Improvements of Fibrosis and Disease Activity Are Associated With Improvement of Patient-Reported Outcomes in Patients With Advanced Fibrosis Due to Nonalcoholic Steatohepatitis

Zobair M. Younossi,1,2 Maria Stepanova,3 Mazen Noureddin,4 Kris V. Kowdley,5 Simone I. Strasser,6 Anita Kohli,7 Peter Ruane,8 Mitchell L. Shiffman,9 Aasim Sheikh,10 Nadege Gunn,11 Stephen H. Caldwell,12 Ryan S. Huss,13 Robert P. Myers,13 Vincent Wai-Sun Wong,14 Naim Alkhouri,15 Zachary Goodman,1,2 and Rohit Loomba4

Patient-reported outcomes (PROs) are important endpoints for clinical trials. The impact of investigational drugs on PROs of patients with advanced nonalcoholic steatohepatitis (NASH) was investigated. Patients with NASH with bridging fibrosis or compensated cirrhosis were enrolled in a phase 2, randomized, placebo-controlled study of selonsertib, firsocostat, or cilofexor, alone or in two-drug combinations (NCT03449446). PROs included Short Form 36 (SF-36), Chronic Liver Disease Questionnaire (CLDQ)-NASH, EuroQol Five Dimension (EQ-5D), Work Productivity and Impairment (WPAI), and 5-D Itch before and during treatment. A total of 392 patients with NASH (mean ± SD, 60 ± 9 years old; 35% men; 89% white; 72% diabetes; and 56% compensated cirrhosis) were included. Baseline Physical Functioning (PF) and Bodily Pain of SF-36 and Fatigue and Worry of CLDQ-NASH were significantly lower in patients with cirrhosis (total CLDQ-NASH score mean ± SD, 4.91 ± 1.06 with cirrhosis vs. 5.16 ± 1.14 without cirrhosis; P < 0.05). Lower baseline PRO scores were independently associated with age, female sex, greater body mass index, diabetes, clinically overt fatigue, and comorbidities (all P < 0.05). After 48 weeks of treatment, patients with ≥1-stage fibrosis improvement without worsening of NASH experienced improvement in EQ-5D and five out of six CLDQ-NASH domains (P < 0.05). Patients with ≥2-point decrease in their nonalcoholic fatty liver disease activity score (NAS) also had improvements in PF and Role Physical scores and all domains of CLDQ-NASH (P < 0.05). Progression to cirrhosis was associated with a decrease in PF scores of SF-36 (P ≤ 0.05). Fibrosis regression was independently associated with greater improvements in PF and EQ-5D scores, while NAS improvement was associated with improvement in fatigue and pruritus (all P < 0.05). Conclusion: Patients with advanced NASH experienced improvement in their PROs after fibrosis regression or improvement in disease activity. (Hepatology Communications 2021;5:1201-1211).

The global prevalence of nonalcoholic fatty liver disease (NAFLD) among adults is 25% but ranges from 13% in some African countries to almost 40% in South America.1 Nonalcoholic steatohepatitis (NASH) is the progressive subtype of NAFLD that can lead to progressive fibrosis and...
cirrhosis as well as impairment of health-related quality of life (HRQoL). Within the United States, the prevalence of NASH in the general population is estimated to range from 1.5% to 6.5%. These rates are higher among at-risk populations, including people with type 2 diabetes mellitus (T2DM) or obesity. Also, Hispanic Americans have higher rates of NASH and fibrosis than Caucasian and African American individuals. Although not uniformly progressive, some patients with NASH, particularly those with histologic fibrosis, can progress to end-stage liver disease, hepatocellular carcinoma (HCC), and ultimately, liver-related death. Largely due to the growing epidemics of obesity and T2DM,
NASH is becoming one of the most common causes of liver-related death, liver transplantation, and HCC in the United States and globally.\(^9,10\)

This report focuses on the potential new treatments for NASH as they relate to patient-reported outcomes (PROs). Recent reports have found that NASH can lead to significant morbidity and impairment of HRQL and other PROs.\(^11-13\) Physical health-related scores appear to be the most negatively affected, especially in those with NASH-related cirrhosis.\(^11\) Indeed, studies have shown that factors independently associated with lower PRO scores in patients with NASH include presence of cirrhosis, female sex, higher body mass index (BMI), smoking, T2DM, as well as psychiatric and other comorbidities.\(^14\) While some evidence suggests that fibrosis improvement may be followed by improvement in PROs,\(^15\) some therapies may negatively impact PROs due to their adverse-effect profile. Therefore, alongside efforts to reduce the clinical burden of NASH using preventive efforts and new treatment regimens, it is important to understand the impact of new therapies on PROs in this disease. Given this, the aim of this study was to assess PRO scores among patients with advanced fibrosis due to NASH before and after treatment with various investigational antifibrotic drugs in the context of a randomized controlled trial.

**Patients and Methods**

**PATIENTS AND STUDY DESIGN**

This analysis used data collected in the Study to Evaluate the Safety and Efficacy of Selonsertib, Firsocostat, Cilofexor, and Combinations in Participants With Bridging Fibrosis or Compensated Cirrhosis Due to Nonalcoholic Steatohepatitis (NASH) (ATLAS) study, a phase 2, randomized, double-blind, placebo-controlled study that evaluated the apoptosis signal-regulating kinase 1 inhibitor selonsertib, the acetyl-coenzyme A carboxylase inhibitor firsocostat, and the farnesoid X receptor (FXR) agonist cilofexor, alone or in two-drug combinations, in patients with advanced fibrosis due to NASH (NCT03449446). The methods and primary results of this study are reported elsewhere.\(^16\) Briefly, the ATLAS study enrolled and treated 392 patients with biopsy-confirmed NASH (defined as the presence of at least grade 1 steatosis, hepatocellular ballooning, and lobular inflammation according to the NAFLD activity score [NAS]) with either bridging fibrosis (F3) or compensated cirrhosis (F4) based on the NASH Clinical Research Network classification.\(^17\) Subjects with grade 0 steatosis and compensated cirrhosis (F4) were also enrolled if they had at least one risk factor for NASH (diabetes, insulin resistance, overweight, obesity, dyslipidemia, hypertension). In lieu of a biopsy, approximately 20% of the cohort was enrolled based on noninvasive markers (liver stiffness by vibration-controlled transient elastography [VCTE; FibroScan; Echosens, Paris, France] and enhanced liver fibrosis test [ELF; Siemens, Tarrytown, NY]) consistent with advanced fibrosis.

Exclusion criteria included a history of decompensated liver disease, Child-Pugh score >6, Model of End-Stage Liver Disease score >12, other causes of liver disease (e.g., alcoholic liver disease, hepatitis B, hepatitis C), liver transplantation, HCC, human immunodeficiency virus infection, recent excessive alcohol or illicit drug use, and any major or unstable comorbidities other than NASH and metabolic syndrome. Comorbidities were recorded using the Medical Dictionary for Regulatory Activities system and summarized by System Organ Class.\(^18\)

The study was conducted at 105 sites in the United States, Canada, Hong Kong, Australia, and New Zealand. Eligible patients were randomized to one of seven treatment groups: placebo; monotherapy with selonsertib (18 mg), cilofexor (30 mg), or firsocostat (20 mg); or combination therapy with cilofexor/selonsertib, firsocostat/selonsertib, or cilofexor/firsocostat. Study drugs were administered orally once daily for 48 weeks. The selonsertib monotherapy group was discontinued following reporting of negative results of the Statin Therapies for Elevated Lipid Levels Compared Across Doses to Rosuvastatin (STELLAR) studies; their baseline data were included in this study.\(^19,20\) The study was approved by the institutional review boards at participating sites, and all participants provided informed consent.

**OUTCOME MEASURES**

**Histology and Noninvasive Tests**

Liver biopsies obtained at screening and week 48 were evaluated in a blinded manner by a single central
The primary efficacy endpoint of the trial was the proportion of patients with one or more stage improvement in fibrosis without worsening of NASH (defined as a ≥1-point increase in ballooning or lobular inflammation) at week 48. Secondary endpoints included fibrosis improvement (without regard to NASH worsening), NASH resolution (defined as a reduction of lobular inflammation to 0 or 1 and hepatocellular ballooning to 0) without worsening of fibrosis, histologic progression to cirrhosis (in subjects without cirrhosis at baseline), and improvement in NASH activity defined as a ≥2-point reduction in NAS.

Due to the limited sensitivity of conventional histologic staging for detecting fibrosis regression, we also evaluated changes in two noninvasive markers of fibrosis, ELF and liver stiffness by VCTE. Specifically, an ELF response was defined as a ≥0.5-unit reduction and a liver stiffness by VCTE response was defined as a ≥25% relative reduction, both from baseline to week 48. Changes of this magnitude have been associated with reduced disease progression in this patient population.

PROs

PROs were collected on the first day of treatment and every 12 weeks thereafter. Patients self-administered PRO instruments before initiation of any treatment–related activities at each visit. This included any discussion about liver biopsy findings. PRO instruments included Short Form 36 (SF-36), the EuroQol Five-Dimension (EQ-5D), the Chronic Liver Disease Questionnaire–NASH (CLDQ–NASH), and the Work Productivity and Activity Impairment: Specific Health Problem (WPAI:SHP). In addition, pruritus was evaluated using the 5-D Itch and a visual analog scale for pruritus (pruritus VAS). These instruments have been validated in various clinical populations and used in trials of patients with chronic liver and other disorders.

The SF-36 is a generic HRQL instrument used to assess HRQL in eight domains: Physical Functioning (PF), Role Physical, Bodily Pain, General Health, Vitality, Social Functioning, Role Emotional, and Mental Health (all range from 0 to 100). It also includes two summary scores, physical component summary and mental component summary scores, which are linear combinations of the domain scores. Additionally, SF-6D health utility scores (range, 0 to 1) were calculated from the SF-36 responses using a described nonparametric Bayesian algorithm.

The EQ-5D is a generic instrument widely used for the calculation of health utility scores. In this instrument, health status is measured in terms of five dimensions: mobility, self-care, usual activities, pain/discomfort, and anxiety/depression; the resulting 5-digit number is then converted into a single weighted index score (range, 0 to 1) using a described crosswalk algorithm.

The CLDQ–NASH is a disease-specific PRO instrument that assesses HRQL of patients with NASH by specifically addressing its most frequent manifestations. The instrument includes 36 items grouped into six domains (Abdominal, Activity/Energy, Emotional, Fatigue, Worry, and Systemic; all range from 1 to 7) and a total score that is an average of the domain scores. The CLDQ–NASH has been validated in patients with biopsy-proven NASH.

The WPAI:SHP instrument evaluates impairment in daily activities and work productivity associated with a specific health problem. It includes a Work Productivity Impairment score, which is a sum of scores for Absenteeism (loss of work productivity owing to missing work hours) and Presenteeism (loss of work productivity owing to decreased productivity while working), and an activity impairment domain (all range from 0 to 1). Unlike the other instruments used in this study, WPAI:SHP returns greater scores for more impairment.

The 5-D Itch instrument assesses the severity of pruritus using five dimensions (degree, duration, direction, disability, and distribution; each on a scale from 1 to 5). The total score is the sum of the dimension scores (range from 5 to 25), with higher scores indicative of more pruritus. In addition, the pruritus VAS was used to assess the severity of itching. With this instrument, patients are asked to describe the severity of their itching by placing a mark on a 10-cm-long scale; the resulting score is the position of the mark in millimeters from the beginning of the scale (0, labeled as “no itching”).
scales to a universal scale ranging from 0 to 100, for presentation purposes.

**Patient and Public Involvement Statement**

Due to the clinical trial nature of this study, patients were not involved in the development of the study but were free to participate as they deemed reasonable for themselves.

**STATISTICAL ANALYSES**

All clinical and demographic parameters and PRO scores were summarized as n (%) or mean ± SD. These parameters were compared between patient subgroups using the chi-square test and Mann-Whitney test, as appropriate; *P* ≤ 0.05 was considered statistically significant. Subgroups for comparison were defined by baseline cirrhosis status, treatment group, and whether or not the specific efficacy endpoint was met. Only observed PRO data were used; no imputation was performed. In addition, we calculated changes from baseline in PRO scores with reference to patients’ own baseline levels. Treatment effects on the changes in PRO scores were evaluated by the treatment group for each study visit after the baseline. For this purpose, least-square mean estimates were returned by mixed regression models that included treatment, visit, treatment-visit interaction, baseline PRO value, and stratification factors involved in randomization (diabetes and cirrhosis) used as fixed effects and subjects used as a random effect. Changes in PRO scores were also summarized as arithmetic mean ± SEM when grouping other than by treatment group was used and were compared to zero by the matched pairs signed-rank test and between the groups of interest by the Kruskal-Wallis rank-sum test.

Independent predictors of the baseline PRO scores were assessed in a series of generalized linear regression models with stepwise bidirectional selection from the list of baseline parameters, including demographics (age, sex, race, location), clinical parameters (smoking status, BMI, the presence of type 2 diabetes, and other comorbidities), presence of cirrhosis, and other histologic parameters (steatosis, hepatocellular ballooning, lobular inflammation). All analyses were conducted using SAS 9.4 (SAS Institute, Cary, NC).

**Results**

In total, 392 patients with NASH were included, of whom the majority (56%) had cirrhosis. The mean age ± SD and BMI were 60 ± 9 years and 34.5 ± 6.9 kg/m², respectively; 35% were men, 89% white, 6% Asian, 86% enrolled in the United States, 49% employed, and 72% had diabetes. Compared with patients without cirrhosis, those with cirrhosis were on average, 2 years older, had a higher prevalence of diabetes and gastrointestinal and vascular comorbidities, less steatosis, and substantially higher liver stiffness by VCTE and serum markers of fibrosis (all *P* < 0.05) (Table 1).

**PROs AT BASELINE**

Baseline scores indicated better PROs in patients without versus with cirrhosis; this was primarily in the physical health-related domains (Supporting Table S1), including PF (mean ± SD, 71.1 ± 27.4 vs. 66.4 ± 26.5, respectively; *P* = 0.0335), Bodily Pain (67.4 ± 26.6 vs. 62.1 ± 25.5; *P* = 0.0480), and physical component summary (46.2 ± 10.1 vs. 44.4 ± 9.7; *P* = 0.0410) of the SF-36; and Fatigue (4.72 ± 1.50 vs. 4.39 ± 1.32; *P* = 0.0105), Worry (5.16 ± 1.51 vs. 4.76 ± 1.61; *P* = 0.0113), and total score (5.16 ± 1.14 vs. 4.91 ± 1.06; *P* = 0.0204) of the CLDQ-NASH. The mean pruritus VAS was lower in patients without versus with cirrhosis, indicative of less itch (12.3 ± 20.9 vs. 15.2 ± 20.9, respectively; *P* = 0.0136). In multivariate analysis, lower baseline PRO scores at baseline were independently associated with older age (physical health-related scores), younger age (mental health-related scores), female sex, higher BMI, and the presence of diabetes, clinically overt fatigue, psychiatric, musculoskeletal, nervous system, gastrointestinal, and cardiac comorbidities (all *P* < 0.05) (Supporting Table S2).

**CHANGES IN PROs DURING TREATMENT**

During treatment, cilofexor-containing regimens were associated with some PRO improvement, especially in scores measured by the disease-specific CLDQ-NASH (Fig. 1A), although the same trends in health utility scores (Fig. 1B) and physical health-related scores measured by generic instruments (Fig. 1C) were less pronounced. On the other hand, selonsertib-containing regimens were associated with...
### TABLE 1. CLINICODEMOGRAPHIC CHARACTERISTICS OF PATIENTS WITH NASH INCLUDED IN THIS STUDY

| Demographics | Noncirrhotic (<F4) (n = 171) | Compensated Cirrhosis (F4) (n = 221) | P Value | Total (N = 392) |
|--------------|------------------------------|--------------------------------------|---------|-----------------|
| **Age, years** | 58.1 ± 9.4                  | 60.8 ± 8.5                           | 0.0036  | 59.6 ± 9.0      |
| **Male sex**  | 65 (38.0%)                  | 74 (33.5%)                           | 0.35    | 139 (35.5%)     |
| **White race**| 150 (87.7%)                 | 199 (90.0%)                          | 0.46    | 349 (89.0%)     |
| **Black race**| 3 (1.8%)                    | 4 (1.8%)                             | 0.97    | 7 (1.8%)        |
| **Asian race**| 14 (8.2%)                   | 11 (5.0%)                            | 0.20    | 25 (6.4%)       |
| **Enrolled in the United States** | 145 (84.8%) | 191 (86.4%)                         | 0.65    | 336 (85.7%)     |
| **Current smoker** | 9 (5.3%)       | 19 (8.6%)                            | 0.20    | 28 (7.1%)       |
| **Employed**  | 81 (48.5%)                  | 107 (49.5%)                          | 0.84    | 188 (49.1%)     |
| **BMI, kg/m²**| 34.4 ± 7.0                  | 34.5 ± 6.8                           | 0.95    | 34.5 ± 6.9      |

| **Comorbidities** | | | | |
|-------------------| | | | |
| **Diabetes mellitus** | 114 (66.7%) | 168 (76.0%) | 0.0410 | 282 (71.9%) |
| **Blood and lymphatic system disorders** | 21 (12.3%) | 42 (19.0%) | 0.07 | 63 (16.1%) |
| **Cardiac disorders** | 25 (14.6%) | 47 (21.3%) | 0.09 | 72 (18.4%) |
| **Ear and labyrinth disorders** | 23 (13.5%) | 16 (7.2%) | 0.0416 | 39 (9.9%) |
| **Endocrine disorders** | 39 (22.8%) | 54 (24.4%) | 0.71 | 93 (23.7%) |
| **Asthemic conditions or fatigue** | 27 (15.8%) | 28 (12.7%) | 0.38 | 55 (14.0%) |
| **Gastrointestinal disorders** | 124 (72.5%) | 184 (83.3%) | 0.0101 | 308 (78.6%) |
| **Immune systemic disorders** | 83 (48.5%) | 103 (46.6%) | 0.70 | 186 (47.4%) |
| **Infections and infestations** | 63 (36.8%) | 95 (43.0%) | 0.22 | 158 (40.3%) |
| **Musculoskeletal, connective tissue disorders** | 101 (59.1%) | 150 (67.9%) | 0.07 | 251 (64.0%) |
| **Neoplasms, benign or malignant** | 39 (22.8%) | 71 (32.1%) | 0.0417 | 110 (28.1%) |
| **Nervous system disorders** | 73 (42.7%) | 99 (44.8%) | 0.68 | 172 (43.9%) |
| **Psychiatric disorders** | 78 (45.6%) | 113 (51.1%) | 0.28 | 191 (48.7%) |
| **Renal and urinary disorders** | 38 (22.2%) | 57 (25.8%) | 0.41 | 95 (24.2%) |
| **Respiratory disorders** | 73 (42.7%) | 102 (46.2%) | 0.49 | 175 (44.6%) |
| **Skin and subcutaneous tissue disorders** | 43 (25.1%) | 72 (32.6%) | 0.11 | 115 (29.3%) |
| **Vascular disorders** | 123 (71.9%) | 180 (81.4%) | 0.0257 | 303 (77.3%) |
| **Vision disorders** | 57 (33.3%) | 71 (32.1%) | 0.80 | 128 (32.7%) |

| **Liver histology** | | | | |
|-------------------| | | | |
| **Steatosis grade 0** | 2 (1.2%) | 35 (15.8%) | <0.0001 | 37 (9.4%) |
| **Steatosis grade 1** | 155 (90.6%) | 181 (81.9%) | 0.0142 | 336 (85.7%) |
| **Steatosis grade 2** | 14 (8.2%) | 5 (2.3%) | 0.0068 | 19 (4.8%) |
| **Lobular inflammation grade 1** | 17 (9.9%) | 16 (7.2%) | 0.34 | 33 (8.4%) |
| **Lobular inflammation grade 2** | 50 (29.2%) | 66 (29.9%) | 0.89 | 116 (29.6%) |
| **Lobular inflammation grade 3** | 104 (60.8%) | 139 (62.9%) | 0.67 | 243 (62.0%) |
| **Hepatocyte ballooning grade 0** | 2 (1.2%) | 4 (1.8%) | 0.61 | 6 (1.5%) |
| **Hepatocyte ballooning grade 1** | 24 (14.0%) | 26 (11.8%) | 0.50 | 50 (12.8%) |
| **Hepatocyte ballooning grade 2** | 145 (84.8%) | 191 (86.4%) | 0.65 | 336 (85.7%) |
| **NAS** | 5.42 ± 1.03 | 5.27 ± 1.05 | 0.13 | 5.33 ± 1.04 |
| **CPA, %** | 4.15 ± 2.39 | 10.4 ± 5.8 | <0.0001 | 7.63 ± 5.58 |

| **NITs** | | | | |
|-------------------| | | | |
| **Liver stiffness by VCTE, kPa** | 15.7 ± 9.9 | 22.2 ± 13.5 | <0.0001 | 19.4 ± 12.5 |
| **ELF score** | 9.72 ± 0.90 | 10.4 ± 1.1 | <0.0001 | 10.1 ± 1.1 |
| **NFS** | −0.209 ± 1.297 | 0.471 ± 1.255 | <0.0001 | 0.175 ± 1.316 |
some PRO decrease in mental health scores (Fig. 1D). No other PRO decrements were observed, while the Worry score of CLDQ-NASH improved in all treatment groups (all \( P < 0.05 \)). Finally, there was no association of any treatment regimen with changes in the two studied pruritus assessments (all \( P > 0.05 \)).

**ASSOCIATIONS BETWEEN CHANGES IN PROs AND HISTOLOGIC RESPONSES**

After 48 weeks of treatment, patients who achieved the primary endpoint of the study, a ≥1-stage improvement in fibrosis without worsening of NASH (observed in 16% of cases with paired histology overall), experienced improvement in their EQ-5D utility score and five out of six CLDQ-NASH domain scores (all \( P < 0.05 \)) (Fig. 2A). Similar changes in PROs were observed in patients who experienced improvement in fibrosis without regard to NASH activity (19% of observed cases) (Fig. 2B). Patients who had a ≥2-point decrease in their NAS (19% of cases) had improvements in PF and Role Physical scores and all domains of CLDQ-NASH (all \( P < 0.05 \)) (Fig. 2C). As NASH resolution without worsening of fibrosis was rarely

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**TABLE 1. Continued**

|                         | Noncirrhotic (<F4) (n = 171) | Compensated Cirrhosis (F4) (n = 221) | \( P \) Value | Total (N = 392) |
|-------------------------|-----------------------------|------------------------------------|--------------|---------------|
| APRI                    | 0.718 ± 0.507               | 0.792 ± 0.537                      | 0.0404       | 0.759 ± 0.525 |
| Fibrosis-4 score        | 1.90 ± 0.97                 | 2.45 ± 1.17                        | <0.0001      | 2.21 ± 1.12  |
| FibroTest score         | 0.416 ± 0.229               | 0.540 ± 0.210                      | <0.0001      | 0.486 ± 0.227 |

All clinical and demographic parameters and PRO scores were summarized as n (%) or mean ± SD. Abbreviations: APRI, aspartate aminotransferase to platelet ratio index; CPA, collagen proportionate area; NFS, NAFLD fibrosis score.
observed in this cohort of patients with advanced fibrosis (n = 6), analyses of PROs were not conducted according to this endpoint. Patients without cirrhosis at baseline who progressed to cirrhosis at week 48 (19% of cases) experienced a decrease in their PF scores (mean ± SD, −6.0 ± 3.0; \( P = 0.05 \)).

In multivariate analyses with adjustment for clinicalodemographic parameters (location, age, sex, race, smoking status, baseline BMI, type 2 diabetes, cirrhosis, NAS), treatment regimen, and a change in BMI and hemoglobin A1c (HbA1c) from baseline (in order to account for potential improvement in diabetes control while in the trial), improvement of fibrosis at week 48 of treatment was independently associated with a greater improvement in PF (mean ± SD, beta = +5.37 ± 2.47; \( P = 0.0306 \)), physical component summary (+2.24 ± 0.96; \( P = 0.0199 \)), and EQ-5D (+0.036 ± 0.017; \( P = 0.0308 \)) scores (Supporting Table S3). In addition, a ≥2-point improvement in NAS was associated with improvement in Fatigue (mean ± SD, +0.32 ± 0.16; \( P = 0.0435 \)), Systemic Symptoms (+0.25 ± 0.12; \( P = 0.0423 \)), and total (+0.21 ± 0.10; \( P = 0.0405 \)) scores by CLDQ-NASH (Supporting Table S3). The improvements in CLDQ-NASH scores seem to be driven primarily by improvement in hepatocyte ballooning (by ≥1 point mean ± SD, beta = +0.41 ± 0.14 for Activity, +0.27 ± 0.12 for Systemic Symptoms). At the same time, there was no significant association of PRO changes with improvement in lobular inflammation or steatosis (all \( P > 0.05 \)). Interestingly, we also found that the only trends in PRO scores associated with changes in HbA1c were both pruritus metrics, namely, 5D-Itch (mean ± SD, beta = +0.67 ± 0.24) and pruritus VAS (beta = +5.21 ± 1.54; both \( P < 0.01 \)) (Supporting Table S3). Notably, there was no association of
histologic improvement with improvement in pruritus after adjustment for improvement in HbA1c (all $P > 0.05$).

**ASSOCIATIONS BETWEEN CHANGES IN PROs AND NONINVASIVE TESTS OF FIBROSIS RESPONSES**

Patients who had their ELF score decreased by at least 0.5 points (28%) had improvements in four out of six domains of CLDQ-NASH (up to +8.4% of a range size, $P \leq 0.05$) (Fig. 2D). However, the association with ELF improvement was found to be strongest at the cutoff of 0.3 as statistically significant improvements were detected in all domains of CLDQ-NASH (up to +10.6% of a range size, $P < 0.05$) (Supporting Fig. S1A). A reduction in liver stiffness of at least 25% (37% of observed cases) was similarly associated with significant improvements in five out of six CLDQ-NASH domains and in EQ-5D utility scores (up to +9.6% of a range size, $P < 0.05$) (Supporting Fig. S1B).

**Discussion**

In this study, we analyzed PRO scores from patients with advanced NASH within the phase 2 ATLAS trial. The approach was to evaluate therapies targeting different pathogenic mechanisms of NASH. Serial measurement of PROs and liver histology before and during treatment with these active therapies have enabled an assessment of the impact of NASH on PROs as well as the impact of these treatment regimens on PROs and the histologic changes that may be associated with PRO improvement in NASH.

At baseline, we confirmed that patients with compensated NASH cirrhosis had lower PRO scores than patients without cirrhosis before the initiation of treatment, especially in the domains related to physical health. In addition, one of the pruritus scores (the pruritus VAS) was worse in patients with cirrhosis. Although this finding requires validation, it may be clinically relevant because some classes of medication, notably FXR agonists, may cause pruritus as a side effect. Given this observation, assessment of pruritus at baseline may be needed to better control the side-effect burden of these regimens. Finally, patients with cirrhosis had worse scores related to Worry on the CLDQ-NASH, likely due, at least in part, to uncertainty regarding their prognosis.

Consistent with reports in patients with advanced fibrosis due to NASH, predictors of lower PRO scores at baseline included older age, higher BMI, the presence of type 2 diabetes, as well as musculoskeletal, cardiovascular, and psychiatric comorbidities. Indeed, after adjustment for these factors, the presence of cirrhosis was not independently associated with PRO scores compared with bridging fibrosis (except for the Worry score of CLDQ-NASH), likely due to the high prevalence of these comorbidities in advanced NASH. The impact of treatment of these comorbidities on PROs in patients with NASH requires further study.

In this regard, we evaluated changes in PRO scores in the context of specific therapies for NASH and did not observe statistically significant differences between treatment regimens. However, promising trends in CLDQ-NASH scores were noted with cilofexor-containing regimens, including the combination with firsocostat. Importantly, adverse changes in pruritus-related PROs were not observed in patients treated with the FXR agonist cilofexor, supporting the tolerability of this therapy from an itch perspective. Indeed, only 1 of 195 patients treated with cilofexor (<1%) in the trial discontinued therapy due to pruritus. Interestingly, the Worry scores of the CLDQ-NASH improved in all treatment groups, potentially attesting to the benefits of patient participation in clinical trials (e.g., due to close follow-up and monitoring).

Of importance was the observation of improvements in PROs in patients with evidence of histologic improvement. Specifically, PRO gains were noted in patients with improvements of both NASH activity, as measured by the NAS and in particular its hepatocyte ballooning component, and fibrosis, as evaluated histologically by transient elastography or with the noninvasive serum marker ELF. In this context, the most significant gains were observed as expected in the PRO domains assessed by the disease-specific CLDQ-NASH, which included improvements in abdominal symptoms, physical activity, as well as NASH-related emotional health and fatigue. Interestingly, in multivariate analysis, fibrosis regression was found to be independently associated with
greater improvements in generic physical health-related scores while a decrease of the NAS score was associated with improvement in some disease-specific PRO scores. Conversely, patients without cirrhosis at baseline who experienced histologic progression to cirrhosis experienced a decrease in their PF scores on the SF-36, supporting the impact of cirrhosis on PROs. All these findings are consistence with reports from clinical trials on the association of PRO improvement with achieving histologic and other treatment endpoints, such as improvement of NAS score, fibrosis stage, and noninvasive test (NITs) in patients with NASH and different stages of baseline fibrosis.\(^{(20,31,33)}\) In fact, a similar association of PRO improvement with improvement in NITs was recently shown using a large cohort of patients with advanced NASH.\(^{(33)}\) The exact mechanism(s) behind all these observations are unclear as we do not have in-depth cytokine measurements in this study,\(^{(34)}\) but taken together they underscore the importance of reversing fibrosis and reducing the inflammatory milieu of NASH.

Assessment of PROs in the clinical trials of NASH can provide a number of valuable pieces of information. First, the assessment could shed light whether histologic or NIT improvement can lead to PRO improvement. In this context, the main findings of our study are in line with published literature.\(^{(33)}\) Second, PRO assessment could provide patient-centric information whether the side-effect profile of the regimen has a negative impact on patients’ experience.\(^{(35)}\) This is especially important when combination regimens, such as those combining cilofexor and firsocostat, are used. In this context, our data provide new information that the combination regimens used in this clinical trial may not have a significant negative impact on PROs.

This study has several limitations that warrant discussion. First, the patients enrolled in this clinical trial had advanced fibrosis due to NASH. The generalizability of these findings to patients with milder disease and those seen in real-world settings requires confirmation. In addition, the PROs evaluated in this study were self-reported and thus subject to recall or perception biases. However, the latter bias should be mitigated by the double-blind nature of this study and the fact that patients completed the PRO instruments while unaware of their most recent clinical results. Moreover, consistency of the findings across multiple PRO domains describing similar aspects of patients’ well-being could indirectly support their validity. Finally, the study was not powered to detect differences in PRO scores between treatment regimens. Hence, any inferences regarding differential effects of these therapies on PROs in patients with NASH should be made cautiously.

As research progresses on treatments for patients with NASH-related fibrosis, understanding the impact of NASH therapies toward HRQL and other PROs is important. In this study of several investigational regimens, we found that patients with NASH-related cirrhosis at baseline reported significantly lower PRO scores, especially in physical health-related domains, compared with patients without cirrhosis. At the same time, we found that achieving clinically relevant endpoints, including fibrosis regression and improvement in NASH disease activity, was associated with improvement in some PROs captured by the disease-specific CLDQ-NASH. Finally, while statistically significant differences between the effects of treatment regimens on PROs were not observed, cilofexor-containing regimens, including the combination with firsocostat, appeared to have the most positive effects on PROs. Longer term research is needed to confirm the sustainability of the reported PRO gains in patients with NASH with histologic improvement during treatment.

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