parameters including serum ferritin, insulin, type4 collagen 7S can help us pick up NASH in NAFLD.

Scoring systems for predicting severe fibrosis in NAFLD

Since patients with severe fibrosis (stage3 or 4) are at high risk for the development of HCC [6,7], noninvasive diagnosis of severe fibrosis is one of the fields that has evolved most rapidly in recent years [8]. The NAFLD fibrosis score (NFS) [9], BARD score [10], APRI (AST to platelet ratio index) [11], and FIB4 index [12] are among the more widely investigated non-invasive tools to cross-sectionally predict advanced fibrosis in NAFLD. NFS is comprised of six parameters such as diabetes/impaired fasting glucose, age, AST, ALT, platelets, BMI, and albumin (http://www.nafldscore.com/). Of the four scoring systems, NFS has received the most extensive validation [4,13,14] and it has been recommended for clinical use in the recent practice guideline on the diagnosis and management of NAFLD [1]. The BARD score developed by Harrison et al. represents the weighted sum of three easily available variables (BMI ≥28 kg/m2 [=1 point], AAR ≥0.8 [=2 points], and DM [=1 point]), and the authors demonstrated that a score of 2–4 was associated with an odds ratio of 17 for predicting advanced fibrosis [10]. In Japan, however, Fuji and colleagues reported significantly poorer applicability of BARD in Japanese patients with NAFLD, compared with Caucasian subjects [15]. BARD score is less predictive of advanced fibrosis in Japanese NAFLD patients because they are less obese than those in western countries. Another disadvantage of BARD score is low PPV. The BARDI score, an enhanced model of BARD, was created by adding the international normalized ratio (INR) to the BARD score [16]. The BARDI score had an improved PPV over the BARD score and maintained an excellent NPV. APRI was calculated as the following formula: APRI=(AST/upper limit)/platelet count [17]. The area under receiver operating characteristic (ROC) curve for APRI was 0.85 with an optimal cut-off of 0.98, giving a sensitivity of 75% and a specificity of 86%. The usefulness of APRI is equal to that of NFS for predicting severe fibrosis in NAFLD [11]. In contrast, the diagnostic accuracy of APRI or BARD score seems to be inferior to other complex invasive scoring systems [18]. FIB4 index (http://gihep.com/...
fibrosis in NAFLD. NAFLD patients with platelet count >19.5×10^4/μL and AAR can be reliable and simple parameters for predicting severe fibrosis. Thus, we have hypothesized that the combination of platelet count and AAR can be useful and easily determined even in routine clinical practice or health checkups. PAAR index was 0 in the NAFL case (Figure 2a) and 2 in the NASH case (Figure 2b). The platelet count and AAR (PAAR index) has been known to decrease with increasing hepatic fibrosis in diabetic patients, because it is simple, easy to measure and handle, cost-effective, and accurate for diagnosis of the severity of fibrosis.

### Scoring systems for predicting cirrhosis in NAFLD

At first, Fujii and colleagues who are members of JSG-NAFLD, confirmed the clinical usefulness of two scoring systems such as cirrhosis discriminant score (CDS), and the hepatitis C antiviral long-term treatment against cirrhosis (HALT-C) model [31]. These scores consist of platelet count, AAR, and PT-INR. Our study group (JSG-NAFLD) tried to originally develop a simple scoring system to differentiate cirrhosis from non-cirrhosis in NAFLD patients. The three variables platelet, albumin, and AAR were combined in an unweighted sum (platelet < 15.3×10^4/μL; 1 point, serum albumin <4.0 g/dL; 1 point, and AAR >0.9; 1 point) and formed an easily calculated composite score for predicting cirrhosis in NAFLD patients, called the PLALA score. A PLALA score (2 and 3) was useful for detecting severe fibrosis even in NAFLD patients with normal ALT levels [27,28].

### Table 1. Formula of noninvasive scoring systems for detecting NASH or severe fibrosis.

| Formula | Equation |
|---------|----------|
| NAFIC score | Ferritin ≥ 200 (female) or 300 (male) ng/ml = 1 point | 
| | Iron ≥ 10 μU/ml = 1 point | 
| | Type IV collagen 7s > 5.0 ng/ml = 2 points | 
| | Scale: 0-4 | 
| Modified NAFIC | Ferritin ≥ 200 (female) or 300 (male) ng/ml = 1 point | 
| | Iron ≥ 10 μU/ml = 1 point, ≥ 15 μU/ml = 2 points | 
| | Type IV collagen 7s > 5.0 ng/ml = 2 points | 
| | Scale: 0-5 | 
| NAFLD fibrosis score | -1.675 + 0.037 × age (years) + 0.094 × BMI (kg/m^2) + 1.13 × IFG/diabetes (yes = 1, no = 0) + 0.99 × AST/ALT ratio (AAR) − 0.013 × platelet count (×10^9/L) − 0.66 × albumin (g/dL) | 
| FIB4 index | (Age [years] × AST [IU/L])/(platelet count [×10^9/L] ×√ALT [IU/L]) | 
| PAAR index | Platelet < 19.5×10^4/μL = 1 point | 
| | AST/ALT ratio (AAR) ≥ 0.8 = 1 point | 
| | Scale 0-2 | 
| PLALA score | Platelet < 15.3×10^4/μL = 1 point | 
| | Albumin < 4.0 g/dL = 1 point | 
| | AST/ALT ratio (AAR) ≥ 0.9 = 1 point | 
| | Scale 0-3 | 

The distribution of FIB4 index was examined in 5,410 people with NAFLD who were diagnosed at health checkups. Overall, 4729 (87.4%) had FIB4 index between indeterminate range (1.45-3.25) and 27 (0.5%) had FIB4 index below the low cut-off point (<1.45), 654 (12.1%) had FIB4 index >3.25 and 11 (0.2%) had FIB4 index >6.5. The distribution of FIB4 index is shown in Figure 3. On the basis of these results, the guideline committee of Japan Society of Gastroenterology recommended the FIB4 index as a useful simple method of identifying patients with fibrosis. Platelet count has been known to decrease with increasing hepatic fibrosis in chronic liver disease. Previously we reported that the platelet count may be an ideal biomarker of the severity of fibrosis also in NAFLD patients on the basis of a multicenter study by the JSG of NAFLD [30]. Thus, we have hypothesized that the combination of platelet count and AAR can be useful and easily determined even in routine clinical practice or health checkups. PAAR index is shown in Figure 2b. The platelet count and AAR (PAAR index) can be used as a simple scoring system to differentiate cirrhosis from non-cirrhosis in NAFLD patients. The three variables platelet, albumin, and AAR were combined in an unweighted sum (platelet < 15.3×10^4/μL; 1 point, serum albumin <4.0 g/dL; 1 point, and AAR >0.9; 1 point) and formed an easily calculated composite score for predicting cirrhosis in NAFLD patients, called the PLALA score. A PLALA score (2 and 3) was useful for detecting liver cirrhosis in NAFLD patients (sensitivity, 86.8%; specificity, 90.8%; NPV, 99.5%; PPV, 26.2%) (Figure 3) [32].
Scoring systems for predicting all-cause or liver related mortality and systemic complications in NAFLD

A major limitation of these scoring systems is that they have largely investigated in cross-sectional studies and thus their utility in monitoring disease natural history, predicting outcomes or response to therapeutic intervention remains unknown. Ideal non-invasive scoring systems can predict all-cause or liver related mortality and systemic complications in NAFLD. It was recently reported that the scores derived from the fibrosis-predicting scoring systems NFS, APRI, and FIB4 index also serve as prognostic factors [33]; the prognostic values of these scores still need to be verified in Japan. In Japan, Kawamura et al. reported that the annual liver carcinogenic rate in NAFLD patients was 0.043%, and that APRI was useful in predicting liver carcinogenesis [34]. Among previously established scoring systems, NFS is the best predictor for mortality in US and China [35-37]. NFS and FIB4 index could identify patients at risk of onset of DM and cerebral-cardiovascular diseases [38]. In the future, we should clarify if our proposed scoring systems (NAFIC score, PAAR index, and PLALA score) predict cancer incidence (hepatic/extrahepatic) and all-cause or liver-related mortality.

Conclusions

In this article, we extensively reviewed several scoring systems to detect NASH or severe fibrosis in NAFLD, mainly on the basis of data from the JSG-NAFLD study in Japan. Although external validation or longitudinal studies are required for practical use, these scoring systems can be easily determined and useful for the management of NAFLD (Figure 3).

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