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Prognostic utility of magnetic resonance elastography and MEFIB index in predicting liver-related outcomes and mortality in individuals at risk of and with nonalcoholic fatty liver disease

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Veeral Ajmera, Khang Nguyen, Nobuharu Tamaki, Suzanne Sharpton, Ricki Bettencourt and Rohit Loomba

Prognostic utility of magnetic resonance

liver-related outcomes and mortality in

elastography and MEFIB index in predicting

individuals at risk of and with nonalcoholic

Abstract

Background: Magnetic resonance elastography (MRE) is an accurate biomarker of liver fibrosis; however, limited data characterize its association with outcomes. We aimed to evaluate the association between liver stiffness (LS) on MRE and liver-related outcomes. **Methods:** This is a longitudinal, retrospective analysis of subjects at risk of NAFLD who had MRE assessment. LS was estimated using MRE, and liver fat was assessed using magnetic resonance imaging proton density fat fraction. Univariable and multivariable survival and regression analyses were used to assess the association between LS on MRE and liver-related outcomes including a cumulative primary outcome of hepatic decompensation, hepatocellular carcinoma (HCC), or death.

Results: In all, 265 patients (68% women) with a mean age of 50 (±18) years and 44% Hispanic ethnicity and 45.3% with NAFLD were included. A total of 76 liver-related events or death occurred, and there was 453 person-years of follow-up time in 97 patients with available follow-up. Each 1-kPa increase in LS was associated with 2.20-fold (95% CI: 1.70–2.84, p < 0.001) increased odds of prevalent hepatic decompensation or HCC. A positive MEFIB index, a combination of MRE \geq 3.3 kPa and FIB-4 \geq 1.6, had a strong association with the primary outcome compared with those without, HR = 21.8 (95% CI: 4.28–111.4, p < 0.001). The MEFIB index had a sensitivity of 75% and specificity of 90%, and a negative score was associated with 98% negative predictive value for incident liver-related events or death. **Conclusion:** LS assessed by MRE is associated with hepatic decompensation and death, and the MEFIB combination of MRE with FIB-4 may have high negative predictive value for liver-related events.

Keywords: ascites, biomarker, cirrhosis, nonalcoholic fatty liver disease, noninvasive, portal hypertension, varices

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Introduction

Nonalcoholic fatty liver disease (NAFLD) is emerging as a leading cause of chronic liver disease globally, and given the affected population, great emphasis has been placed on noninvasive assessment of disease.^{1,2} Despite the high prevalence of NAFLD, only a subset of patients develop decompensated cirrhosis and hepatocellular carcinoma (HCC).³ Fibrosis stage on liver biopsy is strongly associated with outcomes;⁴ however, Correspondence to: Rohit Loomba

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liver biopsy is impractical to scale to the affected population, leaving a great need for noninvasive biomarkers associated with liver-related outcomes.

Magnetic resonance elastography (MRE) is a reproducible,⁵ continuous measure of liver stiffness, the most accurate noninvasive test (NIT) for fibrosis stage on liver biopsy,6,7 and examines a much larger volume of the liver than a biopsy. An increase in liver stiffness on MRE is associated with fibrosis progression,8 and MRE is increasingly being used as an end point in early stage NAFLD treatment trials.9,10 Emerging data from recent studies have suggested that liver stiffness on MRE may be associated with liver-related outcomes.^{11–14} Other NITs including the FIB-4 score (fibrosis index based on the 4 factor) and NAFLD Fibrosis Score (NFS) have demonstrated associations with liver-related events and death;15-17 however, when used alone, they have only modest discriminatory ability. Similarly, vibration-controlled transient elastography (VCTE) is associated with liver-related events among patients with chronic advanced liver disease,18 but has limited discriminatory ability for liver-related events in patients with less advanced disease.

Recently, the combination of NITs has demonstrated improved accuracy for diagnosing signifiactivity.19,20 cant fibrosis and NAFLD Furthermore, the combination of liver stiffness on VCTE and platelet count have been used to identify patients with chronic liver disease with clinically significant portal hypertension.²¹ А prospective study demonstrated that the combination of MRE \ge 3.3 kPa and FIB-4 \ge 1.6 (MEFIB index) provided excellent positive predictive value (PPV) of 97.1% for significant fibrosis and was validated in an independent cohort²⁰ to identify candidates for pharmacologic treatment for nonalcoholic (NASH) who have NASH with stage 2 fibrosis or higher. To date, studies evaluating the association between MRE and liver-related outcomes have not evaluated a combination of NITs to optimize diagnostic performance in clinical practice. Therefore, we hypothesized that liver stiffness on MRE would be associated with liverrelated events and death and that the combination of MRE and FIB-4 would improve the ability to identify low- and high-risk groups. Using a wellcharacterized cohort of patients at risk of NAFLD, we evaluated the association between the liver

stiffness on MRE and the use of the MEFIB index to predict hepatic decompensation or death.

Materials and methods

Study design

This study includes a cross-sectional and longitudinal analysis derived from two cohorts of patients who had a research MRE assessment and were at risk of NAFLD, with other causes of chronic liver disease excluded, with a retrospective evaluation for liver-related events and death. This study included 265 patients who underwent a standardized research visit including history, physical exam, and MRE assessment between 2011 and 2020 at the UCSD NAFLD Research Center.²²⁻²⁶ All patients provided written informed consent prior to enrolling in the study, and the study was approved by the UCSD Institutional Review Board. The reporting of this study conforms to the STROBE statement²⁷ with a completed checklist available in supplemental material.

Inclusion and exclusion criteria

The study population was derived from two cohorts at risk of NAFLD, defined as having other causes of chronic liver disease ruled out through history and laboratory testing. The Twin and Family Study $(N=141)^{24}$ recruited participants who were twins, siblings, or parent–off-spring pairs at least 18 years old residing in Southern California. Participants were excluded from the study if they met any of the following criteria: significant alcohol intake (defined as ≥ 14 drinks/week for men or ≥ 7 drinks/week for women) within the previous 2-year period or biochemical evidence of liver disease other than NAFLD.

Participants were also derived from the Familial Cirrhosis Study (N=124),²⁸ which included probands with NAFLD cirrhosis defined by American Association for the Study of Liver Diseases (AASLD) and their first-degree family members (sibling, child, or parent). Exclusion criteria included significant alcohol intake (defined as ≥ 14 drinks/week for men or ≥ 7 drinks/week for women) within the previous 2-year period or biochemical evidence of liver disease other than NAFLD.

Clinical assessment and laboratory tests

All patients underwent a standardized clinical evaluation including a detailed history and a physical examination, which included vital signs, height, weight, and anthropometric measurements, performed by a trained clinical investigator. Body mass index (BMI) was defined as the body weight (in kilograms) divided by height (in meters) squared. Alcohol consumption was documented outside clinical visits and confirmed in the research clinic using the Alcohol Use Disorders Identifications Test (AUDIT) and the Skinner questionnaire. Other causes of liver disease were systematically ruled out based on history and laboratory tests. Patients underwent the following biochemical tests: glucose, albumin, hemoglobin A1c, alanine aminotransferase, aspartate aminotransferase, total bilirubin, alkaline phosphatase, fasting lipid panel, platelets, insulin, and international normalized ratio. FIB-4²⁹ and NFS³⁰ were calculated as described previously. Participants were instructed to fast for a minimum of 8h prior to collection of laboratory tests.

Magnetic resonance imaging

Abdominal magnetic resonance imaging (MRI) was obtained on a single 3T MR scanner (GE Signa EXCITE HDxt, GE Healthcare, Waukesha, WI) at the UCSD MR3T Research Laboratory using previously described methods.^{31–35} Magnetic resonance imaging proton density fat fraction (MRI-PDFF) was used to measure hepatic steatosis, and NAFLD is defined as having an MRI-PDFF \ge 5% or a liver stiffness \ge 3.63 kPa plus a metabolic risk factor after exclusion of other causes of liver disease. Liver stiffness was estimated using 2-dimensional MRE, which is the most accurate biomarker for the quantitative assessment of liver stiffness as a surrogate for hepatic fibrosis.20,36,37 A passive driver was fitted around the body over the liver and connected to an acoustic active driver that delivered continuous vibrations at 60 Hz to produce shear waves in the liver, which were processed to generate elastograms depicting liver stiffness. Four slices were assessed, and co-localized regions of interest (ROIs) were manually specified as previously described.38 Assessment of the mean MRE value was performed independent of knowledge of clinical outcomes with the exception of those apparent on imaging.

Justification for evaluating MRE cut points and MEFIB index

The cut point of ≥ 3.63 kPa for advanced fibrosis is the most well-validated MRE cut point and has been evaluated in multiple studies^{6,39} and an individual patient meta-analysis⁷ with excellent diagnostic accuracy (*c*=0.93) for advanced fibrosis (stage 3–4). Furthermore, a liver stiffness measurement (LSM) cut point of 5 kPa on MRE has high specificity (90%) for the presence of cirrhosis and was included in the analysis for liverrelated outcomes.⁷

The MEFIB combination of MRE and FIB-4 has identified a cut point that leverages MRE to identify patients with stage 2 fibrosis or higher with a high PPV.^{20,40} The index includes patients who are at intermediate risk by FIB-4 and accurately identifies those who actually have stage 2 fibrosis or higher. The population identified by NITs, including the MEFIB index, may be used to select candidates for clinical trials. Assessing their longterm risk and documenting low risk for those excluded is an important clinical as well as validation question, which this study is designed to address.

Patient follow-up

Two modes of follow-up were employed to adjudicate patient outcomes. Patients were contacted by telephone, and an outcome questionnaire was administered. Three attempts were made to contact each patient prior to stating that the patient was lost to follow-up. In addition, the electronic health record was reviewed for each patient by two hepatologists (V.A. and N.T.) to assess for hepatic decompensation, HCC, or death. Patients with no follow-up time were only evaluated in cross-section for outcomes present at baseline.

Outcome measures

The *primary outcome* was a cumulative end point including ascites, hepatic encephalopathy, varices needing treatment, HCC, and death and for assessment of incident events was the time to the earliest event. Ascites was defined per AASLD guidance by imaging or physical examination.⁴¹ Hepatic encephalopathy was defined as brain dysfunction caused by liver dysfunction and/or portosystemic shunting per practice guidelines.⁴² Varices needing treatment were defined as

medium/large varices, small varices with high-risk stigmata, decompensated patients with small varices, or variceal hemorrhage per AASLD guidance.⁴³ HCC was defined by histology or Liver Reporting and Data Systems (LI-RADS) for definite HCC.

Secondary outcomes included each individual component of the primary outcome and liver transplantation.

Statistical analysis

We hypothesized that there would be an association between liver stiffness on MRE and the cumulative outcome. Power analysis was performed assuming that 15% of the cohort had $MRE \ge 3.63 \text{ kPa}$, consistent with advanced fibrosis, and that the presence of MRE \ge 3.63 kPa was associated with a 15% event rate over the follow-up period compared with an event rate of 2% for the cumulative outcome in those without advanced fibrosis. A sample size of 264 would provide 80% power with a two-tailed alpha of 0.05; therefore, we had adequate power to test and confirm our hypothesis. For patient characteristics, an ANOVA was performed on continuous variables presented as mean [standard deviation (SD)], and Kruskal-Wallis was performed on those presented as median [interquartile range (IQR)]. Chi-square or Fisher's exact test was conducted as appropriate on all categorical variables. Kaplan-Meier curves were generated to evaluate for the cumulative incidence of liver-related events or death for the subset of patients with longitudinal follow-up. For the cumulative outcome, time was measured to the first event. Patients were censored at the last known follow-up in the clinical record. Logistic regression was used to assess for the prevalent association between liver-related events and liver stiffness at baseline. For patients with follow-up and without the event at baseline, Cox proportional hazard models were used to calculate hazard ratios (HRs). Dichotomous thresholds on MRE at cut points of 3.63 and 5 kPa were also evaluated. The combination of FIB-4 and MRE in the MEFIB index (MRE \ge 3.3 kPa and FIB-4 \ge 1.6) was evaluated for its association with outcomes. All statistical analyses were performed using SAS 9.4 (SAS Institute), and a p value less than 0.05 was considered statistically significant.

Results

Characteristics of the study population

Two hundred sixty-five patients who underwent MRE were included. Participants had a mean age of 50 (± 18) years, were predominantly female (68%), and were 46% White and 44% Hispanic. The prevalence of NAFLD was 45.3%. Advanced fibrosis (MRE \ge 3.63 kPa) on MRE was present in 18% of the cohort and 12% had MRE > 5 kPa. The mean BMI was 29 (\pm 7) kg/m² (Table 1). A total of 29 patients had 76 events [ascites (n=17), hepatic encephalopathy (n=17), varices needing treatment (n=18), HCC (n=10), and death n=14] and 453 person-years of follow-up among 97 patients with longitudinal follow-up data. Participants without follow-up data were only assessed for prevalent events at the time of MRE (Supplemental Figure 1). Four patients underwent subsequent liver transplantation. Higher liver stiffness was associated with older age, higher BMI, Hispanic ethnicity, the presence of diabetes, and hypertension (Table 1).

Factors associated with prevalent liver-related outcomes

Twenty-one participants had 40 events at baseline. Higher LSM was associated with prevalent liver-related outcomes [odds ratio (OR) = 2.20per 1 kPa increase; 95% confidence interval (CI): 1.70-2.84, p < 0.0001]. The association was amplified at higher LSM with MRE of 3.63-5.0 kPa and >5 kPa associated with an OR = 13.4 (95% CI: 2.7-66.6, p=0.002) and OR=41.6 (95% CI: 12.4–139.7, p<0.001), respectively, for the cumulative outcome compared with MRE < 3.63 kPa and remained significant in multivariable analysis adjusted for age and sex: OR=7.4 (95% CI: 1.3-41.5, p=0.022) and OR = 26.2 (95% CI: 7.6–90.5, p < 0.001). In sensitivity analysis limited to subjects with NAFLD at baseline (n=120), higher LSM with MRE of 3.63-5.0 kPa and >5 kPa associated with an OR = 12.7 (95% CI: 1.06–150.70, p = 0.045) and OR=70.93 (95% CI: 8.66–580.87, p < 0.001), respectively, for the cumulative outcome compared with MRE < 3.63 kPa. Similarly, in the subset of patients who did not meet the study definition for NAFLD, the results remained consistent with MRE \ge 3.63 kPa associated with a higher risk of the primary outcome, OR=15.33 (95% CI: 1.21-193.6, p=0.035), compared with

 Table 1. Clinical, demographic, and imaging characteristics by liver stiffness on MRE.

MRE liver stiffness	Total N	<3.63 kPa N=218	3.63–5 kPa N=15	>5 kPa N=32	p
Demographic					
Age in years, mean (SD)	265	47.4 (17.8)	60.9 (14.6)	62.5 (10.1)	<0.0001
Male, <i>n</i> (%)	265	67 (30.7%)	8 (53.3%)	9 (28.1%)	0.1716
BMI (kg/m²), mean (SD)	255	28.5 (7.3)	32.3 (4.7)	31.0 (6.1)	0.0350
White, <i>n</i> (%)	256	107 (50.9%)	3 (21.4%)	8 (25.0%)	0.0038
Hispanic, n (%)	256	84 (40.0%)	8 (57.1%)	21 (65.6%)	0.0149
Diabetes, n (%)	265	22 (10.1%)	6 (40.0%)	22 (68.8%)	<0.0001
Hypertension, <i>n</i> (%)	265	50 (22.9%)	8 (53.3%)	16 (50.0%)	0.0005
Hyperlipidemia, <i>n</i> (%)	265	41 (18.8%)	4 (26.7%)	7 (21.9%)	0.7165
Current smoker, <i>n</i> (%)	236	17 (8.8%)	1 (7.7%)	3 (10.3%)	0.8956
Family history					
Liver disease, n (%)	265	20 (9.2%)	1 (6.7%)	3 (9.4%)	1.0000
Biochemical profile					
HbA1c (%), median (IQR)	259	5.6 (0.6)	6.2 (1.2)	6.2 (1.6)	0.0001
AST (U/l), median (IQR)	260	22.0 (9.0)	32.0 (17.0)	53.0 (38.0)	<0.0001
ALT (U/l), median (IQR)	259	19.5 (13.0)	31.0 (31.0)	32.0 (33.0)	<0.0001
Alkaline phosphatase (U/l), median (IQR)	260	71.0 (25.0)	102.0 (46.0)	107.0 (49.5)	<0.0001
Total bilirubin (mg/dl), median (IQR)	260	0.4 (0.3)	0.6 (0.4)	0.8 (1.0)	<0.0001
Albumin (g/dl), median (IQR)	260	4.5 (0.4)	4.3 (0.2)	4.1 (0.6)	<0.0001
HOMA-IR median (IQR)	243	2.5 (2.6)	7.3 (4.4)	10.3 (13.4)	<0.0001
Triglycerides (mg/dl), median (IQR)	259	98.0 (71.0)	138.0 (99.0)	105.0 (81.0)	0.1785
HDL (mg/dl), median (IQR)	257	56.0 (22.0)	43.0 (10.0)	41.0 (21.0)	<0.0001
LDL (mg/dl), median (IQR)	255	106.0 (48.0)	85.0 (43.0)	84.0 (35.0)	0.0010
Platelet count (10º/l), median (IQR)	259	257.0 (82.5)	189.0 (86.0)	119.0 (90.5)	<0.0001
INR, median (IQR)	259	1.0 (0.1)	1.1 (0.2)	1.2 (0.2)	<0.0001
Clinical scores					
FIB-4, median (IQR)	256	0.9 (0.7)	1.7 (1.7)	4.2 (3.6)	<0.0001
NAFLD Fibrosis Score, median (IQR)	246	-2.5 (1.7)	0.3 (2.5)	1.2 (2.1)	<0.0001
MELD score, median (IQR)	256	6.0 (1.0)	7.0 (1.0)	8.0 (5.0)	<0.0001

(Continued)

THERAPEUTIC ADVANCES in

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Table 1. (Continued)

3.63–5 kPa p N=15 N=32	
9.5 (6.5) 5.9 (5.0) 0.1	426
4.3 (0.4) 7.3 (2.1) -	
14 (93.3%) 29 (90.6%) < 0	.0001
	N=15 N=32 9.5 (6.5) 5.9 (5.0) 0.1 4.3 (0.4) 7.3 (2.1) -

AST, aspartate aminotransferase; ALT, alanine aminotransferase; BMI, body mass index; FIB-4, fibrosis index based on the 4 factors; HbA1c, hemoglobin A1c; HDL, high-density lipoprotein; HOMA-IR, homeostatic model assessment method for insulin resistance [calculated as (fasting insulin (μ U/ml) × fasting glucose (mmol/l)/22.5]; IQR, interquartile range; INR, international normalized ratio; LDL, low-density lipoprotein; MRI-PDFF, MRI-based proton density fat fraction; MRE, magnetic resonance elastography; SD, standard deviation.

ANOVA performed on continuous variables presented as mean (SD), and Kruskal–Wallis was performed on all other continuous variables. Chi-square or Fisher's exact test was conducted as appropriate on all categorical variables. Statistically significant *p* values are denoted in bold.

> MRE < 3.63 kPa. A 1-unit increase in FIB-4 [OR=1.94 (95% CI: 1.40–2.68, p < 0.001)] and NFS [OR=1.73 (95% CI: 1.16–2.66, p=0.01)] was associated with higher odds of liver-related outcome or death (Table 2).

Higher liver stiffness on MRE was associated with an increased risk for each liver-related event (Table 3). Liver-related events increased in frequency with higher categories of liver stiffness (<3.63 kPa, 3.63-5 kPa, and >5 kPa) for varices needing treatment (1.9%, 20%, and 34%), ascites (1.4%, 6.7%, and 41%), hepatic encephalopathy (1.4%, 13.3%, and 37.5%), and death (0.5%, 20%, and 18.8%). The risk of HCC was low in MRE < 3.63 (0.5%) but similar in 3.63-5kPa (20.0%) and >5 kPa (18.8%). The results remained consistent on sensitivity analysis of a composite outcome of only hepatic decompensation (varices needing treatment or ascites or hepatic encephalopathy) [OR=2.39 per 1kPa increase (95% CI: 1.82–3.13, $p \le 0.0001$)]. The ordinal categories 3.63-5.0 kPa and >5 kPa were associated with an OR=13.19 (95% CI: 2.65-65.71, *p*=0.0016) and OR=52.75 (95% CI: 15.77–176.51, $p \le 0.0001$), respectively, for the hepatic decompensation compared with MRE < 3.63 kPa. Similarly, in sensitivity analysis evaluating the association between MRE and risk of HCC, higher liver stiffness was associated with increased risk of HCC [OR=1.55 per 1kPa increase (95% CI: 1.26–1.91, $p \le 0.0001$)]. The ordinal categories 3.63-5.0 kPa and >5 kPa were associated with an OR=53.5 (95% CI: 5.17-553.55, p = 0.0008) and OR = 49.39 (95% CI:

5.72–426.42, p=0.0004) for HCC compared with < 3.63 kPa, respectively.

MEFIB index and liver-related outcomes or death

A positive MEFIB index, defined as a combination of MRE \geq 3.3 kPa and FIB-4 \geq 1.6, was present in 39 patients and had a strong association with prevalent and future incident liver-related outcomes or death compared with those with a negative MEFIB (59.0% *versus* 2.7%). The OR for the primary outcome was OR = 42.9 (95% CI: 13.3–138.7, p < 0.001) for a positive MEFIB *versus* a negative MEFIB and remained significant in multivariable analysis adjusted for age and sex, OR = 24.1 (95% CI: 7.2–81.4, p < 0.001).

Factors associated with incident liver-related outcomes or death

Among participants without liver-related events at baseline with longitudinal follow-up after MRE (n=97 and 40.2% with NAFLD), 12 incident cumulative events occurred over 453 person-years of follow-up in eight patients. In addition, 24 incident events occurred in 21 patients with at least one event at baseline. The incidence of the cumulative outcome was 0.5%, 10.0%, and 10.3% per person-year in the <3.63 kPa, 3.63– 5 kPa, and >5 kPa groups, respectively. The 3-year risk of incident decompensation, HCC, or death was 0% for MRE < 3.63 kPa, 16.7% for MRE=3.63–5 kPa, and 25% for MRE >5 kPa among those without a decompensating event at

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Table 2. Factors associated with prevalent liver-related outcomes.

	Liver-related outcomes or death OR (95% CI)	p value
MRE		
<3.63 kPa (referent)	Ref	
3.63-5 kPa	13.37 (2.68–66.64)	0.0015
>5kPa	41.61 (12.40–139.66)	<0.0001
Demographic and biochemical		
BMI (kg/m²)	1.04 (0.99–1.10)	0.1397
HbA1c (%)	1.42 (1.03–1.95)	0.0309
AST (U/l) (per 5-unit increase)	1.20 (1.10–1.30)	<0.0001
ALT (U/l) (per 5-unit increase)	1.04 (0.98–1.10)	0.1881
Alkaline phosphatase (U/l) (per 5-unit increase)	1.25 (1.15–1.36)	<0.0001
Total bilirubin (mg/dl)	18.23 (6.55–50.77)	<0.0001
Albumin (g/dl)	0.007 (0.001–0.046)	<0.0001
HOMA-IR	1.07 (1.03–1.12)	0.0026
Triglycerides (mg/dl) (per 5-unit increase)	1.02 (0.99–1.04)	0.1409
HDL (mg/dl) (per 5-unit increase)	0.78 (0.65–0.94)	0.0083
LDL (mg/dl) (per 5-unit increase)	0.88 (0.81–0.95)	0.0017
Platelet count (10%) (per 10-unit increase)	0.77 (0.71–0.84)	<0.0001
Clinical score		
FIB-4	2.39 (1.77–3.22)	<0.0001
NAFLD Fibrosis Score	3.07 (2.08–4.53)	<0.0001
MELD score (per 1-unit increase)	1.63 (1.36–1.96)	<0.0001
Positive MEFIB	42.89 (13.26–138.73)	<0.0001
Imaging		
MRI-PDFF	0.90 (0.80–1.02)	0.0959

ALT, alanine aminotransferase; AST, aspartate aminotransferase; BMI, body mass index; FIB-4, fibrosis index based on the 4 factors; HbA1c, hemoglobin A1c; HDL, high-density lipoprotein; HOMA-IR, homeostatic model assessment method for insulin resistance (calculated as (fasting insulin (μ U/ml)× fasting glucose (mmol/l)/22.5); INR, international normalized ratio; LDL, low-density lipoprotein; MRI-PDFF, MRI-based proton density fat fraction. Statistically significant *p* values are denoted in bold.

baseline. The incident risk of liver-related outcomes and death increased significantly across the three liver stiffness categories, p < 0.001(Figure 1). A positive MEFIB index was also associated with a higher risk of incident events, HR=21.84 (95% CI: 4.28–111.4, p < 0.001) (Table 4; Figure 2). Two incident events in patients with a negative MEFIB index occurred at

	Total number	MRE liver stiffness			p valueª
	of events	<3.63 kPa N=218	3.63–5 kPa <i>N</i> = 15	>5 kPa N=32	
Varices needing treatment, N (%)	18	4 (1.9%)	3 (20.0%)	11 (34.4%)	<0.0001
Ascites, N (%)	17	3 (1.4%)	1 (6.7%)	13 (40.6%)	<0.0001
Hepatic encephalopathy, <i>N</i> (%)	17	3 (1.4%)	2 (13.3%)	12 (37.5%)	<0.0001
Hepatocellular carcinoma, N (%)	10	1 (0.5%)	3 (20.0%)	6 (18.8%)	<0.0001
Liver transplant, N (%)	4	0	0	4 (12.5%)	0.0003
Death, <i>N</i> (%)	14	5 (2.3%)	1 (6.7%)	8 (25.0%)	<0.0001

Table 3. Liver-related outcomes and death by liver stiffness on MRE.

^ap value calculated using Fisher's exact test. Statistically significant p values are denoted in bold.

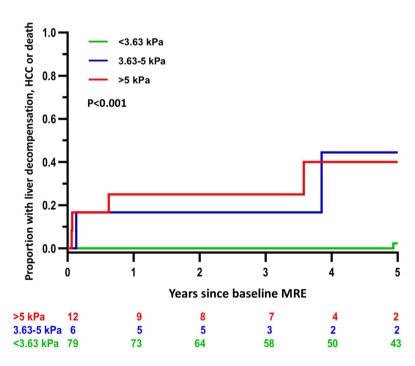


Figure 1. Five-year cumulative incidence of hepatic decompensation, hepatocellular carcinoma, or death by liver stiffness cut points on MRE < 3.63, 3.63-5 kPa, and >5 kPa.

4.93 and 8.38 years after baseline. Importantly, the sensitivity and specificity of MEFIB for the cumulative outcome was 75% and 90%, and a negative score was associated with a high negative predictive value (NPV) of 98% for liver-related events or death.

Discussion

Using a well-phenotyped cohort of subjects with MRE assessment and liver-related outcome assessment, we demonstrate that liver stiffness on MRE is strongly associated with liver-related outcomes or death. The risk of decompensation and death increased with liver stiffness greater than or equal to the established cut point of 3.63 kPa and was highest among those with liver stiffness greater than 5 kPa where there was a 25% risk of decompensation within 1 year. Importantly, MEFIB index, which is positive when MRE \geq 3.3 kPa and FIB-4 \geq 1.6, had high NPV, 98%, to rule out future liver-related events.

In context with published literature

MRE is the most accurate biomarker of liver fibrosis in NAFLD;⁴⁴ however, it has been evaluated primarily against liver biopsy as the reference standard. Liver biopsy interpretation suffers from only modest inter- and intra-observer agreement for the assessment of characteristics of histologic NAFLD including fibrosis stage.^{45,46} Instead, assessment of the direct association between MRE and liver-related outcomes and death is critical to determining its value as a biomarker of liver disease.

Matsui and colleagues performed a cross-sectional analysis evaluating the association between liver stiffness on MRE and gastroesophageal varices and demonstrated that a combination of Table 4. Factors associated with incident liver-related outcomes or death.

	Liver-related outcomes or death Hazard ratios (95% CI)	<i>p</i> value
MRE		
<3.63 kPa (referent)	ref	
3.63-5 kPa	17.09 (2.38–122.75)	0.0048
>5 kPa	16.58 (2.90–94.79)	0.0016
Demographic and biochemical		
BMI (kg/m²)	1.02 (0.92–1.13)	0.6660
HbA1c (%)	1.69 (0.91–3.14)	0.0941
AST (U/l) (per 5-unit increase)	1.17 (1.07–1.28)	0.0007
ALT (U/l) (per 5-unit increase)	1.13 (1.05–1.21)	0.0008
Alkaline phosphatase (U/l) (per 5-unit increase)	1.11 (0.99–1.26)	0.0823
Total bilirubin (mg/dl)	5.90 (1.25–27.82)	0.0249
Albumin (g/dl)	0.16 (0.01–1.94)	0.1512
HOMA-IR	1.03 (0.98–1.07)	0.2637
Triglycerides (mg/dl) (per 5-unit increase)	1.03 (0.99–1.07)	0.1790
HDL (mg/dl) (per 5-unit increase)	0.60 (0.40–0.89)	0.0107
LDL (mg/dl) (per 5-unit increase)	0.96 (0.86–1.07)	0.4486
Platelet count (10º/l) (per 10-unit increase)	0.85 (0.76–0.96)	0.0077
Clinical score		
FIB-4	1.94 (1.40–2.68)	<0.0001
NAFLD Fibrosis Score	1.73 (1.13–2.66)	0.0115
MELD score (per 1-unit increase)	1.34 (1.07–1.69)	0.0122
Positive MEFIB	21.84 (4.28–111.40)	0.0002
Imaging		
MRI-PDFF	1.03 (0.94–1.13)	0.5187

ALT, alanine aminotransferase; AST, aspartate aminotransferase; BMI, body mass index; FIB-4, fibrosis index based on the 4 factors; HbA1c, hemoglobin A1c; HDL, high-density lipoprotein; HOMA-IR, homeostatic model assessment method for insulin resistance [calculated as (fasting insulin (μ U/ml) × fasting glucose (mmol/l)]/22.5]; INR, international normalized ratio; LDL, low-density lipoprotein; MRI-PDFF, MRI-based proton density fat fraction. Statistically significant *p* values are denoted in bold.

MRE (<4.2 kPa) and platelet count (>180,000/ µl) can rule out varices but did not report on other liver-related outcomes or conduct longitudinal analysis for incident events.⁴⁷ Han *et al.*¹¹ also evaluated the cross-sectional association between

MRE and cirrhosis as well as decompensation and identified an optimal cut point of 6.48 kPa with moderate diagnostic accuracy, c=0.71. Finally, Gidener *et al.*¹² performed a longitudinal study evaluating the ability to predict future

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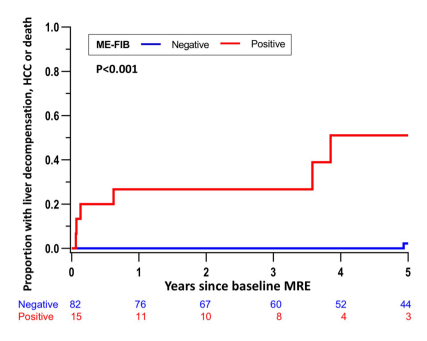


Figure 2. Five-year cumulative incidence of hepatic decompensation, hepatocellular carcinoma, or death by MEFIB index, with positive defined as a combination of MRE \ge 3.3 kPa and FIB-4 \ge 1.6.

cirrhosis development among noncirrhotic patients and the ability to predict hepatic decompensation among 194 patients with compensated NAFLD cirrhosis with longitudinal follow-up. While the authors report only a small increase in diagnostic accuracy in models combining FIB-4 and liver stiffness on MRE, the article did not evaluate the combination of the two tests to maximize sensitivity or specificity.

In this study, we evaluate prevalent liver-related outcomes and evaluate the incidence rate for future outcomes in a subset of patients with longitudinal follow-up without decompensation at baseline. Importantly, we explored the previously validated MEFIB, which had demonstrated high PPV for significant fibrosis and, in this study, demonstrated a high NPV for liver-related events in those with a negative score. The risk of incident decompensation was highest in those with MRE > 5 kPa with a stepwise increase for all hepatic decompensation except HCC, which had a similar risk in those with MRE = 3.63-5 kPa and those with MRE > 5 kPa. If validated, this finding may have implications for future thresholds for HCC screening guided by NITs. Importantly, two incident events did occur in the low-risk group approximately 5 and 8 years after MRE

assessment supporting the need for ongoing follow-up in at-risk populations.

Strengths and limitations

Although the study evaluated a diverse, wellcharacterized cohort, certain limitations are acknowledged. Importantly, the cohort was single center and only a subset of patients had longitudinal follow-up with a limited number of events. However, the association between MRE and incident outcomes has been limited with only one other single center study published to date.¹² While our findings corroborate the prior study, we also evaluate a previously validated index: the MEFIB with practical implications for how a negative score could be used to effectively exclude liver-related events or death (NPV 98%). Furthermore, the study was retrospective; however, our study participants all had a standardized research visit at baseline to systematically evaluate for other causes of liver disease including alcohol, whereas previous studies have relied completely on chart review. Patients with NAFLD are at risk of extrahepatic events including major adverse cardiovascular events, which we did not capture. However, all-cause mortality among those without hepatic decompensation or HCC was low at 3 deaths and represented 21% of deaths in the study cohort. Finally, the study did not assess the impact of change in MRE value on liver-related outcomes, and patients may have participated in clinical trials during the study period affecting their risk of future outcomes. However, there is no US Food and Drug Administration (FDA)-approved treatment for NAFLD, and the impact of an effective treatment in a clinical trial would only attenuate the association between a single MRE and outcomes. Future, multicenter, prospective studies will be required to corroborate the findings and the association between liver stiffness on MRE and liverrelated events and death.

Implications for future research

This study corroborates the importance a single LSM on MRE; however, future longitudinal studies of the implications of changes in liver stiffness over time are needed. In this study, no liver-related events happened within 4 years of a negative MEFIB index. Larger studies or metaanalysis may be required to better define the best interval for follow-up after a low liver stiffness value on MRE alone and in combination with other NITs. Future studies may consider evaluating other combinations of blood-based tests with MRE, including the Enhanced Liver Fibrosis (ELF) score, which has demonstrated an association with NAFLD progression.48 Importantly, while the risk of portal hypertensive complications increased with higher liver stiffness, the risk of HCC appeared to plateau with LSM \ge 3.63 kPa, and future studies should evaluate how to incorporate liver stiffness on MRE into identifying high-risk NAFLD populations for HCC screening who do not have biopsy-proven cirrhosis. In conclusion, liver stiffness on MRE has a strong association with future liver-related events and death, and the combination of MRE and FIB-4 using the MEFIB threshold demonstrated high accuracy to exclude liver-related events or death.

Author contributions

Veeral Ajmera: Conceptualization; Data curation; Formal analysis; Investigation; Methodology; Supervision; Validation; Writing – original draft; Writing – review & editing.

Khang Nguyen: Data curation; Writing – review & editing.

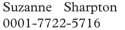
Nobuharu Tamaki: Data curation; Writing – review & editing.

Suzanne Sharpton: Writing – review & editing.

Ricki Bettencourt: Data curation; Formal analysis; Writing – review & editing.

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Conflict of interest statement

The authors declared the following potential conflicts of interest with respect to the research, authorship, and/or publication of this article: Dr Loomba serves as a consultant or advisory board member for Bird Rock Bio, Celgene, Enanta, GRI Bio, Madrigal, Metacrine, NGM, Sanofi, Arrowhead Research, Galmed, GNI, Novo Nordisk, Merck, Siemens, Pfizer, Gilead, and Glympse Bio. In addition, his institution has received grant support from Allergan, BMS, BI, Daiichi-Sankyo Inc., Eli-Lilly, Galectin, Galmed, GE, Genfit, Intercept, Janssen Inc, Madrigal, Merck, NGM, Pfizer, Prometheus, Siemens, and Sirius. He is also the co-founder of Liponexus Inc.

Supplemental material

Supplemental material for this article is available online.

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