Assessing liver fibrosis without biopsy in patients with HCV or NAFLD

ABSTRACT

Staging of liver fibrosis is increasingly done using non-invasive methods, in some cases obviating the need for liver biopsy. Scores based on laboratory values and demographic variables have been developed and validated for assessing fibrosis in patients with hepatitis C virus (HCV) infection and nonalcoholic fatty liver disease (NAFLD), as have several imaging methods that measure shear-wave velocity, a reflection of fibrosis severity.

KEY POINTS

Liver biopsy remains the gold standard for determining fibrosis stage but is expensive and entails risk of complications.

For patients infected with HCV, fibrosis stage should be determined with transient elastography, a transthoracic ultrasonographic technique that measures shear-wave velocity.

For patients with cirrhosis, transient elastography combined with a platelet count can detect developing portal hypertension and determine whether to screen for esophageal varices.

For NAFLD, combined elastography and NAFLD fibrosis score—which incorporates patient characteristics and laboratory test results—should be used to determine the need for liver biopsy.

Staging of liver fibrosis, important for determining prognosis in patients with chronic liver disease and for the need to start screening for complications of cirrhosis, was traditionally done only by liver biopsy. While biopsy is still the gold standard method to stage fibrosis, noninvasive methods have been developed that can also assess disease severity.

This article briefly reviews the epidemiology and physiology of chronic liver disease and the traditional role of liver biopsy. Pros and cons of alternative fibrosis assessment methods are discussed, with a focus on their utility for patients with nonalcoholic fatty liver disease (NAFLD) and hepatitis C virus (HCV) infection.

■ CHRONIC LIVER DISEASE: A HUGE HEALTH BURDEN

Chronic liver disease is associated with enormous health and financial costs in the United States. Its prevalence is about 15%,1 and it is the 12th leading cause of death.2 Hospital costs are estimated at about $4 billion annually.3

The most common causes of chronic liver disease are NAFLD (which may be present in up to one-third of the US population and is increasing with the epidemic of obesity), its aggressive variant, nonalcoholic steatohepatitis (NASH) (present in about 3% of the population), and HCV infection (1%).4,5

Since direct-acting antiviral agents were introduced, HCV infection dropped from being the leading cause of liver transplant to third place.6 But at the same time, the number of patients on the transplant waiting list who have NASH has risen faster than for any other cause of chronic liver disease.7
ASSESSING LIVER FIBROSIS

FIBROSIS: A KEY INDICATOR OF DISEASE SEVERITY

With any form of liver disease, collagen is deposited in hepatic lobules over time, a process called fibrosis. Both HCV infection and NASH involve necroinflammation in the liver, hepatocyte apoptosis, and activation of stellate cells, leading to progressive collagen deposition in hepatic lobules. Fibrosis typically starts in the region of the central vein and portal tracts and eventually extends to other areas of the lobule.

Determining fibrosis severity is critical when a patient is diagnosed with chronic liver disease, as it predicts long-term clinical outcomes and death in HCV and NAFLD. Different staging systems have been developed to reflect the degree of fibrosis, based on its distribution as seen on liver biopsy (Table 1, Figure 1).

In HCV infection, advanced fibrosis is defined as stage 4 to 6 using the Ishak system or stage 3 to 4 using the Meta-analysis of Histological Data in Viral Hepatitis (META-VIR) system. In NAFLD, advanced fibrosis is defined as stage 3 to 4 using the NASH Clinical Research Network system.

Staging fibrosis is also important so that patients with cirrhosis can be identified early to begin screening for hepatocellular carcinoma and esophageal varices to reduce the risks of illness and death. In addition, insurance companies often require documentation of fibrosis stage before treating HCV with the new direct-acting antiviral agents.

LIVER BIOPSY IS STILL THE GOLD STANDARD

Although invasive, liver biopsy remains the gold standard for determining fibrosis stage. Liver biopsies were performed “blindly” (without imaging) until the 1990s, but imaging-guided biopsy using ultrasonography was then developed, which entailed less pain and lower complication and hospitalization rates. Slightly more hepatic tissue is obtained with guided liver biopsy, but the difference was deemed clinically insignificant. Concern initially

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**TABLE 1**

<table>
<thead>
<tr>
<th>Stage</th>
<th>Ishak staging for HCV (^{10})</th>
<th>META-VIR staging for HCV (^{11})</th>
<th>NASH Clinical Research Network staging system for NAFLD (^{12})</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No fibrosis</td>
<td>No fibrosis</td>
<td>No fibrosis</td>
</tr>
<tr>
<td>1</td>
<td>Expansion of some portal areas with or without septa</td>
<td>Portal fibrosis without septa</td>
<td>Perisinusoidal or periportal fibrosis</td>
</tr>
<tr>
<td>2</td>
<td>Expansion of most portal areas with or without septa</td>
<td>Portal fibrosis with rare septa</td>
<td>Perisinusoidal and portal/periportal fibrosis</td>
</tr>
<tr>
<td>3</td>
<td>Expansion of most portal areas with occasional portal-portal bridging</td>
<td>Portal fibrosis with numerous septa</td>
<td>Bridging fibrosis</td>
</tr>
<tr>
<td>4</td>
<td>Expansion of portal areas with marked bridging (portal-portal, portal-central)</td>
<td>Cirrhosis</td>
<td>Cirrhosis</td>
</tr>
<tr>
<td>5</td>
<td>Marked bridging with occasional nodules (incomplete cirrhosis)</td>
<td>Cirrhosis</td>
<td>Cirrhosis</td>
</tr>
</tbody>
</table>

HCV = hepatitis C virus; META-VIR = Meta-analysis of Histological Data in Viral Hepatitis; NAFLD = nonalcoholic fatty liver disease; NASH = nonalcoholic steatohepatitis

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Fibrosis: a process of collagen deposition in hepatic lobules

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Figure 1. Findings on liver biopsy in nonalcoholic fatty liver disease and hepatitis C virus infection.

Expansion of portal tract by fibrosis (trichrome stain, × 100)

In the setting of steatohepatitis, fibrosis starts in the centrilobular perisinusoidal region (trichrome stain, × 100)

A subset of nonalcoholic steatohepatitis cases reveals only portal-based fibrosis, consistent with stage 1C (left); the centrilobular region seen on the right is intact (trichrome stain, × 100)

Periportal fibrosis is characterized by expansion of portal tracts with irregular border and focal entrapment of hepatocytes (trichrome stain, × 200)

Bridges of fibrous tissue connecting two portal tracts in a hepatitis C case, supporting stage 3 (bridging fibrosis) (trichrome stain, × 100)

End-stage liver disease with multiple cirrhotic nodules, surrounded by fibrous bands (trichrome stain, × 100)
arose about the added cost involved with imaging, but imaging-guided biopsy was actually found to be more cost-effective.14

In the 2000s, transjugular liver biopsy via the right internal jugular vein became available. This method was originally used primarily in patients with ascites or significant coagulopathy. At first, there were concerns about the adequacy of specimens obtained to make an accurate diagnosis or establish fibrosis stage, but this limitation was overcome with improved techniques.15,16 Transjugular liver biopsy has the additional advantage of enabling one to measure the hepatic venous pressure gradient, which also has prognostic significance; a gradient greater than 10 mm Hg is associated with worse prognosis.17

Disadvantages of biopsy: Complications, sampling errors
Liver biopsy has disadvantages. Reported rates of complications necessitating hospitalization using the blind method were as high as 6% in the 1970s,18 dropping to 3.2% in a 1993 study.19 Bleeding remains the most worrisome complication. With the transjugular method, major and minor complication rates are less than 1% and 7%, respectively.15,16 Complication rates with imaging-guided biopsy are also low.

Liver biopsy is also prone to sampling error. The number of portal tracts obtained in the biopsy correlates with the accuracy of fibrosis staging, and smaller samples may lead to underestimating fibrosis stage. In patients with HCV, samples more than 15 mm long led to accurate staging diagnosis in 65% of patients, and those longer than 25 mm conferred 75% accuracy.20 Also, different stages can be diagnosed from samples obtained from separate locations in the liver, although rarely is the difference more than a single stage.21

Histologic evaluation of liver biopsies is operator-dependent. Although significant interobserver variation has been reported for degree of inflammation, there tends to be good concordance for fibrosis staging.22,23

STAGING BASED ON DEMOGRAPHIC AND LABORATORY VARIABLES
Several scores based on patient characteristics and laboratory values have been developed for assessing liver fibrosis and have been specifically validated for HCV infection, NAFLD, or both. They can serve as inexpensive initial screening tests for the presence or absence of advanced fibrosis.

FIB-4 index for HCV, NAFLD
The FIB-4 index predicts the presence of advanced fibrosis using, as its name indicates, a combination of 4 factors in fibrosis: age, platelet count, and the levels of aspartate aminotransferase (AST) and alanine aminotransferase (ALT), according to the formula:

\[
\text{FIB-4 index} = \frac{(\text{age} \times \text{AST} [\text{U/L}])}{(\text{platelet count} [\times 10^9/\text{L}] \times \sqrt{\text{ALT} [\text{U/L}]}).}
\]

The index was derived from data from 832 patients co-infected with HCV and human immunodeficiency virus.24 The Ishak staging system10 for fibrosis on liver biopsy was used for confirmation, with stage 4 to 6 defined as advanced fibrosis. A cutoff value of more than 3.25 had a positive predictive value of 65% for advanced fibrosis, and to exclude advanced fibrosis, a cutoff value of less than 1.45 had a negative predictive value of 90%.

The FIB-4 index has since been validated in patients with HCV infection25 and NAFLD.26 In a subsequent study in 142 patients with NAFLD, the FIB-4 index was more accurate in diagnosing advanced fibrosis than the other noninvasive prediction models discussed below.27

NAFLD fibrosis score
The NAFLD fibrosis score, constructed and validated only in patients with biopsy-confirmed NAFLD, incorporates age, body mass index, presence of diabetes or prediabetes, albumin level, platelet count, and AST and ALT levels.

A group of 480 patients was used to construct the score, and 253 patients were used to validate it. Using the high cutoff value of 0.676, the presence of advanced fibrosis was diagnosed with a positive predictive value of 90% in the group used to construct the model (82% in the validation group). Using the low cutoff score of –1.455, advanced fibrosis could be excluded with a negative predictive value of 93% in the construction group and 88% in the validation group.28 A score between the cutoff values merits liver biopsy to determine fibrosis stage. The score is more accurate in patients with diabetes.29 When used by prima-
ry care physicians, the NAFLD fibrosis score is more cost-effective than transient elastogra-
phy and liver biopsy for accurately predicting advanced fibrosis.30

AST-to-platelet ratio index score for HCV, NAFLD
The AST-to-platelet ratio index (APRI) score was developed in 2003 using a cohort of 270
patients with HCV and liver biopsy as the standard. A cutoff value of less than or equal
to 0.5 had a negative predictive value of 86%
for the absence of significant fibrosis, while a
score of more than 1.5 detected the presence
of significant fibrosis with a positive predictive
value of 88%.31 The APRI score was subsequently
validated for NAFLD.27,32

FibroSure uses a patented formula
FibroSure (LabCorp; labcorp.com) uses a pat-
tented mathematical formula that takes into
account age, sex, and levels of gamma-glutamyl
transferase, total bilirubin, haptoglobin,
apolipoprotein-A, and alpha-2 macroglobulin
to assess fibrosis. Developed in 2001 for use in
patients with HCV infection, it was reported
to have a positive predictive value of greater
than 90% and a negative predictive value of
100% for clinically significant fibrosis, defined
as stage 2 to 4 based on the META VIR stag-
ing system in the prediction model.33 The use
of FibroSure in patients with HCV was sub-
sequently validated in various meta-analyses
and systematic reviews.34,35 It is less accurate
in patients with normal ALT levels.36
FibroSure also has good accuracy for pre-
dicting fibrosis stage in chronic liver disease
due to other causes, including NAFLD.37

The prediction models discussed above use
routine laboratory tests for chronic liver dis-
ease and thus are inexpensive. The high cost
of additional testing needed for FibroSure,
coupled with the risk of misdiagnosis, makes
its cost-effectiveness questionable.38

■ IMAGING TO PREDICT FIBROSIS STAGE

Conventional ultrasonography (with or with-
out vascular imaging) and computed tomog-
raphy can detect cirrhosis on the basis of cer-
tain imaging characteristics,39,40 including the
nodular contour of the liver, caudate lobe hy-
pertrophy, ascites, reversal of blood flow in the
portal vein, and splenomegaly. However, they
cannot detect fibrosis in its early stages.

The 3 methods discussed below provide
more accurate fibrosis staging by measuring
the velocity of shear waves sent across hepatic
tissue. Because shear-wave velocity increases
with liver stiffness, the fibrosis stage can be
estimated from this information.41

Transient elastography
Transient elastography uses a special ultra-
sound transducer. It is highly accurate for pre-
dicting advanced fibrosis for almost all causes
of chronic liver disease, including HCV infec-
tion42,43 and NAFLD.44 The cutoff values of
wave velocity to estimate fibrosis stage differ
by liver disease etiology.

Transient elastography should not be used
to evaluate fibrosis in patients with acute hepa-
titis, which transiently increases liver stiffness,
resulting in a falsely high fibrosis stage diagno-
sis.45 It is also not a good method for evaluating
fibrosis in patients with biliary obstruction or
extrahepatic venous congestion. Because liver
stiffness can increase after eating,46 the test
should be done under fasting conditions.

A significant limitation of transient elas-
tography has been its poor accuracy in patients
with obesity.47 This has been largely overcome
with the use of a more powerful (XL) probe but
is still a limitation for those with morbid obe-
sity.48 Because many patients with NAFLD are
obese, this limitation can be significant.

Transient elastography has gained populari-
ty for evaluating fibrosis in patients with chron-
ic liver disease for multiple reasons: it is cost-
effective and results are highly reproducible,
with low variation in results among different
observers and in individual observers.49 Com-
bined with a platelet count, it can also be used
to detect the development of clinically signifi-
cant portal hypertension in patients with cir-
rhosis, thus determining the need to screen for
esophageal varices using endoscopy.50 Screen-
ing endoscopy can be avoided in patients whose
liver stiffness remains below 20 kPa or whose
platelet count is above 150 × 10^9/L.

Acoustic radiation force imaging
Unlike transient elastography, which requires
a separate transducer probe to assess shear-
wave velocity, acoustic radiation force im-
aging uses the same transducer for both this
ASSESSING LIVER FIBROSIS

Transient elastography is cost-effective, and results are highly reproducible.

Magnetic resonance elastography
Magnetic resonance elastography uses a special transducer placed under the rib cage to transmit shear waves concurrently with magnetic resonance imaging. It has been tested in patients with HCV and NAFLD and has been found to have better diagnostic accuracy than transient elastography and acoustic radiation force imaging.\(^5\) Patients must be fasting for better diagnostic accuracy\(^5\) and must hold their breath while elastography is performed. The need for breath-holding and the high cost limit the use of this method for assessing fibrosis.

BOTTOM LINE FOR ASSESSING FIBROSIS

Although liver biopsy remains the gold standard for accurately determining fibrosis stage, noninvasive methods, especially imaging techniques, are fast evolving. Guidelines recommend using transient elastography to determine fibrosis stage noninvasively in patients with HCV,\(^6\) but a similar recommendation cannot be made for NAFLD with available data. For NAFLD, combined elastography and NAFLD fibrosis score are recommended to determine the need for a liver biopsy (Figure 2).\(^7\) Currently, we recommend using a combination of the scores discussed above and the imaging tests.
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SINGH AND COLLEAGUES


ADDRESS: Arthur J. McCullough, MD, Department of Gastroenterology and Hepatology, A30, Cleveland Clinic, 9500 Euclid Avenue, Cleveland, OH 44195; mcculla@ccf.org