The feasibility and reliability of transient elastography using Fibroscan®: A practice audit of 2335 examinations

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BACKGROUND: Liver stiffness measurement (LSM) using transient elastography is widely used in the management of patients with chronic liver disease.

OBJECTIVES: To examine the feasibility and reliability of LSM, and to identify patient and operator characteristics predictive of poorly reliable results.

METHODS: The present retrospective study investigated the frequency and determinants of poorly reliable LSM (interquartile range [IQR]/median LSM [IQR/M] >30%) with median liver stiffness ≥7.1 kPa using the FibroScan (Echosens, France) over a three-year period. Two experienced operators performed all LSMs. Multiple logistic regression analyses examined potential predictors of poorly reliable LSMs including age, sex, liver disease, the operator, operator experience (<500 versus ≥500 scans), FibroScan probe (M versus XL), comorbidities and liver stiffness. In a subset of patients, medical records were reviewed to identify obesity (body mass index ≥30 kg/m²).

RESULTS: Between July 2008 and June 2011, 2335 patients with liver disease underwent LSM (86% using the M probe). LSM failure (no valid measurements) occurred in 1.6% (n=37) and was more common using the XL than the M probe (3.4% versus 1.3%; P=0.01). Excluding LSM failures, poorly reliable LSMs were observed in 4.9% (n=113) of patients. Independent predictors of poorly reliable LSM included older age (OR 1.03 [95% CI 1.01 to 1.05]), chronic pulmonary disease (OR 1.58 [95% CI 1.05 to 2.37]), coagulopathy (OR 2.22 [95% CI 1.31 to 3.76]) and higher liver stiffness (OR per kPa 1.03 [95% CI 1.02 to 1.05]), including presumed cirrhosis (stiffness ≥12.5 kPa; OR 5.24 [95% CI 3.49 to 7.89]). Sex, diabetes, the underlying liver disease and FibroScan probe were not significant. Although reliability varied according to operator (P=0.0005), operator experience was not significant. In a subanalysis including 434 patients with body mass index data, obesity influenced the rate of poorly reliable results (OR 2.93 [95% CI 0.95 to 9.03]; P=0.06).

CONCLUSIONS: FibroScan failure and poorly reliable LSM are uncommon. The most important determinants of poorly reliable results are older age, obesity, higher liver stiffness and the operator, the latter emphasizing the need for adequate training.

Key Words: Biopsy; Diagnostic test; Fibrosis; Hepatitis; Stiffness

The evaluation of liver fibrosis is a vital component of the management of patients with chronic liver disease, both for guiding therapy and estimating prognosis. Traditionally, liver biopsy has been used for this purpose (1); however, biopsy is limited by invasiveness, cost, variability in histological interpretation, sampling error (2,3) and difficulty of repetition for monitoring changes in fibrosis over time. Moreover, potentially serious, albeit uncommon, complications including hemorrhage and death may occur (4). Due to these limitations, numerous, noninvasive means for staging liver fibrosis have been developed, including transient elastography (TE) (5-8) using FibroScan (Echosens, France). TE is an ultrasound-based tool for measuring liver stiffness as a surrogate of liver fibrosis. It is widely used due to its high

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accuracy for the diagnosis of advanced fibrosis, thus reducing the use of biopsy for the initial evaluation of patients with liver disease (9,10). In addition, studies have demonstrated that liver stiffness is responsive to changes in fibrosis over time (eg, attributable to treatment or disease progression), thus supporting its use for serial patient monitoring. An emerging body of literature also supports the prognostic significance of liver stiffness, suggesting that TE may be useful for guiding management strategies including the intensity of follow-up (11-17). For example, because the incidence of variceal hemorrhage is low in patients with liver stiffness <21 kPa, restriction of endoscopic screening to cirrhotic patients with higher liver stiffness has been advocated (14).

Because numerous clinical decisions are made based on the results of liver stiffness measurement (LSM) using TE, it is vital to understand the reliability of this tool and factors that influence its accuracy. Traditionally, TE examinations with <10 valid measurements, a success rate <60% and/or a ratio of the interquartile range (IQR) of liver stiffness to the median value (IQR/M) >30% have been classified as unreliable (18). According to this definition, approximately 15% of LSMs are unreliable, with higher rates among older patients, women and patients with diabetes, hypertension, and higher body mass index (BMI) and waist circumference. Limited operator experience (<500 versus ≥500 examinations) may also influence reliability (18). However, this definition has been criticized because no studies have ever demonstrated that reliable results are more accurate than unreliable results. As such, Boursier et al (19) recently proposed a new definition of poorly reliable TE results (IQR/M >30% and median liver stiffness ≥7.1 kPa). In this study, poorly reliable LSMs – which occurred in 9% of patients – were less accurate than reliable LSMs according to multiple indicators of diagnostic test performance (19). The factors that influence the rate of poorly reliable results according to this definition have not been explored.

Accordingly, the objective of the present study was to examine the feasibility and reliability of TE using this revised definition of reliability in a large cohort of patients with various liver diseases and severities to reflect routine clinical practice. We aimed to identify patient and operator characteristics associated with poorly reliable results to better inform our interpretation of LSMs using TE.

METHODS

Study population
In the present retrospective study, 2437 consecutive patients who underwent LSM using TE (FibroScan) at the University of Calgary Liver Unit (UCLU; Calgary, Alberta), between July 2008 and June 2011 were identified. The UCLU is the major referral centre for patients with liver disease who reside in Southern Alberta and serves a catchment population of approximately 1.5 million. LSM using TE is performed routinely in all patients who attend the UCLU without overt evidence of hepatic decompensation. In patients with multiple examinations, only the first LSM was considered to eliminate selection bias (ie, repeated examinations on patients who are easier to scan). For our analysis of the prevalence of FibroScan failure and reliability (see definitions below), the entire study cohort was examined. To identify predictors of poorly reliable LSM, only patients for whom linkage with Alberta administrative databases was possible (n=2335) were included to permit identification of the patients’ underlying liver diseases and comorbidities. Therefore, for these analyses, 36 nonresidents of Alberta, 32 patients with invalid provincial health numbers and 34 patients for whom linkage with the administrative data were unsuccessful were excluded. These patients had demographic characteristics and liver stiffness results similar to the remainder (data not shown). The Conjoint Health Research Ethics Board at the University of Calgary approved the study protocol.

LSM
Two experienced operators (OP1 and OP2) performed all FibroScan examinations as per the manufacturer’s recommendations. OP2 was employed after OP1 and performed her first LSM in September 2010. Between July 2008 and July 2009, the FibroScan M probe was used in all patients; thereafter, the FibroScan XL probe was used in obese patients (BMI ≥30 kg/m²). Briefly, with the patient lying in the dorsal decubitus position and the right arm in maximal abduction, the tip of the FibroScan transducer probe was placed on the skin between the ribs over the right lobe of the liver. Assisted by a sonographic image, a portion of the liver at least 6 cm thick and free of large vascular structures was identified, and an attempt was made to collect at least 10 valid LSMs. The median liver stiffness value (in kPa) was considered to be representative of the elastic modulus of the liver. As an indicator of LSM variability, the IQR/M was calculated.

Administrative data sources
The present study used three Alberta administrative databases to identify the underlying liver disease etiologies and comorbidities of study participants via linkage using their unique personal health number. These databases have been used to examine the epidemiology (20-22), outcomes (22,23) and coding accuracy (20,24-28) of a variety of medical conditions.

1. Physician Claims Database. This database includes claims submitted for payment by Alberta physicians for services provided to registrants of the Alberta Health Care Insurance Plan, a universal plan that covers >99% of Alberta residents (29). Each record in the database includes the service provided, the date and up to three diagnosis fields. The database was queried from April 1, 2001 to March 31, 2011.

2. Inpatient Discharge Abstract Database. This database contains diagnosis, procedure and mortality information on all discharges from hospitals within Alberta. These data are routinely transmitted to the Canadian Institute for Health Information for aggregation with nationwide hospitalization data (29). Chart validation studies have shown rates of agreement exceeding 95% for demographic data and 75% to 96% for most responsible diagnosis codes (30). The database was queried from January 1, 1991 to January 31, 2012.

3. National Ambulatory Care Reporting System (NACRS)/Ambulatory Care Classification System (ACCS) Database. This database contains information on facility-based ambulatory care including clinic and emergency department visits, same-day surgery, day procedures and rehabilitation services (29). The database was queried from July 1, 1996 to December 31, 2011.

Outcomes and predictor variables
The primary outcome variable was poorly reliable LSM, as defined by Boursier et al (19). Specifically, poorly reliable LSMs had an IQR/M >30% and median liver stiffness ≥7.1 kPa. Very reliable LSMs had an IQR/M ≤10%, whereas reliable LSMs had an IQR/M >10% and ≤30% (regardless of liver stiffness) or an IQR/M >30% with median liver stiffness <7.1 kPa. When compared with liver biopsy as the reference standard for staging fibrosis, poorly reliable LSMs are less accurate (ie, have lower areas under the ROC curves [AUROCs] and rates of accurate patient classification for fibrosis) than very reliable and reliable scans (19). As a secondary outcome measure, LSM failure, defined as no valid measurements after at least 10 attempts, was examined. The primary predictor variables included age, sex, the underlying liver disease, median liver stiffness (examined as a continuous variable and categorized as presumed cirrhosis [liver stiffness ≥12.5 kPa]) (31), comorbidities, the FibroScan operator (OP1 versus OP2), operator experience (first 499 scans versus ≥500 scans) (18) FibroScan probe (M versus XL) and the year of the LSM (2008 to 2009 versus 2010 to 2011). The hepatic diagnosis was categorized according to a hierarchy as follows: hepatitis B virus (HBV), hepatitis C virus, autoimmune (including primary biliary cirrhosis, primary sclerosing cholangitis and autoimmune hepatitis), hereditary hemochromatosis, alcoholic liver disease, nonalcoholic fatty liver disease and other (see Appendix I for list of relevant International Classification of Diseases, Ninth Revision,
TABLE 1
Characteristics of the study cohort according to the reliability of liver stiffness measurement

| Variable                          | Entire cohort* (n=2335) | Very reliable (n=659) | FibroScan (n=1526) | Poorly reliable (n=113) | OR (95% CI) for poorly reliable FibroScan |
|----------------------------------|-------------------------|-----------------------|--------------------|--------------------------|------------------------------------------|
| Male sex                         |                         |                       |                    |                          |                                          |
| Age, years, median (IQR)         | 50 (40–58)              | 49 (40–57)            | 50 (40–58)         | 53 (48–61)               | 1.04 (1.02–1.05)                          |
| Body mass index¹, kg/m², median (IQR) | 25.5 (23–30)           | 25 (22–28)           | 25.5 (23–30)       | 32.6 (25–41)             | 1.06 (1.00–1.11)                          |
| <30                              | 75 (332)                | 86 (107)             | 73 (211)           | 48 (10)                  | Reference                                 |
| ≥30                              | 25 (112)                | 14 (18)              | 27 (77)            | 53 (11)                  | 3.68 (1.52–8.94)                          |
| Hepatic diagnosis                |                         |                       |                    |                          |                                          |
| Hepatitis B                      | 27 (638)                | 31 (201)             | 27 (413)           | 18 (20)                  | 0.55 (0.34–0.90)                          |
| Hepatitis C                      | 36 (840)                | 36 (234)             | 35 (541)           | 44 (50)                  | 1.44 (0.99–2.11)                          |
| Autoimmune                       | 5.5 (129)               | 4.3 (28)             | 6.0 (92)           | 7.1 (8)                  | 1.31 (0.62–2.75)                          |
| Hemoschromatosis                 | 3.0 (69)                | 3.2 (21)             | 2.8 (43)           | 2.7 (3)                  | 0.90 (0.28–2.92)                          |
| Alcohol                          | 3.3 (78)                | 3.6 (24)             | 2.8 (42)           | 6.2 (7)                  | 2.12 (0.95–4.73)                          |
| Nonalcoholic fatty liver disease | 6.8 (158)               | 7.9 (52)             | 6.0 (92)           | 8.9 (10)                 | 1.38 (0.70–2.69)                          |
| Other/unknown                    | 18 (423)                | 15 (99)              | 20 (303)           | 13 (15)                  | 0.68 (0.39–1.18)                          |
| Comorbidities                     |                         |                       |                    |                          |                                          |
| Congestive heart failure         | 4.2 (97)                | 3.2 (21)             | 4.0 (61)           | 8.9 (10)                 | 2.49 (1.25–4.94)                          |
| Cardiac arrhythmia               | 6.5 (152)               | 5.6 (37)             | 6.1 (93)           | 12.4 (14)                | 2.24 (1.24–4.02)                          |
| Hypertension                     | 34 (803)                | 33 (218)             | 33 (507)           | 50 (56)                  | 1.98 (1.35–2.89)                          |
| Paralysis                        | 0.8 (18)                | 0.3 (2)              | 0.8 (12)           | 2.7 (3)                  | 4.23 (1.20–14.9)                          |
| Chronic lung disease             | 30 (699)                | 29 (193)             | 29 (445)           | 40 (45)                  | 1.60 (1.09–2.36)                          |
| Diabetes mellitus                | 12 (274)                | 8.4 (62)             | 11 (172)           | 23 (26)                  | 2.49 (1.58–3.94)                          |
| Peptic ulcer                     | 3.0 (69)                | 3.3 (22)             | 2.4 (37)           | 6.2 (7)                  | 2.38 (1.06–5.34)                          |
| Coagulopathy                     | 7.7 (179)               | 8.0 (53)             | 6.6 (100)          | 19 (21)                  | 3.03 (1.84–5.01)                          |
| Fluid/electrolyte disorders      | 15 (340)                | 14 (94)              | 14 (210)           | 23 (26)                  | 1.85 (1.17–2.91)                          |
| Alcohol abuse                    | 13 (306)                | 14 (90)              | 12 (181)           | 22 (25)                  | 2.01 (1.26–3.18)                          |
| Liver stiffness, kPa, median (IQR) | 6.3 (4.7–10.1)         | 6.0 (4.6–8.8)        | 6.1 (4.6–9.6)      | 13.3 (9.0–27.0)          | 1.04 (1.03–1.05)                          |
| Cirrhosis (≥12.5 kPa)            | 19 (430)                | 17 (109)             | 17 (257)           | 57 (64)                  | 6.49 (4.40–9.57)                          |
| M probe (versus XL probe)        | 86 (2009)               | 88 (562)             | 86 (1313)          | 78 (88)                  | 0.54 (0.34–0.85)                          |
| Operator 2 (versus operator 1)   | 53 (1235)               | 53 (361)             | 54 (823)           | 35 (39)                  | 0.45 (0.30–0.66)                          |
| Operator experience              |                         |                       |                    |                          |                                          |
| 1–499 scans                      | 35 (816)                | 38 (250)             | 33 (497)           | 37 (42)                  | Reference                                 |
| ≥500 scans                       | 65 (1519)               | 62 (409)             | 67 (1029)          | 62 (71)                  | 0.88 (0.59–1.30)                          |
| Year of FibroScan                |                         |                       |                    |                          |                                          |
| 2008–2009                        | 24 (550)                | 23 (154)             | 23 (334)           | 38 (43)                  | Reference                                 |
| 2010–2011                        | 76 (1785)               | 77 (505)             | 78 (1192)          | 62 (70)                  | 0.47 (0.32–0.69)                          |

Data presented as % (n) unless otherwise indicated. Bolded values indicate statistical significance (ie, P<0.05). *Entire cohort includes 37 FibroScan (Echosens, France) failures; †Body mass index data from medical record review available in 444 patients; ‡Only comorbidities with statistically significant associations with poorly reliable liver stiffness measurement are included. IQR Interquartile range

Clinical Modification [ICD-9-CM] [32] and ICD-10 [33] codes). For example, according to this approach, a patient with any record of a diagnosis code for HBV with or without other hepatic diagnoses would be categorized as having HBV. Comorbid conditions occurring before the FibroScan examination were defined using the Elixhauser list of 30 comorbidities (34), a well-validated algorithm for predicting outcomes in patients with hepatic (35) and nonhepatic disorders (34,36). Liver diseases were excluded from this algorithm. Also examined were the type of FibroScan probe because use of the XL probe could be considered a surrogate marker for obesity, which is not available in the administrative databases. In a subset of patients, medical records were reviewed to extract the BMI on the day of the FibroScan examination.

Statistical analyses
Between-group comparisons were made using Fisher’s exact and χ² tests for categorical variables, and Wilcoxon rank-sum and Kruskall-Wallis tests for continuous variables. Univariate logistic regression was used to identify predictors of poorly reliable FibroScan examinations including age, sex, underlying liver disease, liver stiffness, comorbidities, the FibroScan probe, the operator, operator experience and year. Variables that were significant in univariate analyses were included in stepwise-forward, multiple logistic regression models in which variables with P<0.1 were retained in the models. In the subset of patients with available BMI data from medical record review, an additional multivariate analysis was performed including BMI. All analyses were performed using Stata version 11.0 (StataCorp, USA); a two-sided P<0.05 was considered to be statistically significant.

RESULTS
Patient characteristics
A total of 2335 patients underwent LSM using TE at the UCLU between July 2008 and June 2011 and met the study inclusion criteria; their characteristics are outlined in Table 1. The median age was 50 years (IQR 40 to 58 years) and 56% were male. The majority of patients had chronic hepatitis C virus (36%) or HBV (27%) infection, while 7% had nonalcoholic fatty liver disease and 6% had autoimmune liver disease. Among 444 patients with BMI data available from medical record review, the median BMI was 25.5 kg/m²; 25% of patients were obese (BMI ≥30 kg/m²). Twelve percent of the cohort had a history of diabetes mellitus, 34% had hypertension, 35% had depression, and 13% and 10% had a history of alcohol and drug abuse, respectively.

LSM results
The majority of LSMs (86%) were obtained using the FibroScan M probe, 47% were performed by OP1 and 76% were performed during
Liver stiffness measurement (ie, failures excluded) This analysis is limited to 434 patients with available BMI data and successful cirrhosis (F4; liver stiffness ≥ 12.5 kPa) and FibroScan probe (M versus XL). This analysis is limited to 434 patients with available BMI data and successful cirrhosis (F4; liver stiffness ≥ 12.5 kPa) and FibroScan probe (M versus XL). This analysis is limited to 434 patients with available BMI data and successful cirrhosis (F4; liver stiffness ≥ 12.5 kPa) and FibroScan probe (M versus XL).

The latter part of the study (2010 to 2011). In total, LSM failure was observed in 37 (1.6%) patients. The incidence of FibroScan failure was greater with the XL probe (3.4% [11 of 326]) than the M probe (1.3% [26 of 2009]; P=0.01) and for OP1 (2.3% [25 of 1100]) compared with OP2 (1.0% [12 of 1235]; P=0.01). Among the 2335 successfully measured patients, the median LSM was 6.3 kPa (IQR 4.7 kPa to 10.1 kPa). The majority (58%) of patients were estimated to have F0 to F1 fibrosis (liver stiffness <7.1 kPa), while 19% had presumed cirrhosis (liver stiffness ≥12.5 kPa) (31).

**Figure 1** Prevalence of poorly reliable FibroScan (Echosens, France) examinations according to obesity (body mass index [BMI] ≥30 kg/m²), presumed cirrhosis (F4; liver stiffness ≥12.5 kPa) and FibroScan probe (M versus XL). This analysis is limited to 434 patients with available BMI data and successful liver stiffness measurement (ie, failures excluded)

**FibroScan reliability**

According to the reliability criteria of Boursier et al (19) and excluding LSM failures, FibroScan examinations were classified as very reliable in 29% (n=659), reliable in 66% (n=1526) and poorly reliable in 4.9% (n=113) of patients (Table 1). According to previously recommended criteria for reliability (18), 15% of these examinations (n=343) would have been classified as unreliable (valid shots <10, success rate <60% and/or IQR/M >30%). Based on the updated definitions of Boursier et al (19), 41% of these ‘unreliable’ examinations would have been classified as very reliable, 22% as reliable and 37% as poorly reliable.

**Predictors of poorly reliable FibroScan results**

Table 1 includes patient and procedural characteristics according to the reliability of the FibroScan examination. In univariate analysis, older age (OR per year 1.04 [95% CI 1.02 to 1.05]) was associated with an increased risk of poorly reliable LSM; male sex was of borderline significance (OR 1.45 [95% CI 0.98 to 2.16]; P=0.06). Patients with HBV were less likely to have a reliably LSM (OR 0.55 [95% CI 0.34 to 0.90]). Several comorbid conditions were associated with an increased risk of poorly reliable LSM including congestive heart failure, cardiac arrhythmias, hypertension, chronic pulmonary disease, diabetes mellitus, paralysis, peptic ulcer disease, coagulopathy, fluid and electrolyte disorders, and alcohol abuse (all P<0.05; Table 1).

Patients with poorly reliable LSM had higher median liver stiffness (13.3 kPa) than those with very reliable (6.0 kPa) and reliable scans (6.1 kPa; P=0.0005). The unadjusted odds of a poorly reliable scan increased by 4% per 1 kPa increase in liver stiffness (OR 1.04 [95% CI 1.03 to 1.05]). Similarly, presumed cirrhosis was more common in patients with poorly reliable LSM (57% versus 17% in patients with very reliable and reliable LSM; P=0.0005). Among 430 patients with cirrhosis, poorly reliable scans were recorded in 15% versus only 2.6% among noncirrhotic patients (OR 6.49 [95% CI 4.40 to 9.57]). In terms of procedural characteristics, poorly reliable LSM was less frequent with the M probe than the XL probe (4.1% versus 7.6%; OR 0.54 [95% CI 0.34 to 0.85]) and in scans performed by OP2 compared with OP1 (OR 0.45 [95% CI 0.30 to 0.66]). However, operator experience (<500 versus ≥500 scans) was not a significant predictor of poorly reliable LSM. Examinations performed in the latter half of the study period were less likely to be poorly reliable than earlier scans (2010 to 2011 versus 2008 to 2009 [4.3% versus 8.1%, respectively]; OR 0.47 [95% CI 0.32 to 0.69]).

Among 434 patients with available BMI data (excluding LSM failures), the median BMI was greater in patients with poorly reliable scans (30.0 kg/m² [IQR 25.5 to 32.0 kg/m²] versus 25.4 kg/m² [IQR 22.9 to 29.0 kg/m²]; P=0.004). Poorly reliable scans were observed in 10% (11 of 106) of obese patients (BMI ≥30 kg/m²) compared with only 3.1% (10 of 328) of nonobese individuals (OR 3.68 [95% CI 1.52 to 8.94]). Using the XL probe, 5.6% (two of 36) of obese patients had poorly reliable results versus 13% (nine of 70) measured using the M probe (P=0.33). The combined influence of obesity and presumed cirrhosis on the risk of poorly reliable LSM according to FibroScan probe is illustrated in Figure 1. In patients with obesity and cirrhosis, the risk of poorly reliable results was high (24% to 25%) with both probes compared with only 0% to 13.3% in patients with none or only one of these risk factors.

**Table 2**

**Independent predictors of poorly reliable FibroScan examinations**

| Variable                      | OR (95% CI) | P     |
|-------------------------------|------------|-------|
| Age†, per year                | 1.03       | (1.01 to 1.05) | <0.0005 |
| Male sex†                     | 1.32       | (0.87–1.99)    | 0.19    |
| Obesity‡ (body mass index ≥30 kg/m²) | 2.93     | (0.95–9.05)     | 0.06    |
| Chronic pulmonary disease     | 1.58       | (1.05–2.37)    | 0.03    |
| Coagulopathy                  | 2.22       | (1.31–3.76)    | 0.003   |
| Liver stiffness†, per kPa     | 1.03       | (1.02–1.05)    | <0.0005 |
| Cirrhosis§                    | 5.24       | (3.49–7.89)    | <0.0005 |
| M probe (versus XL probe)     | 0.64       | (0.38–1.06)    | 0.08    |
| Operator (OP1 versus OP2)     | 2.58       | (1.70–3.93)    | <0.0005 |
| Operator experience† (<500 versus ≥500 scans) | 0.86 | (0.67–1.10)    | 0.23    |

*Data from a stepwise-forward logistic regression model (bolded values represent statistical significance) including age, sex, liver stiffness, FibroScan (Echosens, France) probe, operator (OP), OP experience, year of scan (2008 to 2009 versus 2010 to 2011), liver disease (hepatitis B versus other) and all comorbidities with P=0.05 in Table 1; †Variable forced into the model based on pre-existing literature suggesting an association with FibroScan reliability. If not shown in the Table, variables were dropped from the model due to P>0.10 (eg, liver disease, year and other comorbidities); ‡OR for obesity from a separate model including the same variables as described above in the subset of patients with available body mass index data and excluding FibroScan failures (n=434); §OR for cirrhosis from a separate model including the same variables as described above.
DISCUSSION
In the present large practice audit of >2300 FibroScan examinations, poorly reliable results – as defined by the revised definition of Boursier et al (19) – were uncommon, occurring in approximately 5% of individuals. In contrast, unreliable results according to a previously recommended definition (valid shots <10, success rate <60% and/or IQR/M >30% [18]) were observed in 15% of patients. This difference is relevant because the latter ‘unreliable’ results have never been shown to be less accurate than reliable results. In fact, use of this outdated definition of reliability may have led to the needless discarding of approximately 10% of FibroScan results, with potentially important implications in clinical practice and in research studies. Importantly, approximately two-thirds of these results would have been classified as very reliable or reliable according to the revised definitions. The prevalence of poorly reliable results observed in our study is slightly lower than that of Boursier et al (19) (4.9% versus 9.1%). This likely reflects differences in the study populations, particularly the lower prevalence of advanced fibrosis in our study (median liver stiffness 6.3 kPa versus 8.1 kPa in Boursier et al [19]) because higher liver stiffness (≥7.1 kPa) is a criterion in the definition for reliability (see below). Moreover, liver stiffness was measured using the XL probe in 25% of our patients, whereas only the M probe was used in the French study. Because liver stiffness measured with the XL probe is consistently lower than with the M probe, a lower rate of poorly reliable results could also be anticipated in our study.

Because TE is increasingly being used in clinical decision making, it is important to understand factors that influence its reliability because poorly reliable results are less accurate than reliable examinations. Specifically, in the study by Boursier et al (19), poorly reliable results were only 70% accurate for the diagnosis of cirrhosis compared with 86% and 90% in patients with reliable and very reliable results, respectively. Corresponding AUCROCs for cirrhosis were 0.82, 0.90 and 0.97, respectively. With these facts in mind, we examined several patient- and operator-related characteristics as potential predictors of unreliable examinations. As previously reported, older age was associated with an increased risk of unreliable LSM (18). The exact reasons for this finding have never been identified; however, we speculate that age-related alterations in the chest wall are involved. It is known that chest wall compliance decreases with age due to structural changes of the intercostal muscles, intercostal joints and rib-vertebral articulations. In addition, age-associated osteoporosis may increase kyphosis, resulting in changes in the geometry of the thorax (37). On a related note, we identified a 1.6-fold increase in the risk of poorly reliable results among patients with chronic lung disease, predominantly chronic obstructive pulmonary disease (data not shown). This novel finding may also relate to structural changes in the chest wall (eg, pulmonary hyperinflation) or technical difficulties with the FibroScan procedure due to deep respirations in these patients. Because an increased risk of LSM failure or unreliable results has not been reported in cohorts with cystic fibrosis (38), additional studies are necessary to confirm this finding and to elucidate potential mechanisms.

Although previous studies have reported that women have a higher rate of unreliable FibroScan examinations compared with men (18), we did not observe a significant impact of sex using the updated definition of reliability. As previously reported (18), obesity was associated with a nearly threefold risk of unreliable results in a subanalysis of patients with available BMI data. The importance of obesity as a predisposing factor for poorly reliable results is supported by the borderline effect of XL probe use (P=0.08), considered a surrogate marker for obesity in the absence of BMI data in all patients. Presumably, subcutaneous and prehepatic adipose tissue in obese patients interferes with transmission of the mechanical shear wave and/or the measurement of its propagation by the FibroScan device (7).

Previous studies by Lucidarme et al (39) and Myers et al (40) have demonstrated an impact of fibrosis stage on the rate of discordance between fibrosis estimated by LSM and liver biopsy; however, an impact on poorly reliable results has not been reported. In the current study, elevated liver stiffness was an independent predictor of poorly reliable LSM. Presumed cirrhosis was the most important risk factor, with fivefold higher odds of poorly reliable results in cirrhotic patients. This finding is not surprising because an LSM ≥7.1 kPa is one criterion in the definition of poor reliability (19). The other factor, IQR/M, also likely played a role because LSM variability tends to be greater in patients with cirrhosis (data not shown). The reason for this is unclear, but may relate to the broader range of potential LSMs in cirrhotic patients (ie, approximately 12.5 kPa to 75 kPa) compared with those who have lower liver stiffness (ie, 2.5 kPa to 12.4 kPa) (31). Interestingly, coagulopathy was associated with a twofold risk of poorly reliable results. Because there is no clear physical explanation for this finding, we suspect it reflects more severe liver disease and, therefore, higher liver stiffness in coagulopathic patients. The majority of these patients had diagnosis codes for thrombocytopenia as opposed to hereditary or acquired coagulation defects (data not shown). Because the effect of coagulopathy on poorly reliable results was independent of liver stiffness, it likely reflects the imperfect sensitivity of FibroScan for the diagnosis of cirrhosis.

A strength of our study was our analysis of the impact of specific liver conditions and comorbidities on FibroScan reliability using administrative data. In addition to uncovering a novel association between chronic lung disease and poorly reliable results, this approach revealed several other associations. First, patients with HBV had a lower likelihood of unreliable results. Because HBV-infected patients in our practice tend to be Asian and of smaller stature, this finding is not unexpected. In fact, we previously reported no FibroScan failures with the M probe in a study examining the value of the pediatric (S2) probe in this patient population (41). Second, hypertension and diabetes, components of the metabolic syndrome that have been associated with unreliable LSMs in previous studies (18), were significant in unadjusted, but not adjusted, analyses. Similarly, patients with congestive heart failure or arrhythmias had a twofold higher risk of poorly reliable LSM in univariate analyses. These findings may relate to hepatic congestion due to cardiac dysfunction, a well-described cause of liver stiffness overestimation (42). Because the power of our multivariate analysis may have been limited due to a small number of poorly reliable results (n=113), studies that use this novel definition of reliability in larger patient populations will be necessary to confirm or refute these findings.

In addition to patient-related predictors of FibroScan reliability, we examined the impact of procedural characteristics including the operator and operator experience. Castera et al (18) reported a lower risk of unreliable results among seven operators who had performed at least 500 examinations, challenging previous assertions that that a novice can consistently obtain reliable results after a short training period of only 50 examinations. In our study, however, there was no difference in the proportion of poorly reliable results between the first 500 and subsequent examinations by our two operators. On the contrary, OP1 was twice as likely to produce a poorly reliable LSM as OP2. This finding supports the importance of adequate operator training and ongoing quality control when using the FibroScan in clinical decision making. It is important to note, however, that these results were confounded by the availability of the FibroScan XL probe only during the final two years of the study. Therefore, OP1 – who was employed before OP2 – scanned many obese patients using the M probe during the early part of the study. This likely led to an overestimation of her true rate of unreliable results. In fact, OP1 was twice as likely to have FibroScan failure as OP2, presumably for the same reason.

Our study has several limitations that warrant discussion. First, we did not have histological data to confirm whether poorly reliable LSMs were less accurate for staging fibrosis than reliable examinations. Second, because this was a retrospective study, we did not prospectively collect data regarding BMI and other anthropometric measures (eg, waist circumference, thoracic perimeter, skin-capsular distance)
that may have influenced FibroScan reliability. Similarly, we relied on administrative data to define patient comorbidities and hepatic diagnoses. Although many of these codes have been validated by medical record review (20,24-28), additional validation is necessary.

**CONCLUSION**

Poorly reliable FibroScan results are uncommon, occurring in approximately 5% of individuals. The most important determinants of poorly reliable LSM are older age, obesity, higher liver stiffness and the operator, the latter emphasizing the need for adequate training and quality control. A novel association between chronic pulmonary disease and poorly reliable results requires confirmation. Additional studies should be conducted to identify means of improving the reliability of FibroScan examinations so as to improve the accuracy of this valuable tool.

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**APPENDIX 1**

**Diagnosis codes used to define underlying liver disease etiologies**

| Hepatic diagnoses | ICD-9-CM (32) | ICD-10 (33) |
|-------------------|---------------|-------------|
| Hepatitis B       | 070.2, 070.3  | B16, B19.0  |
|                   |               | B18.1, B19.1|
| Hepatitis C       | 070.41, 070.44, 070.51, B17.1, B18.2, 070.54, 070.7 | B19.2 |
| Nonalcoholic fatty liver disease | 571.8 | K75.81, K76.0 |
| Alcoholic liver disease | 571.0, 571.1, 571.2, 571.3 | K70 |
| Primary biliary cirrhosis | 571.6 | K74.3 |
| Primary sclerosing cholangitis | 576.1 | K83.0 |
| Autoimmune hepatitis | 571.42 | K75.4 |
| Hemochromatosis    | 275.0         | E83.1 |

**ICD International Classification of Diseases**

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