Title: Agreement Between Magnetic Resonance Imaging Proton Density Fat

Fraction Measurements and Pathologist-assigned Steatosis Grades of Liver

Biopsies from Adults with Nonalcoholic Steatohepatitis

Short title: Agreement of MRI PDFF with histology

Article type: Original article

Authors:

Michael S. Middleton, MD PhD¹

Elhamy R. Heba, MD¹

Catherine A. Hooker, BSc¹

Mustafa R. Bashir, MD²

Kathryn J. Fowler, MD³

Kumar Sandrasegaran, MD⁴

Elizabeth M. Brunt, MD³

David E. Kleiner, MD PhD⁵

Edward Doo, MD⁶

Mark L. Van Natta, MHS⁷

James Tonascia, PhD⁷

Joel E. Lavine, MD, PhD⁸

Brent A. Neuschwander-Tetri, MD³

Arun Sanyal, MD⁹

Rohit Loomba, MD, MHSc¹⁰

Claude B. Sirlin, MD¹

for the NASH Clinical Research Network

Affiliations:

- 1 Department of Radiology, UCSD School of Medicine, San Diego, California
- 2 Department of Radiology, Duke University Medical Center, 3808, Durham, North Carolina
- 3 Saint Louis University School of Medicine, St. Louis, Missouri
- 4 Department of Radiology, Indiana University School of Medicine, Indianapolis, Indiana
- 5 Laboratory of Pathology, National Cancer Institute
- 6 Liver Diseases Section, Digestive Diseases Branch, National Institute of Diabetes and Digestive and Kidney Diseases
- 7 Johns Hopkins Bloomberg School of Public Health, Baltimore, Maryland
- 8 Division of Pediatric Gastroenterology, Hepatology and Nutrition, Department of Pediatrics,
 Columbia University Medical Center, New York, New York
- 9 Virginia Commonwealth University School of Medicine, Richmond, Virginia
- 10 NAFLD Translational Research Unit, Division of Gastroenterology, UCSD School of Medicine, San Diego, California

Acknowledgement

We would like to acknowledge support from the following sources: NIDDK U01 DK061718, U01 DK061728, U01 DK061731, U01 DK061732, U01 DK061734, U01 DK061737, U01 DK061738, U01 DK061730, and U01 DK061713; NCATS UL1 TR000439, UL1 TR000436,

UL1 TR000006, UL1 TR000448, UL1 TR000100, UL1 TR000004, UL1 TR000423, and UL1 TR000058); NCI.

Abbreviations:

AUROC area under receiving operating curve

BMI body mass index

CAP NAFLD activity score

CI confidence interval

CRN clinical research network

DCC Data Coordinating Committee

DICOM Digital Imaging and Communications in Medicine

EOT end of treatment

FLINT Farnesoid X Receptor Ligand Obeticholic Acid in NASH Treatment Trial

GRE gradient-recalled echo

H&E Hematoxylin and eosin

MRI magnetic resonance imaging

MRS magnetic resonance spectroscopy

NAS NAFLD activity score

NAFLD nonalcoholic fatty liver disease

NASH nonalcoholic steatohepatitis

NPV negative predictive value

PDFF proton density fat fraction

PPD positive predictive value

ROC receiver operating characteristic

SD standard deviation

RCC Radiology Coordinating Center

ROI region of interest

TE time to echo

TR repeat time

Corresponding author contact information:

Michael S. Middleton, MD PhD

Liver Imaging Group, Department of Radiology

UCSD School of Medicine

408 Dickinson Street

San Diego, CA 92103-8226

msm@ucsd.edu

Disclosures:

Michael S. Middleton, MD PhD

Alexion (Contracted work through university,

through Biomedical Systems as CRO)

AstraZeneca (Contracted work through university,

through Profil as CRO)

Bioclinica (Contracted work through university;

Genzyme/Sanofi as sponsor)

Biomedical Systems (Contracted work through

university)

Bracco (Consultant)

Bristol-Myers Squibb (Contracted work through university)

Celgene (CDA to discuss contracted work through university)

Galmed (Contracted work through university)

Genentech (CDA to discuss contracted work through university)

General Electric (Stockholder, contracted work through university, grant)

Genzyme (Contracted work through university)

Gilead (Grant, and contracted work through university)

Guerbet (Grant)

Icon (CDA to discuss contracted work through university)

Intercept Pharmaceuticals (Contracted work through university)

Isis (Contracted work through university)

Janssen (Contracted work through university)

Merge Healthcare (Consultant)

NuSirt (Contracted work through university)

Perspectum (CDA to discuss contracted work through university)

Pfizer (Contracted work through university, stockholder)

Profil (Contracted work through university, for sponsor AstraZeneca)

Quantitative Insights (Consultant)

Sanofi (Contracted work through university)

Shire (CDA with Icon, with Shire as sponsor, to discuss contracted work through university)

Siemens (Contracted work through university)

Synageva (Contracted work through university,

through Biomedical Systems as CRO)

Takeda (Contracted work through university)

Virtualscopics (Contracted work through university)

Zydus (Discussions leading to CDA to discuss contracted work through university)

Elhamy R. Heba, MD no COI

Catherine A. Hooker, BSc no COI

Mustafa Bashir, MD RadMD (Consultant)

Siemens Healthcare (Research grant)

NGM Biopharmaceuticals (Research grant)

TaiwanJ Pharmaceuticals (Research grant)

Kathryn J. Fowler, MD no COI

Kumar Sandrasegaran, MD no COI

Elizabeth M. Brunt, MD no COI

David E. Kleiner, MD PhD no COI

Edward Doo, MD no COI

Mark L. Van Natta, MHS no COI

James Tonascia, PhD no COI

Joel E. Lavine, MD, PhD no COI

Brent A. Neuschwander-Tetri, MD Nimbus Therapeutics (Consultant)

Enanta (Consultant)

Novartis (Consultant)

Galmed (Consultant)

Zafgen (Consultant)

Receptos (Consultant)

Pfizer (Consultant)

Allergan (Consultant)

MedImmune/AstraZeneca (Consultant)

Arun Sanyal, MD AbbVie (Consultant)

Astra Zeneca (Institutional grant support)

Boehringer Ingelhiem (Consultant)

Bristol-Myers Squibb (Institutional grant support)

Astra Zeneca (Consultant)

Echosens (Unpaid consultant)

Eli Lilly (Unpaid consultant)

Exalenz Bioscience (Consultant)

FibroGen (Consultant)

Genfit (Stock options, consultant)

Gilead (Institutional grant support)

Hemoshear (Consultant)

Immune Pharma (Consultant)

Immuron (Consultant)

Intercept Pharmaceuticals (Unpaid consultant)

Merck (Institutional grant support)

Nimbus Therapeutics (Consultant)

Nitto Denko (Consultant)

Novartis (Institutional grant support)

Pfizer (Consultant)

Salix Pharmaceuticals (Consultant, institutional

grant support)

Sanyal Biotechnologies (President)

Takeda (Consultant)

Tobira Therapeutics (Consultant, institutional grant

support)

Rohit Loomba, MD, MHSc Adheron (Research funding)

Alnylam (Consultant)

Arisaph (Research funding)

Arrowhead Research (Advisory board)

Boehringer Ingelheim (Consultant)

Bristol-Myers Squibb (Research funding, advisory

board, consultant)

Celgene (Consultant)

CNI (Consultant)

Conatus (Advisory board, consultant)

Daiichi-Sankyo Inc (Research funding)

DeuteRx (Consultant)

Eli Lilly (Consultant)

Enanta (Consultant)

Fibrogen (Consultant)

Galmed (Research funding, advisory board)

General Electric (Research funding)

Gilead (Research funding, advisory board,

consultant)

Immuron (Research funding)

Intercept Pharmaceuticals (Clinical trial steering

committee)

Isis (Consultant)

Janssen (Consultant)

Kinemed (Research funding)

Liponexus Inc (Co-founder)

Merck (Research funding, consultant)

Metacrine (Consultant)

MS (Research funding, advisory board, consultant)

NGM Biopharmaceuticals (Research funding,

consultant)

Nimbus (Advisory board)

Pfizer (Consultant)

Promedior (Research funding)

Receptos (Consultant)

RuiYi (Consultant)

Scholar Rock (Consultant)

Shire (Consultant)

Siemens (Research funding)

Tobira (Research funding, advisory board)

Viking (Consultant)

Zafgen (Consultant)

Alexion (Contracted work through university,

(through Biomedical Systems as CRO)

AstraZeneca (Contracted work through university,

through Profil as CRO)

Bioclinica (Contracted work through university;

Genzyme/Sanofi as sponsor)

Biomedical Systems (Contracted work through

Claude B. Sirlin, MD

university)

Bracco (Consultant)

Bristol-Myers Squibb (Contracted work through university)

Fibrogen (Consultant)

Galmed (Contracted work through university)

General Electric (Contracted through university, grant)

Genzyme (Contracted work through university)

Gilead (Contracted work through university)

Guerbet (Grant)

Icon (CDA to discuss contracted work through university)

Intercept Pharmaceuticals (CDA to discuss contracted work through university)

Isis (Contracted work through university)

Janssen (Contracted work through university)

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discuss contracted work through university)

Siemens (Contracted work through university)

Synageva (Contracted work through university,
through Biomedical Systems as CRO)

Takeda (Contracted work through university)

Tobira (Consultant)

Virtualscopics (Contracted work through university)
```

Author contributions:

```
Michael S. Middleton, MD PhD

study concept and design

acquisition of data

analysis and interpretation of data

drafting of manuscript

critical revision of manuscript for important intellectual content

study supervision

Elhamy R. Heba, MD

analysis and interpretation of data

drafting of manuscript

critical revision of manuscript for important intellectual content
```

```
Catherine A. Hooker, BSc
       analysis and interpretation of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
Mustafa Bashir, MD
       acquisition of data
       analysis and interpretation of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
Kathryn J. Fowler, MD
       acquisition of data
       analysis and interpretation of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
Kumar Sandrasegaran, MD
       acquisition of data
       analysis and interpretation of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
Elizabeth M. Brunt, MD
       analysis and interpretation of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
```

```
David E. Kleiner, MD PhD
       analysis and interpretation of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
Edward Doo, MD
       study concept and design
       obtained funding
       drafting of manuscript
       critical revision of manuscript for important intellectual content
Mark L. Van Natta, MHS
       analysis and interpretation of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
       statistical analysis
James Tonascia, PhD
       study concept and design
       analysis and interpretation of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
       study supervision
       obtaining funding
       statistical analysis
Joel E. Lavine, MD, PhD
```

```
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       drafting of manuscript
       critical revision of manuscript for important intellectual content
       obtaining funding
Brent A. Neuschwander-Tetri, MD
       acquisition of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
Arun Sanyal, MD
       acquisition of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
Rohit Loomba, MD, MHSc
       study concept and design
       acquisition of data
       analysis and interpretation of data
       drafting of manuscript
       critical revision of manuscript for important intellectual content
       study supervision
Claude B. Sirlin, MD
       study concept and design
       acquisition of data
       analysis and interpretation of data
```

drafting of manuscript

critical revision of manuscript for important intellectual content

study supervision

obtaining funding

for the NASH Clinical Research Network

study concept and design

technical support

administrative support

ABSTRACT

Background & Aims: We assessed the diagnostic performance of magnetic resonance imaging (MRI) proton density fat fraction (PDFF) in grading hepatic steatosis and change in hepatic steatosis in adults with nonalcoholic steatohepatitis (NASH) in a multi-center study, using central histology as reference.

Methods: We collected data from 113 adults with NASH participating in a multi-center, randomized, double-masked, placebo-controlled, phase 2b trial to compare the efficacy cross-sectionally and longitudinally of obeticholic acid vs placebo. Hepatic steatosis was assessed at baseline and after 72 weeks of obeticholic acid or placebo by liver biopsy and MRI (scanners from different manufacturers, at 1.5T or 3T). We compared steatosis estimates by PDFF vs histology. Histologic steatosis grade was scored in consensus by a pathology committee. Cross-validated receiver operating characteristic (ROC) analyses were performed.

Results: At baseline, 34% of subjects had steatosis grade 0 or 1, 39% had steatosis grade 2, and 27% had steatosis grade 3; corresponding mean PDFF values were 9.8%±3.7%, 18.1%±4.3%, and 30.1%±8.1%. PDFF classified steatosis grade 0–1 vs 2–3 with an area under the ROC curve (AUROC) of 0.95 (95% CI, 0.91–0.98), and grade 0–2 vs grade 3 steatosis with an AUROC of 0.96 (95% CI, 0.93–0.99). PDFF cut-off values at 90% specificity were 16.3% for grades 2–3 and 21.7% for grade 3, with corresponding sensitivities of 83% and 84%. After 72 weeks' of obeticholic vs. placebo, 42% of subjects had a reduced steatosis grade (mean reduction in PDFF from baseline of 7.4%±8.7%), 49% had no change in steatosis grade (mean increase in PDFF from baseline of 0.3%±6.3%), and 9% had an increased steatosis grade (mean increase in PDFF

from baseline of 7.7%±6.0%). PDFF change identified subjects with reduced steatosis grade with an AUROC of 0.81 (95% CI, 0.71–0.91) and increased steatosis grade with an AUROC of 0.81 (95% CI, 0.63–0.99). A PDFF reduction of 5.15% identified subjects with reduced steatosis grade with 90% specificity and 58% sensitivity, whereas a PDFF increase of 5.6% identified those with increased steatosis grade with 90% specificity and 57% sensitivity.

Conclusions: Based on data from a phase 2 randomized controlled trial of adults with NASH, PDFF estimated by MRI scanners of different field strength and at different sites, accurately classifies grades and changes in hepatic steatosis when histologic analysis of biopsies is used as a reference.

KEY WORDS: FLINT, NAFLD, direct comparison, non-invasive

INTRODUCTION

Nonalcoholic fatty liver disease (NAFLD) is estimated to affect a billion people in the world today [1,2]. Reported population-based prevalence estimates of nonalcoholic steatohepatitis (NASH), in which hepatocellular injury and inflammation are present in addition to steatosis, have ranged from 1.1% to 18.5% [3], with even larger percentages for higher-risk subpopulations [4,5]. Long-term outcomes of patients diagnosed with NASH include diabetes, cardiovascular disease, cirrhosis, hepatocellular carcinoma, and death [6,7].

Magnetic resonance imaging (MRI)-estimated proton density fat fraction (PDFF) is a non-invasive biomarker of hepatic steatosis, which is a key feature of NAFLD and NASH [8]. In single-site cross-sectional studies, PDFF has been shown to be accurate using magnetic resonance spectroscopy (MRS) as a reference standard [9,10,11,12], and to be reproducible across scanner manufacturers and field strengths [13,14]. Moreover, other single-center crosssectional studies using liver biopsy as reference have shown that PDFF correlates with hepatic steatosis grade [15,16,17,18,19], and accurately classifies dichotomized steatosis grade with areas under receiver operating characteristic (ROC) curves ranging from 0.825 to 0.989 [15,20]. The performance of PDFF as a biomarker for hepatic steatosis has also been investigated in single-center longitudinal treatment trials. In the MOZART trial (35 subjects, diagnosed NASH, treated with ezetimibe vs placebo for 24 weeks), histologic responders (reduction in NAFLD Activity Score [NAS] of two or more points) had greater relative PDFF reduction (29.3%) than non-responders (2.0%) [21]. In another randomized trial (34 subjects, diagnosed NASH, treated with N-3 fish oil vs placebo for one year), subjects treated with N3 lost more weight and showed a greater decrease in PDFF than those treated with placebo, although there were no differences in histologic response [22]. Until now, however, diagnostic performance of PDFF has not been

investigated in a well-controlled, multi-center treatment trial. Assessing multi-center diagnostic performance is necessary to further validate PDFF prior to its widespread adoption and implementation [20].

Therefore, the purpose of this study was to assess cross-sectional and longitudinal diagnostic performance of PDFF to grade hepatic steatosis in adults using centrally-scored histology as reference. To do this, we performed a prospectively-designed study of PDFF estimated by MRI in the Farnesoid X Receptor Ligand Obeticholic Acid in NASH Treatment (FLINT) trial [23]. The FLINT trial was a multi-center, randomized, double-masked, placebo-controlled, phase 2b clinical trial of treatment with either obeticholic acid or placebo in patients with NASH. Hepatic steatosis was assessed at baseline and after 72 weeks of obeticholic acid vs placebo by both liver biopsy and MRI, allowing comparison of PDFF to histologic hepatic steatosis grade in a well-characterized, longitudinal multi-center study.

MATERIALS AND METHODS

Study design

MRI was offered at baseline and end-of-treatment (EOT) to adults participating in the FLINT trial. PDFF was a secondary imaging endpoint, for which histologic hepatic steatosis grade served as the reference standard.

Eligibility criteria for the FLINT trial are published elsewhere [23]; all subjects were diagnosed with NASH based on local histology review of a standard-of-care liver biopsy. Inclusion criteria for the liver MRI portion of the FLINT trial were that the subject was enrolled, and that the subject was willing and able to complete both MRI exams (the baseline exam prior to randomization and within 90 days of baseline biopsy, and the EOT MRI exam within 90 days of EOT biopsy). MRI exclusion criteria were contraindication to MRI, extreme claustrophobia,

pregnancy or trying to become pregnant, weight or girth exceeding MRI scanner capability, or any condition or circumstance that, in the opinion of the clinical trial site investigator would interfere with completion of the MRI exam.

The FLINT trial, including the MRI portion, was approved by an Institutional Review Board at each participating clinical trial site, and was in compliance with the Health Insurance Portability and Accountability Act. All subjects enrolled in the FLINT trial signed informed consent for MRI exams.

All authors had access to the study data and have reviewed and approved the final manuscript.

MRI clinical trial site qualification

The MRI portion of this study was managed by the NASH CRN Radiology Coordinating Center (RCC) in conjunction with the NASH CRN Data Coordinating Center (DCC). The RCC ensured that each participating clinical trial site was qualified to participate in the MRI portion of the FLINT trial based on submission of technically adequate phantom or volunteer MRI data, and provided central imaging quality control.

MRI exams

Seven of the eight participating FLINT clinical trial sites contributed MRI data to this study using 1.5T (three sites) and 3T (four sites) MR scanners (**Table 1**). At each clinical trial site, only a single scanner was used so that longitudinal instrument-dependent measurement variability would be minimized. PDFF reproducibility has been demonstrated in two studies across different clinical trial sites, and for both field strengths used in this study [**13,14**].

For each MRI exam, an advanced, non-contrast, breath-hold, magnitude-based, gradient-recalled-echo (GRE), two-dimensional axial sequence was obtained of the entire liver using a

torso array coil centered over the liver. T1 effects that could cause subject- or scanner-dependent bias were minimized using a low flip angle (ten degrees) and a repetition time (TR) of \geq 120 ms. Six echoes were collected at alternately out-of-phase and in-phase echo times (TEs) for the water and main fat peaks to permit PDFF quantification while accounting for subject-based T2* effects [9,10,11,12] (Table 1). Images in Digital Imaging and Communications in Medicine (DICOM) format were transferred from clinical trial sites to the RCC for analysis.

Data analysts at the RCC placed one circular 1-cm radius region of interest (ROI) on 5th echo (out-of-phase) images in each of the nine anatomical liver segments; those ROIs were propagated to images for the other echoes. Mean signal intensities at those ROI locations were recorded. A custom MATLABTM (The MathWorks, Natick, MA, USA) multi-peak, non-linear, least-squares fitting algorithm [9,10,11,12] that accounted for T2* signal decay and multi-frequency interference effects of the various proton locations in triglyceride [24] was used to estimate PDFF for each ROI location.

Liver biopsy

MRI analysis

Percutaneous core liver biopsies were performed by NASH CRN investigators as research procedures or for clinical care. Hematoxylin and eosin (H&E), Masson's trichrome, and iron stains were prepared from formalin-fixed tissue specimens. The interval between performance of MRI and biopsy was calculated. Local clinical trial site histological analysis was performed to determine eligibility for FLINT trial enrollment; those local scores were not used in the current analysis.

Comparison of histologic steatosis grade and PDFF

PDFF, and histologic steatosis percentage (and grade) both assess liver fat, but they do so

in very different ways. Histologic steatosis assessment yields a percentage of hepatocytes that show fat globules on H&E stain (from whence is derived a steatosis grade from 0 to 3), whereas PDFF estimates a ratio of observable MRI signal from fat, compared to all observable MRI signal (from both fat and water). Thus, if all examined hepatocytes were seen to be filled with 50% fat globules, the histologic steatosis percentage would be 100% since all cells show fat globules, and the steatosis grade would be 3, but the MRI PDFF percentage would be, ignoring technical factors for the moment, something like 50%. Thus, we see in practice that PDFF percentages typically are somewhat less than half of histologic steatosis percentages.

Central histological analysis

Biopsies were reviewed by the pathologists of the NASH CRN Pathology Committee who provided central histology assessment by consensus at a multi-head microscope. Pathologists were blinded to all clinical information, including participation in this study. Steatosis grade was provided according to the standardized NASH CRN histological NAFLD scoring system (proportion of hepatocytes containing fat macrovesicles: grade 0 for < 5%, grade 1 for 5 to 33%, grade 2 for > 33 to 66%, and grade 3 for > 66% [25]). Other centrally scored features that were evaluated in this study were lobular inflammation, portal inflammation, hepatocellular ballooning, fibrosis stage, and NAFLD Activity Score (NAS). *Blinding*

RCC analysts and other staff at the RCC were blinded to histology results, and pathologists were blinded to imaging results. RCC analysts were blinded to treatment assignment.

Other data

Subject demographics, laboratory, anthropomorphic measurements, and medical history

were collected at each clinical trial site.

Statistical analysis

Statistical analyses were done with SAS (SAS Institute 2011, Base SAS 9·3 Procedures Guide) and Stata (StataCorp 2013, Stata Statistical Software: release 13).

A single PDFF value was calculated for each MRI exam as the mean of the PDFF values for the nine anatomical liver segments. Demographic, histologic and imaging information were summarized with categorical variables expressed as numbers and percentages and continuous variables expressed by mean (± standard deviation [SD]). The proportion of subjects who had, vs did not have MRIs at baseline was compared with regard to treatment group, study site, demographics, liver enzymes, lipids, metabolic factors, co-morbidities, concomitant liver medications, and histology findings. Pearson partial correlation coefficients and 95% confidence intervals were estimated between PDFF and histologic variables (steatosis, lobular inflammation, hepatocellular ballooning, fibrosis, and portal inflammation) at baseline and for changes from baseline to 72 weeks.

Diagnostic accuracy of PDFF to classify hepatic steatosis grade at baseline was tested for grades 0-1 vs 2-3, and grades 0-2 vs 3. Diagnostic accuracy of change in PDFF to classify change in hepatic steatosis grade from baseline to EOT was tested for: reduction vs no change/increase and increase vs no change/decrease. Cross-validated area under ROC (AUROC) curves using a jack-knife procedure and 95% CIs were estimated for each of these dichotomizations [26]. Cut-off PDFF values were estimated using the lowest threshold value for which there was \geq 90% specificity to distinguish between these dichotomized categories. Sensitivity, positive predictive values (PPVs), and negative predictive valves (NPVs) were calculated along with 95% confidence intervals (CIs) fixing specificity at 90%.

RESULTS

Of 283 adults enrolled in the FLINT trial from March 16, 2011 to December 3, 2012 at eight participating FLINT clinical trial sites, 113 (40%) had MRI and liver biopsy at baseline, 85 (30%) had MRI and liver biopsy at EOT, and 78 (28%) had MRI and liver biopsy at both time points. One subject with a baseline MRI but without a centrally-read baseline liver biopsy was excluded from the analysis. All baseline MRIs were performed prior to randomization and occurred an average of 51 days following baseline biopsy. Follow-up MRIs were performed an average of two days after follow-up biopsy, and were performed from 29 days before to 78 days after biopsy.

Cohort baseline population characteristics are summarized in **Table 2**. Subjects who had, vs did not have MRIs had significantly (p < 0.05) lower mean aspartate aminotransferase, alkaline phosphatase, weight, BMI and prevalence of hyperlipidemia and hypertension. Additionally, the 78 subjects with both paired MRIs and paired biopsies vs the 204 subjects without, had lower (p < 0.05) mean AST, ALT and alkaline phosphatase at baseline. $Cross-sectional\ analysis$

The distribution of PDFF at baseline is shown in **Figure 1**: PDFF mean was 18.6%±9.6%, and ranged from 3.7% to 49.3%.

At baseline, 34% (38/113) of biopsies had steatosis grade 0 or 1, 39% (44/113) had grade 2, and 27% (31/113) had grade 3 with corresponding mean PDFF values of 9.8%±3.7%, 18.1%±4.3%, and 30.1%±8.1%, respectively. PDFF values for each histologic steatosis grade at baseline are shown in **Figure 2**. Higher steatosis grade corresponded to higher mean PDFF.

Partial correlation of baseline PDFF values with histologic features is summarized in **Table 3**. PDFF is highly correlated with steatosis controlling for other histologic features

(Pearson Partial Correlation = 0.80 [95% CI, 0.73-0.86]; p < 0.001).

Diagnostic accuracy of PDFF for classifying steatosis is summarized in **Table 4**. The area under ROC (AUROC) from logistic regression using PDFF as a surrogate for classifying steatosis grade 0-1 vs 2-3 was 0.95 (95% CI, 0.91–0.98) and for classifying steatosis grade 0-2 vs 3 was 0.96 (95 % CI, 0.93–0.99). PDFF cut-off values at 90% specificity were 16.3% for grades 2-3, and 21.7% for grade 3 discrimination. Diagnostic accuracy of PDFF for classifying steatosis was similar between the 1.5T vs. 3T scanners (data not shown).

Longitudinal analysis

PDFF values for each histologic hepatic steatosis grade change category (reduction, no change, and increase in steatosis grade) are shown in the box plots in **Figure 3**. Changes in PDFF were positive in 71% (5/7) of subjects in whom steatosis grade increased and were negative in of subjects in 91% (30/33) of subjects in whom steatosis grade decreased. The mean (SD) change in MRI-PDFF in 38 subjects with no change in histology grade category was 0.3%±6.3%, ranging from -21.1 to +21.0%.

Correlation of change from baseline to EOT in PDFF values with change from baseline to EOT in histologic features is summarized in **Table 3**. Correlation between change in PDFF and change in histologic steatosis grade controlling for other histologic features was high (Pearson Partial Correlation = 0.63 [95% CI, 0.47-0.75]; p < 0.001).

Diagnostic accuracy of change in PDFF vs change in histologic hepatic steatosis grade from baseline to EOT is summarized in **Table 4**. At EOT, 42% of paired biopsies had improvement in steatosis grade, 49% had no change, and 9% had worsening. The AUROC using PDFF change to classify histologic hepatic steatosis grade improvement and worsening, respectively, were 0.81 (95% CI, 0.71–0.91) and 0.81 (95% CI, 0.63–0.99). Cut-off values for

PDFF change at 90% specificity were -5.1% for improvement and +5.6% for worsening hepatic steatosis grade.

DISCUSSION

We compared PDFF to histologic hepatic steatosis grade at baseline, and change in PDFF to change in histologic hepatic steatosis grade from baseline to EOT. We found that PDFF correlates well with histologic hepatic steatosis grade cross-sectionally, and that change in PDFF correlates well with change in histologic hepatic steatosis grade longitudinally.

ROC analysis at baseline yielded PDFF cutoffs predicting histologic hepatic steatosis grade that are similar to those reported in the literature for a single-center study. We found that at 90% specificity, PDFF cut-off values were 16.3% (83% sensitivity) and 21.7% (84% sensitivity) for discrimination of histologic steatosis grades 0-1 from 2-3, and grades 0-2 from 3, respectively. Tang et al [15] reported PDFF cut-off values for those discriminations in their single-center study of 77 children and adults with known or suspected NAFLD of 17.4% (64% sensitivity and 96% specificity) and 22.1% (71% sensitivity and 92% specificity), respectively. In a subsequent single-center study of 89 adults, Tang et al [27] subsequently reported high sensitivity and specificity for those cutoffs to distinguish steatosis grades in another clinical trial. As emphasized previously by others [28], PDFF is a quantitative marker of MRI-visible fat content while histology scoring is a semi-quantitative marker of proportion of steatotic hepatocytes. Since PDFF and histology do not attempt to measure the same quantity, perfect agreement is not expected.

The AUROC of PDFF for classifying longitudinal change in steatosis is 0.81, which is inferior to its cross-sectional performance at baseline. The lower AUROC for change may reflect the 'noisiness' of histologic grading as a marker of steatosis, an effect which is amplified when differences in PDFF are compared to differences in histologic steatosis grade.

Hepatic steatosis may also be assessed non-invasively by CAP (a transient elastography-derived controlled attenuation parameter), which is an estimate of total ultrasound attenuation [29]. Although we did not perform a direct comparison of CAP and PDFF, the accuracy of MRI to estimate PDFF has been reported to be higher than that of CAP, using histology as reference standard [30 31].

Strengths of this study are that the cohort of patients that was studied was well-characterized, with a racial/ethnic makeup representative of patients diagnosed with NASH in the United States; that subjects enrolled in this study had both pre- and post-treatment biopsy; that histology was scored centrally by the NASH CRN Pathology Committee; and that a range of MRI scanner manufacturers at two field strengths and across multiple study sites were utilized. Thus, our study results are likely to be generalizable to the entire adult NASH population in the United States, while also providing compelling evidence that MRI-PDFF can be applied successfully in multi-center trials.

A major limitation of this study was that not all subjects enrolled in the FLINT trial had MRIs. Only 40% of randomized subjects had an MRI at baseline and only 28% had paired EOT and baseline MRIs. Participation in the MRI study was reduced because the MRI qualification process at each clinic was completed after the start of enrollment into the trial. Participation rates into the MRI study varied widely at the nine study sites, ranging from 0 to 84%. In addition, subjects who had the MRI weighed significantly less than those who did not participate; this may be because the heaviest FLINT participants were too large to undergo MRI. Also, significantly lower mean alanine aminotransferase and aspartate aminotransferase and lower rates of hyperlipidemia and hypertension were seen at baseline in those who did vs did not participate in the MRI study suggesting a less severe group in those who participated. Our results could be

biased if the relationships between MRI-PDFF and histologic features varied in those who participated vs did not participate in the MRI study.

Another limitation was that relatively few subjects of non-White race were included. This was unavoidable since NASH is relatively uncommon in non-Whites, and the racial and ethnic diversity of our study population reflects that seen in the national prevalence of NASH. Our study also did not include children, as the FLINT trial enrolled only adults; thus multi-center validation studies of MRI PDFF in children are needed. Finally, potentially confounding factors such as those in **Table 2** were not investigated because the study was not powered to permit those investigations. It is also not likely that there are meaningful confounders to our results, given that none have been reported for single-site studies [14,15].

Steatosis grade was determined in this study according to the standardized NASH CRN histological NAFLD scoring system based on the proportion of hepatocytes containing fat macrovesicles. The performance of MRI PDFF compared to microvesicular steatosis remains unclear, especially since microvesicular steatosis is often under-staged. The relationship of microvesicular steatosis and PDFF was not examined in this paper because all work to date validating MRI PDFF against MR spectroscopy and histology has considered only macrovesicular steatosis, and because microvesicular steatosis is an uncommon finding in NASH tending to be seen in the more severe cases [32].

Finally, since an inclusion criterion for the FLINT trial was the presence of NASH as assessed by a local pathologist, only one subject had histologic hepatic steatosis grade 0 at central reading. Hence, grades 0 and 1 were considered as a single category in our analyses. Further multi-center studies in populations including subjects with lower amounts of hepatic

steatosis (i.e., grades 0 and 1) are needed to investigate correlations of PDFF with hepatic steatosis in that range.

In conclusion, in a well-controlled, multi-center study in adults, PDFF showed high agreement with histologic hepatic steatosis grade based on MRI data at sites using scanners from different manufacturers and of different field strengths. Importantly, change in PDFF accurately classified change in EOT histologic hepatic steatosis grade. These findings support wider use of PDFF in multi-center trials in adults as a biomarker for hepatic steatosis at baseline, and for change in hepatic steatosis with treatment.

FIGURE CAPTIONS

- **Figure 1.** PDFF distribution of study population at baseline.
- Bar and whisker plot at baseline of PDFF vs steatosis grade. For each steatosis grade, maximum and minimum PDFF values are indicated by horizontal lines at the bottom and top of each steatosis grade entry, the darkened box in the middle represents PDFF data points in the 25% to 75% interquartile range, and the line through the middle of the central darkened box represents the median PDFF value.
- Bar and whisker plot of change in PDFF vs change in histologic hepatic steatosis grade. Mean PDFF change values for steatosis grade reduction, no change in steatosis grade, and increase in steatosis grade were, respectively: -7.4% ± 8.7% (n=42), 0.3% ±6.3% (n=49), and 7.7% ± 6.0% (n=9). Explanations of each barand-whisker plot entry are as noted for Figure 2, except that in this figure values represent PDFF change, the maximum and minimum PDFF change values that are indicated by horizontal lines at the bottom and top of each steatosis grade entry exclude outliers, and the isolated dots represent outliers.

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TABLES

 Table 1.
 MRI scanners and techniques

Clinical trial site MR scanners	1.5T	3T
Site A	-	Siemens TIM Trio
Site B	-	Siemens TIM Trio
Site C	Siemens Avanto	- 7
Site D	-	GE Signa HDxt
Site E	-	GE Signa HDxt
Site F	Siemens Avanto) -
Site G	Siemens TIM Symphony	-
	(7
Scanning parameters		
Repetition time (ms)	≥ 120	≥ 120
First time-to-echo (ms)	2.3	1.15
Delta time-to-echo (ms)	2.3	1.15
Number of echoes	6	6
Flip angle (°)	10	10
Bandwidth (Hz/Px)	≥ 500	≥ 1,000
Slice thickness (mm)	8 or 10	8 or 10
Slice gap (mm)	0	0
Phase encoding steps	192	128
Frequency encoding steps	192	128

 Table 2.
 Baseline characteristics of those obtaining vs not obtaining baseline MRI

	Baseline MRI (n=113)	No Baseline MRI (n=169)	p-value
Study site* – no.	1		
Site A	14	15	7
Site B	15	12	
Site C	17	10	
Site D	32	6	
Site E	7	21	
Site F	9	33	
Site G	19	18	
Treatment group			
Obeticholic acid – no. (%)	55 (48.7)	86 (50.9)	0.81
Demographics			
Age (yrs)	51±11	51±11	0.98
Male sex - no. (%)	43 (38.0)	53 (31.2)	0.25
White race - no. (%)	93 (82.3)	140 (82.8)	1.00
Hispanic ethnicity - no. (%)	19 (16.8)	24 (14.2)	0.61
Liver enzymes			
Aspartate aminotransferase - U/L	55±32	65±39	0.03
Alanine aminotransferase – U/L	76±43	87±54	0.08
γ-Glutamyltransferase – U/L	80±115	75±71	0.71
Alkaline phosphatase - U/L	78±26	85±27	0.03
Total bilirubin – mg/dL	0.6±0.3	0.7±0.4	0.49
Metabolic factors			
Weight - kg	95±17	100±23	0.03
Body-mass index - kg/m ²	33.6±5.2	35.3±6.9	0.02
Waist circumference - cm	109±12	112±16	0.16
Waist to hip ratio	0.95±0.07	0.95±0.09	0.99
Fasting serum glucose - mg/dL	113±31	118±40	0.19
Insulin - umol/mL	28±36	28±27	0.94
HOMA-IR - (mg/dL * umol/mL) / 405	8.5±12.1	8.3±8.7	0.90
Hemoglobin A1c - %	6.3±1.0	6.6±1.1	0.06
Systolic blood pressure - mmHg	130±16	133±16	0.15
Diastolic blood pressure - mmHg	78±11	77±10	0.64
Comorbidities			
Hyperlipidemia - no. (%)	58 (51.3)	115 (68.0)	0.006
Hypertension - no. (%)	59 (52.2)	112 (66.3)	0.02
Diabetes - no. (%)	57 (50.4)	92 (54.4)	0.54
Cardiovascular disease - no. (%)	4 (3.5)	11 (6.5)	0.42
Liver histology findings			
Definite steatohepatitis - no. (%)	91 (80.5)	134 (79.3)	0.88

Fibrosis - stage ¹	1.7±1.1	1.9±1.0	0.23
NAFLD activity score (NAS) ²	5.2±1.3	5.2±1.3	0.62
Hepatocellular ballooning - score	1.4±0.7	1.4 ± 0.8	0.88
Steatosis - score	1.9±0.8	2.1±0.8	0.08
Lobular inflammation - score	1.8±0.7	1.8 ± 0.7	0.34
Portal inflammation - score ³	1.1±0.6	1.2±0.6	0.86

Notes: Plus-minus values are means±SD; BMI = body mass index; PDFF = proton density fat fraction; SD = standard deviation.

- * Two study sites with combined 55 randomized subjects did not obtain MRIs
- Fibrosis was assessed on a scale of 0 to 4, with higher scores indicating more severe fibrosis
- 2 NAS was assessed on a scale of 0 to 8, with higher scores indicating more severe disease; the components of this measure are steatosis (assessed on a scale of 0 to 3), lobular inflammation (assessed on a scale of 0 to 3), and hepatocellular ballooning (assessed on a scale of 0 to 2)
- Portal inflammation was assessed on a scale of 0 to 2 with higher scores indicating more severe inflammation

Table 3. Partial correlation of MRI-PDFF with histologic components at baseline and change in MRI-PDFF with change in histologic components at EOT

Histologic component	MRI-PDFF at baseline (n=113)			Change in MRI-PDFF at EOT (n=78)		
	r	95% CI	p-value	r	95% CI	p-value
Steatosis	0.80	0.80-0.90	< 0.001	0.63	0.47-0.75	< 0.001
Lobular inflammation	-0.21	-0.28-0.09	0.03	-0.14	-0.36-0.09	0.23
Ballooning	-0.02	-0.510.19	0.85	0.17	-0.06-0.39	0.14
Fibrosis	-0.11	-0.470.14	0.24	-0.18	-0.39–0.05	0.13
Portal inflammation	0.02	-0.30-0.07	0.84	-0.13	-0.35-0.10	0.26

Notes: MRI = magnetic resonance imaging; PDFF = proton density fat fraction; EOT = end of treatment

 Table 4.
 Diagnostic accuracy of PDFF for classifying steatosis

Cross- sectional steatosis grade classification (n=113)	(%) a	F threshold at 90% ificity	Sensitivity	PPV	NPV	AUROC (95% CI)
0-1 vs 2-3	10	5.3	83%	95%	73%	0.95 (0.91–
			(62/75)	(62/65)	(35/48)	0.98)
0-2 vs 3	2	1.7	84%	76%	94%	0.96 (0.93-
			(26/31)	(26/34)	(74/79)	0.99)
					, ,	
Longitudinal steatosis grade change classification (n=78)	Mean (SD) PDFF change (%)	Cutoff PDFF change at 90% specificity (%)	Sensitivity	PPV	NPV	AUROC (95% CI)
Decrease in steatosis grade (improvement) (42%)	-7.4±8.7	- 5.1	58% (19/33)	83% (19/23)	75% (41/55)	0.81 (0.71– 0.91)
No change in steatosis grade (49%)	0.3±6.3	-		-	-	-
Increase in steatosis grade (worsening) (9%)	7.7±6.0	5.6	57% (4/7)	36% (4/11)	96% (64/67)	0.81 (0.63– 0.99)

Notes: PDFF = proton density fat fraction; MRI = magnetic resonance imaging; PPV = positive predictive value; NPV = negative predictive value; AUROC = area under ROC curve; CI = confidence interval





