Accurate assessment of the degree of liver fibrosis is important for estimating prognosis and deciding on an appropriate course of treatment for cases of chronic liver disease (CLD) with various etiologies. Because of the inherent limitations of liver biopsy, there is a great need for non-invasive and reliable tests that accurately estimate the degree of liver fibrosis. Ultrasound (US) elastography is considered a non-invasive, convenient, and precise technique to grade the degree of liver fibrosis by measuring liver stiffness. There are several commercial types of US elastography currently in use, namely, transient elastography, acoustic radiation force impulse imaging, supersonic shear-wave imaging, and real-time tissue elastography. Although the low reproducibility of measurements derived from operator-dependent performance remains a significant limitation of US elastography, this technique is nevertheless useful for diagnosing hepatic fibrosis in patients with CLD. Likewise, US elastography may also be used as a convenient surveillance method that can be performed by physicians at the patients’ bedside to enable the estimation of the prognosis of patients with fatal complications related to CLD in a non-invasive manner.

Keywords: Elasticity imaging techniques; Liver cirrhosis; Hypertension, portal; Ultrasonography
issue of intra- or inter-observer variability among pathologists in evaluating the grade of fibrosis is an additional limitation because the interpretation process is a subjective and semi-quantified method [4]. According to previous research on chronic hepatitis C, agreement among pathologists regarding the fibrosis grade is not excellent ($\kappa$, about 0.5) [5]. Although the rate of complications is very low and the risk has declined with the use of ultrasonographic guidance [6], liver biopsy is somewhat invasive and post-biopsy bleeding can be serious. With respect to non-invasive alternatives to liver biopsy, several serological or biochemical methods for the estimation of liver fibrosis have been validated primarily in patients with chronic hepatitis C, but still lack the ability to identify and classify the intermediate stages of fibrosis [7].

Introduced in 1991, elastography is another non-invasive technique for evaluating the elastic properties of soft tissue either quantitatively or qualitatively [8]. The elastography of the liver is theoretically not easy to determine compared with that of superficial organs because the liver is located deep and under the rib cage. Nevertheless, various techniques of ultrasound (US) elastography have been developed for repeatedly measuring hepatic fibrosis. From a technical standpoint, two types of US elastography for the measurement of liver stiffness are under development: shear wave-based elastography and real-time tissue elastography (Fig. 1, Table 1). This review addresses the principles and clinical usefulness of US elastography for the diffuse liver disease with an emphasis on shear wave-based elastography.

**Basic Principles of Elastography**

Elastography is a promising imaging technique because the elastic modulus of tissues measured by this technique provides the most broad-banded properties compared with other quantitative values measured by computed tomography (attenuation value), magnetic resonance (MR) imaging (T1 relaxation time), and conventional ultrasonography (bulk modulus). The order of magnitude of the elastic modulus is approximately five times larger than that for other imaging modalities [9], meaning that the use of the elastic modulus can maximize the discrimination between different tissues or between normal tissue and lesions. The elastic modulus is defined as the slope of the stress-strain curve during elastic deformation. Therefore, a stiffer object has a higher elastic modulus. There are various approaches to elastic imaging, all of which consist of three basic steps: excitation (stress) application, tissue response (strain) measurement, and mechanical parameters estimation [9].

**Excitation Application**

In its most basic form, shear wave-based elastography applies a perpendicular stress force on the target organ to induce “shear” on the tissue (Fig. 2). By definition, shear is the change of shape (displacement)—without a change in volume—produced by a pair of forces acting in opposite directions. At this point, transversely propagating waves with a very low velocity develop in the tissue,

![Shear wave-based elastography](image1)

![Real-time tissue elastography](image2)

**Fig. 1. Classification of ultrasound elastography of the liver. ARFI, acoustic radiation force impulse.**

| Technique | Transient elastography | ARFI imaging | Supersonic shear-wave imaging | Real-time tissue elastography |
|-----------|------------------------|--------------|-------------------------------|------------------------------|
| Type of force | Dynamic | Dynamic | Dynamic | Quasi-static |
| Applied force | Mechanical impulse | US-induced radiation force impulse | US-induced radiation force impulse | Intrinsic (heartbeat) |
| Measurement of strain | Single measure, beam-line average | Single image within a box | Image within a color box | Full area image |
| Estimated parameter | Elastic modulus converted from shear wave velocity (kPa) | Velocity of shear wave (cm/sec) | Elastic modulus converted from shear wave velocity (kPa) | Strain ratio |
| Qualitative or quantitative | Quantitative | Qualitative/quantitative | Quantitative | Qualitative |
| Clinical evidence | Very much | Much | Little | Scanty |

ARFI, acoustic radiation force impulse.

Modified from Bamber et al. [54] with permission from Elsevier.
which are called shear waves. In the case of transient elastography, a mechanical push is used for excitation application, which produces transient shear waves in the target tissue. This type of excitation application is classified as a dynamic elastography technique with an external source. In addition, dynamic techniques induce vibrations and comprise the basic method for shear wave-based US elastography and MR elastography. In particular, rather than a mechanical push with transient elastography, focused US beams from a US transducer make shear waves through the absorption of acoustic energy; acoustic radiation force impulse (ARFI) imaging and supersonic shear-wave imaging (SSI) belong to this category [10]. With respect to shear wave elastography (Supersonic Imagine, Aix-en-Provence, France), a type of SSI, multiple acoustic radiation forces are successively focused at different depths in a tissue to generate a strong shear wave to propagate along the tissue at a safe level of acoustic power, which is then coherently summed in a Mach cone shape and improves the propagation distance [11]. On the other hand, real-time tissue elastography methods are derived from the static elastography technique used for the measurement of breast tissue elasticity, and employ the quasi-static or intrinsic stress derived from heartbeats [12,13].

**Tissue Response Measurement**

Measurement of tissue response is the most critical component of elastography. The basic measurement method consists of a comparison of successively obtained images and a reference image (Fig. 2). Through either a mechanical push or acoustic radiation force, the A-axis (direction of force=depth direction) displacement of the target tissue occurs, and the shear waves are generated simultaneously. These are very slow (1−10 m/sec) compared with an US beam and travel perpendicular to the direction of the stress force. To detect a shear wave, two methods using US have been introduced. Transient elastography causes a single transient shear wave to propagate along the A-axis direction by using an M-mode US technique and calculates Young modulus of the tissue by using this information [14]. Another method is the Doppler technique, in which radiofrequency (RF) images including the information of the propagating shear waves are measured using the echo of the transmitted US beams at a very high frame rate, which can be used to generate a tissue displacement map [8,15,16]. Using the tissue displacement maps obtained during the period of shear wave propagation (i.e., less than 14 ms), it is possible to calculate the velocity of a shear wave by analyzing the movement of the peak of the shear wave. In this way, the elastic modulus can be calculated by $E=\frac{3}{2} \rho V_s^2$ where $\rho$ denotes the density of the tissue and $V_s$ represents the velocity of the shear wave.

**Mechanical Parameter Estimation**

Both qualitative and quantitative methods are used to perform mechanical parameter estimation (Fig. 2). Liver stiffness is usually

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**Fig. 2.** Diagram depicting the process of shear wave-based ultrasound elastography. Shear wave-based elastography applies a perpendicular stress force to a target organ in order to induce shear on the tissue. The information on the propagating shear wave including the velocity of the shear wave could be measured by obtaining radiofrequency images with a high frame rate, which can be used to generate a tissue displacement map. Then, the elastic property for quantitative estimation is calculated by the propagating velocity of the shear wave. ARFI, acoustic radiation force impulse.
measured quantitatively. The elastic properties for quantitative estimation are expressed as Young modulus (E) or the shear modulus (μ). For most soft tissues, Young modulus and the shear modulus are related by a simple scale factor of 3: i.e., E = 3 μ [9]. Transient elastography and SSI display tissue stiffness as the elastic modulus (kPa), while ARFI imaging shows it by shear wave velocity (cm/sec). In the case of real-time elastography, tissue elasticity cannot be measured directly from the reflected US echo. Relative tissue elasticity is thus calculated and displayed as a color overlay of the conventional B-mode image, and the strain ratio between two different points can be obtained instead of elastic modulus or shear wave velocity.

**Methods of Shear Wave-based Elastography**

**Transient Elastography**

Transient elastography was the first commercialized elastography method developed to noninvasively assess the stiffness of deep soft tissues such as the liver. Transient elastography consists of two parts: a mechanical vibrator and a single-channel US transducer [14]. The mechanical vibrator generates a low-frequency wave at 50–500 Hz to generate shear stress in the target tissue at a length of 4 cm, and the velocity of the shear wave can then be measured using an US signal (Fig. 3). The most superior advantage of transient elastography is that it has been extensively validated by numerous investigations targeted at patients with CLD, and the results of transient elastography are generally accepted to be well correlated with different stages of liver fibrosis. The validity of liver stiffness measurements are determined by the success rate and the interquartile range divided by the median (IQR/M) in cases with more than 10 valid measurements. Here, the success rate is the ratio of the number of valid measurements to the total number of measurements and should be greater than 60%; IQR/M should be lower than 30% [17]. Despite these advantages, there are several disadvantages of transient elastography. They are as follows: first, transient elastography does not provide a B-mode image, which is essential for accurate targeting. Second, transient elastography is not performed for the patient with ascites. In addition, transient elastography exhibits a relatively high measurement failure rate of 4.5%–6%. Major factors related to this failure rate include a body mass index greater than 28, age over 50 years, non-alcoholic steatohepatitis, diabetes, and a γ-glutamyl-transpeptidase level higher than 57 IU/L [18].

**Acoustic Radiation Force Impulse Imaging and Shear Wave Imaging**

Both ARFI imaging and SSI use focused high-intensity, short-duration acoustic pulses instead of the mechanical vibration of transient elastography in order to produce shear waves in the target tissue [19]. ARFI imaging generates shear waves by a single pushing beam, while shear wave propagation is monitored using conventional pulse-echo US at multiple off-axis lateral locations. By collecting displacement through time information at multiple lateral locations separated by a known distance from the excitation source, the speed of the propagating shear waves can be estimated (Fig. 4). The region of interest (ROI) of ARFI imaging is relatively small (i.e.,...
10 mm × 5 mm) as compared to that of SSI.

SSI is a new shear wave-based US elastography technique. SSI generates push beams at multiple axial depths to create a near-supersonic plane wave shear and transmits the unfocused beam (plane wave) to monitor the shear wave propagation throughout the ROI (Fig. 5). The ROI of SSI is fan-shaped and larger than other modalities (up to 50 mm × 50 mm) [20,21]. A remarkable feature of SSI is that it can show viscoelastic properties in all areas in an ROI with a color look-up table and thus, is expected to overcome the limitations of transient elastography by which liver stiffness cannot be measured accurately in patients with severe obesity, thick subcutaneous fat, and ascites [22]. Moreover, the same technique can be used to display a grayscale US image on the background of the elastogram, so it is more reliable and familiar to a physician who uses conventional ultrasonography. However, for as wide a use of SSI for the diagnosis of liver fibrosis as that of transient elastography, further clinical experiences and evidence are needed.

**Real-Time Tissue Elastography**

As mentioned above, real-time elastography is a method derived from the static elastography technique used for the measurement of breast tissue elasticity. Real-time tissue elastography uses a grayscale US machine, incorporating elastography into the conventional US scanner. This approach uses a quasi-static method.
consisting of an excitation application in which the examiner has to compress and relax the body by a transducer or utilize intrinsic stress derived from the heartbeat [12,13]. Simultaneously, echo signals are captured in real time, and the relative stiffness (elastic ratio) of the liver can be calculated along with a real-time color map. Like SSI, real-time tissue elastography can display tissue elasticity images and conventional grayscale US images at the same time but is unable to calculate the elastic modulus (Fig. 6).

**Clinical Applications of US Elastography**

**Hepatic Fibrosis**

In terms of the diagnostic performance of US elastography for measuring liver stiffness, there have been many clinical studies regarding the diagnosis of hepatic fibrosis using transient elastography [23,24]. According to a recent meta-analysis aimed at chronic hepatitis C patients, the pooled estimate of the cut-off value for significant fibrosis (≥F2 on the META VIR score system) was 7.11 kPa with a sensitivity of 72% and specificity of 82%. In the case of cirrhosis (F4), the results showed a cut-off of 15.08 kPa with a sensitivity of 84% and specificity of 95%. Another meta-analysis of 40 eligible studies showed that the summary sensitivity and specificity were 78% and 80% for significant fibrosis, and 83% and 90% for cirrhosis, respectively (Table 2). In addition, this meta-analysis suggested that transient elastography could be used as a good screening test for cirrhosis, but not for accurately diagnosing fibrotic stages other than cirrhosis because no optimal cut-offs of liver stiffness for individual fibrosis stages have been validated. There are relatively fewer clinical studies of ARFI imaging and SSI compared with transient elastography studies, although some studies showed that the performance and reliability of ARFI imaging and SSI are comparable to those of transient elastography [19,20,25]. According to a recent meta-analysis study for ARFI imaging, the mean diagnostic accuracy of ARFI expressed as areas under the receiver operating characteristic curves (AUROCs) was 0.87 for ≥F2, 0.93 for ≥F3, and 0.93 for F4 [26]. In the case of real-time elastography, the elastic ratio (ratio of the value in the intrahepatic venous small vessels to the value in the parenchyma) was calculated instead of liver stiffness, and the diagnostic performance of this technique was superior to that of non-invasive biochemical markers including aspartate aminotransferase to the platelet ratio index (APRI) and the Forns Index [12].

**Table 2. Summary sensitivity, specificity, and diagnostic odds ratio (DOR) to diagnose F2 and F4 fibrosis by using transient elastography extracted from a meta-analysis**

| Causes           | Stage | Studies | Cut-off liver stiffness | Sensitivity (%) | Specificity (%) | DOR  |
|------------------|-------|---------|-------------------------|-----------------|-----------------|------|
| Chronic hepatitis C | ≥F2   | 14      | 7.6 (5.1–10.1)          | 78 (71–84)      | 80 (71–86)      | 13.9 |
|                  | F4    | 11      | 15.3 (11.9–26.5)        | 83 (77–88)      | 90 (87–93)      | 46.5 |
| Chronic hepatitis B | ≥F2   | 4       | 7.0 (6.9–7.2)           | 84 (67–93)      | 78 (68–85)      | 17.9 |
|                  | F4    | 6       | 11.3 (9.0–13.4)         | 80 (61–91)      | 89 (82–94)      | 34.3 |

Modified from Tschochatzis et al. [24]
The diagnostic performance of US elastography in patients with an HBV infection has not yet been well addressed. Marcellin et al. [27] first reported the diagnostic accuracy of transient elastography in the patients with chronic hepatitis B. Although the clinical evidence for fibrosis staging in patients with chronic hepatitis B is relatively weak and the overall diagnostic performance is slightly worse than for chronic hepatitis C patients, the diagnostic accuracy of transient elastography is acceptable [28]. In a recent meta-analysis of the performance of transient elastography for the staging of liver fibrosis in chronic hepatitis B, the mean AUROCs for the diagnosis of significant fibrosis, severe fibrosis, and liver cirrhosis were 0.859, 0.887, and 0.929, respectively, as determined from 18 studies comprising 2,772 patients [29]. Compared with HCV patients, the cut-off value for cirrhosis in HBV patients tends to be lower because the liver fibrosis area in macronodular cirrhosis is relatively smaller. In contrast, alanine aminotransferase flares can

**Fig. 7. Acute exacerbation of hepatitis B virus-induced hepatitis.**
Grayscale ultrasonography (A) shows mild parenchymal coarseness of the liver; the liver stiffness is 10.3 kPa according to the shear wave elastography (B). Six months later, the patients recovered from liver function deterioration. The grayscale ultrasonographic feature (C) is similar to the previous study, but the liver stiffness decreases to 6 kPa (D).
result in overestimating the fibrosis grade, and thus, the increased liver stiffness should be interpreted carefully in patients with acute exacerbation of chronic HBV hepatitis [30] (Fig. 7).

Non-Alcoholic Fatty Liver Disease
Non-alcoholic fatty liver disease (NAFLD) actually represents an emerging disease of great clinical interest because of the increasing incidence of metabolic diseases and obesity in recent decades. The disease spectrum of NAFLD is very wide, ranging from simple fatty liver to non-alcoholic steatohepatitis, and liver fibrosis can develop and progress to liver cirrhosis [31]. Although transient elastography is difficult to perform in cases of obesity because subcutaneous fatty tissue attenuates the pushing pulse, its role in causing NAFLD has recently been highlighted owing to the development of a new technique utilizing a vibration-controlled transient elastography device, which allows the calculation of the new controlled attenuation parameter (CAP). CAP is known to be useful for the non-invasive and accurate estimation of liver steatosis [32]. In addition, a new XL probe designed to measure shear waves at deeper positions by using a lower central US frequency (2.5 MHz) can be applied, thereby allowing more reliable results to be obtained compared with conventional M probes [33]. According to a diagnostic algorithm introduced by a recent review article [34], transient elastography could be useful to stratify patients at indeterminate risk; if liver stiffness was lower than 7.9 kPa, advanced fibrosis could be excluded with a 97% negative predictive value. On the other hand, if liver stiffness was higher than 9.6 kPa, the diagnosis of advanced fibrosis could be made with a 72% positive predictive value [35]. Thus, only the remaining patients (7.9–9.6 kPa) require liver biopsy for diagnosis, and the number of NAFLD patients with the indication of liver biopsy can be decreased, leading to a reduced incidence of biopsy-induced complications [34].

Follow-up after Liver Transplantation
After liver transplantation, US elastography is also useful to diagnose liver fibrosis caused by relapsed chronic hepatitis and the acute rejection of the liver graft. A systematic review of studies comparing US elastography to liver biopsy for the detection of liver fibrosis by a recurrent HCV infection stated that the diagnostic accuracy for significant fibrosis (F2) using transient elastography was generally good, with a sensitivity and a specificity of 83%. Further, with respect to liver cirrhosis, the sensitivity and specificity were improved to 98% and 84%, respectively [36]. Liver stiffness may also be increased by acute cellular rejection following liver transplantation; however, it is important to keep in mind that liver stiffness can increase in transplanted livers without evidence of rejection since it may undergo ischemic or reperfusion injury within 4 weeks from transplantation, which can in turn result in transient hepatocellular ballooning and hepatocanalicular cholestasis with inflammation that may recover within 2–3 weeks without specific treatment. Therefore, US elastography may be useful to detect rejection at follow-up more than 4 weeks after transplantation [37].

Portal Hypertension
Estimation of the severity of portal hypertension in patients with liver cirrhosis is another major use of liver stiffness measurements. Increased portal pressure is the major factor driving the clinical course of cirrhosis. Measurement of the hepatic venous pressure gradient (HVPG) following hepatic venous catheterization was used as a surrogate marker of portal hypertensive stigmata. Recently, there were some investigations concerning the feasibility of the noninvasive measurement of liver stiffness to estimate severe portal hypertension [38,39]. Looking at the results of these studies, we concluded that liver stiffness could be closely correlated with both HVPG and the presence of complications related with portal hypertension (Fig. 8).

Roles for Longitudinal Surveillance
To date, the majority of elastography studies have focused on evaluating the cross-sectional performance with respect to the histological fibrosis grade or HVPG. However, an important but undervalued use of elastography is the ability to repeatedly measure liver stiffness. The roles of elastography as longitudinal perspectives with respect to the prediction of the long-term prognosis of the disease and monitoring of clinical courses with or without treatment are well known. In particular, these approaches can be used to non-invasively estimate the prognosis of the patients with fatal complications related to CLD, such as variceal bleeding and decompensation.

A longitudinal follow-up of elastography has been proposed as a way to establish the tailored management strategies by providing more detailed prognostic information [40]. For example, the concept of cirrhosis has recently changed from dynamic to bidirectional. In other words, cirrhosis patients may recover if antiviral therapy can be applied properly. At this time, the ideal approach to assess histological outcomes during treatment is serial liver biopsy; however, this is not possible in most cases. Instead, the measurement of liver stiffness by elastography is very useful for monitoring the changes in liver fibrosis during the antiviral treatment [41,42]. In terms of portal hypertension, elastography may also be used to predict the development of variceal bleeding by using a hybrid parameter, the liver stiffness-spleen diameter to platelet ratio score (LSPS) defined as the product of liver stiffness and the maximum spleen diameter divided by the platelet count [43]. According to risk stratification
based on LSPS, a different prophylactic treatment for the prevention of variceal bleeding should be considered for patients with an LSPS value higher than 6.5 points [44].

**US Elastography: Weaknesses and Strengths**

The most significant challenge facing US elastography is the issue of measurement reproducibility. A number of studies concerning this issue have been published; however, many investigators have brought up questions about this issue due to the inherent limitations of US such as the operator-dependent performance. Transient elastography is a highly reproducible and user-friendly technique [45], and liver stiffness measurement by transient elastography does not require a learning curve: even a novice can obtain a reliable result after a single training session [46]. However, because liver stiffness measurements can be influenced significantly by steatosis, obesity, lower degrees of hepatic fibrosis [45], necroinflammation of hepatocytes [47], cholestasis [48], elevated central venous pressure [49], and even postprandial conditions [50], it should be carefully applied when used as an alternative measurement of liver stiffness instead of liver biopsy.

In the case of ARFI, the overall reproducibility is also not bad, having an intraclass correlation coefficient (ICC) value for the interrater observation of 0.81 and an ICC for the intrarater observation of 0.90. However, gender (women), high body mass index, ascites, and lower degree of liver disease (noncirrhotic patients) are considered factors that impede the reproducibility of ARFI [51]. In the case of SSI, the inter- and intraobserver agreements

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**Fig. 8. Shear wave elastography of a patient with portal hypertension.**

On grayscale ultrasonography, the cirrhotic change of the liver and the perihepatic ascites (arrows) are noted (A). The measured liver stiffness is 42.6 kPa (B), which is about 8 times the normal value. The hepatic venous pressure gradient was 26.3 mmHg, which indicates severe portal hypertension (C).
have ICC values of 0.88 and 0.94, respectively, which are similar to the results of ARFI imaging [52].

Despite the issues described above, US elastography has many advantages in clinical fields. The most important aspect is convenience, as is the case with most ultrasonography examination techniques. Indeed, US elastography is fast, easy to use, and portable, so much so that it can be performed at the patient’s bedside. Likewise, because it does not use ionizing radiation, US elastography is relatively safe, even in patients who repeatedly undergo the procedure. US elastography is also less expensive than MR elastography [53]. Going forward, the most important strength of US elastography is the availability of a large amount of accumulated clinical data that have demonstrated its clinical usefulness, although most of these data are related to transient elastography.

Conclusions

Measurement of liver stiffness using various technical developments is evolving to overcome its limitations. Recently, the European Federation of Societies for Ultrasound in Medicine and Biology (EFSUMB) published an informative guideline for the use of US elastography [54,55] that deals with the relevant technology and clinical applications. Along with the basic principles for use, these guidelines include the practical advantages and disadvantages of US elastography as well as recommendations for the examination of various body parts. According to these guidelines, US elastography is useful to assess the severity of liver fibrosis in patients with diffuse liver disease and particularly to distinguish patients with nil to mild fibrosis from those with significant fibrosis, although some of the newer techniques must be validated through clinical studies. At present, however, US elastography for the differentiation of focal hepatic lesions is not recommended.

In conclusion, US elastography is useful for diagnosing hepatic fibrosis in patients with CLD and may be used as a convenient and non-invasive surveillance method to estimate the prognosis of patients with fatal complications related to CLD. Accordingly, the development of a standardized method for liver stiffness measurement and technical improvements should be a priority for the clinical application of US elastography. Together, these efforts will significantly enhance the clinical implications of US elastography.

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Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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