Ezetimibe combination therapy with statin for non-alcoholic fatty liver disease: an open-label randomized controlled trial (ESSENTIAL study)

Yongin Cho1,2†, Hyungjin Rhee3†, Young-eun Kim4, Minyoung Lee4,5, Byung-Wan Lee4,5, Eun Seok Kang4,5, Bong-Soo Cha4,5, Jin-Young Choi3 and Yong-ho Lee4,5,6*

Abstract

Background: The effect of ezetimibe, Niemann-Pick C1-like 1 inhibitor, on liver fat is not clearly elucidated. Our primary objective was to evaluate the efficacy of ezetimibe plus rosuvastatin versus rosuvastatin monotherapy to reduce liver fat using magnetic resonance imaging-derived proton density fat fraction (MRI-PDFF) in patients with non-alcoholic fatty liver disease (NAFLD).

Methods: A randomized controlled, open-label trial of 70 participants with NAFLD confirmed by ultrasound who were assigned to receive either ezetimibe 10 mg plus rosuvastatin 5 mg daily or rosuvastatin 5 mg for up to 24 weeks. The liver fat change was measured as average values in each of nine liver segments by MRI-PDFF. Magnetic resonance elastography (MRE) was used to measure liver fibrosis change.

Results: Combination therapy significantly reduced liver fat compared with monotherapy by MRI-PDFF (mean difference: 3.2%; p = 0.020). There were significant reductions from baseline to study completion by MRI-PDFF for both the combination and monotherapy groups, respectively (18.1 to 12.3%; p < 0.001 and 15.0 to 12.4%; p = 0.003). Individuals with higher body mass index, type 2 diabetes, insulin resistance, and severe liver fibrosis were likely to be good responders to treatment with ezetimibe. MRE-derived change in liver fibrosis was not significantly different (both groups, p > 0.05). Controlled attenuation parameter (CAP) by transient elastography was significantly reduced in the combination group (321 to 287 dB/m; p = 0.018), but not in the monotherapy group (323 to 311 dB/m; p = 0.104).

Conclusions: Ezetimibe and rosuvastatin were found to be safe to treat patients with NAFLD. Furthermore, ezetimibe combined with rosuvastatin significantly reduced liver fat in this population.

Trial registration: The trial was registered at ClinicalTrials.gov (registration number: NCT03434613).

Keywords: Niemann-Pick C1-like 1 inhibitor, Statins, Hepatic steatosis, Hepatic fibrosis

Background

The incidence of non-alcoholic fatty liver disease (NAFLD) is increasing globally and is closely related to metabolic syndrome features including obesity, insulin resistance, dyslipidemia, and diabetes mellitus (DM). Cardiovascular disease, which is closely related to dyslipidemia, is a major cause of mortality in patients...
with NAFLD [1]. Furthermore, NAFLD independently increases the risk of cardiovascular complications [2]. Therefore, management of dyslipidemia in patients with NAFLD is an important concern. In recent treatment guidelines for dyslipidemia, more aggressive treatment to target low-density lipoprotein cholesterol (LDL-C) is recommended [3]. Statins, the most potent drugs that lower LDL-C, are known to have several side effects whose risks appear to increase with higher doses [4]. For this reason, concomitant use of second-line drugs such as ezetimibe with a lower statin dose may be beneficial.

Cholesterol alone can contribute to the progression of non-alcoholic steatohepatitis (NASH) [5], and the effects of anti-dyslipidemia therapies on hepatic steatosis and/or fibrosis have been investigated. Evidence of benefit by using a statin in patients with NAFLD is limited [6], and statin was not beneficial in improving NASH in a previous randomized controlled trial [7]. Also, the effect of ezetimibe on hepatic steatosis is still controversial. Ezetimibe exhibits lipid-lowering effects through inhibition of Niemann-Pick C1-like 1 (NPC1-L1) in the intestine and has been reported to reduce visceral fat and improve insulin resistance in several studies [8, 9]. Improvement of hyperinsulinemia may lead to the inhibition of sterol regulatory element-binding protein-1c (SREBP-1c) and blockade of fatty acid synthase [10]. And this mechanism can contribute to improvement in hepatic steatosis. Recently, our group reported that the use of ezetimibe affects autophagy of hepatocytes and improves hepatic steatosis in an animal model [11].

However, the effects of ezetimibe have not been clearly elucidated in human studies. Ezetimibe did not significantly reduce liver fat in a previous randomized controlled trial by Loomba et al. [12]. This study was designed using ezetimibe alone, with the use of concomitant statin drugs not clearly controlled. Considering that the improvement of cardiovascular disease risk through ezetimibe has been identified in combination with statins [13], increased understanding of ezetimibe effect on NAFLD in combination with a statin will be of great clinical significance. Therefore, in this study, we investigated whether ezetimibe contributes to hepatic steatosis improvement in the setting of controlled statin treatment.

Methods
Study design and patient population
The ESSENTIAL (Effects of Statin Monotherapy and Statin/Ezetimibe Combination Therapy on Non-alcoholic Steatohepatitis in patients with Hyperlipidemia and Fatty Liver) study was an investigator-initiated, randomized, open-label, prospective, active-controlled clinical trial to examine the efficacy of ezetimibe 10 mg/day orally combined with rosuvastatin 5 mg/day versus rosuvastatin 5 mg/day orally for up to 24 weeks to improve hepatic steatosis as measured by magnetic resonance imaging-derived proton density fat fraction (MRI-PDFF). Participants of the study were recruited from Severance Hospital from May 2018 to June 2019 in Seoul, Korea.

The trial was registered at ClinicalTrials.gov (registration number: NCT03434613). All participants provided written informed consent and the Ethics Committee of the Yonsei University College of Medicine approved this study (4-2017-1168), which conforms to the ethical principles of the 1975 Declaration of Helsinki.

Inclusion criteria
Patients were required to meet all the following criteria for study inclusion: 19–80 years of age, hepatic steatosis documented by abdominal ultrasound, hyperlipidemia indicated to treat by the domestic dyslipidemia treatment guideline [14], diagnosed with type 2 diabetes (T2DM, HbA1c ≤9.0% with no change in type of oral or injectable hypoglycemic agent/s within 12 weeks of enrollment) or without diabetes, and written informed consent. Exclusion criteria are provided in the Supplementary Appendix (Additional file 1).

Baseline assessment at screening
All patients underwent a baseline assessment before undergoing randomization, including medical history and physical exam. For those who had undergone abdominal ultrasonography within the past 1 year, test results were reviewed for the presence of hepatic steatosis. Otherwise, abdominal ultrasonography was performed for screening purposes.

Randomization
A random allocation sequence was computer-generated elsewhere, and participants were assigned in a 1:1 ratio to receive treatment with ezetimibe plus rosuvastatin or rosuvastatin alone. To equalize the proportion of type 2 diabetes in both groups, patients were stratified according to the presence of diabetes before randomization.

Primary and secondary outcomes
The primary outcome was change in liver fat by MRI-PDFF quantified as the average in colocalized regions of interest (ROI) within each of nine liver segments [15]. Secondary outcomes were change in liver fibrosis by magnetic resonance elastography (MRE), change in insulin sensitivity determined by homeostatic model assessment insulin resistance (HOMA-IR), and changes in parameters including body weight, body mass index (BMI), waist circumference (WC), systolic and diastolic blood pressure, fasting glucose, HbA1c, free fatty acids,
platelets, alkaline phosphatase, total bilirubin, aspartate aminotransferase (AST), alanine aminotransferase (ALT), total cholesterol, triglyceride, high-density lipoprotein cholesterol (HDL-C), LDL-C, high sensitivity C-reactive protein, HOMA-beta, and biomarkers including interleukin (IL)-1 beta, IL-8, and IL-18. Transient elastography (TE) (Fibroscan®; Echosens, Paris, France) was also performed before and after treatment, and changes in the hepatic steatosis index-controlled attenuation parameter (CAP) and hepatic fibrosis index-liver stiffness measurement (LSM) were also included as secondary outcome measures.

MRI-PDFF for fat quantification and MRE for liver fibrosis quantification
For hepatic fat quantification, MRI-PDFF sequence (mDIXON Quant) was used, and a fat fraction map was automatically generated by the manufacturer’s console. For liver fibrosis measurement, MRE was performed using a 2-dimensional gradient-echo sequence and a passive driver placed on the participant’s right upper abdomen (see Additional file 1: Supplementary Appendix for details and Table S1).

Statistical analysis
All categorical variables were expressed as number (proportion) and compared by $\chi^2$ analysis. The independent samples $t$-test or Wilcoxon-Mann-Whitney $U$ test was used for differences between continuous variables. For the primary objective, per-protocol analysis was performed. Wilcoxon signed-rank test or logistic regression was used for additional analysis, when indicated.

We expected the ezetimibe treatment group to have a liver fat reduction of >10% compared to baseline and the statin monotherapy group to have a <5% reduction compared to baseline. We also expected a dropout rate of 15–20%; thus, we planned to enroll 70 participants to achieve power of at least 80% with $\alpha$ and 0.05 with $\beta$ assuming study completion by ≥29 participants per arm.

Additional analyses for treatment response were conducted across multiple subgroups to examine whether the effect of ezetimibe differed on the basis of patient demographics or clinical characteristics. Characteristics for subgroup analysis included age (age 50 or more, under age 50), sex (men, women), body mass index (30 kg/m² or more, under 30 kg/m²), presence of type 2 diabetes, sarcopenia (defined as a skeletal muscle mass index (SMI, %) <2 standard deviations below the gender-specific mean for the Korean population: SMI (%) <29.0 in men and <22.9 in women) [16], HOMA-IR (over median value of the whole study population, under median value), baseline MRI-PDFF (over median value, under median value), and baseline MRE (over median value, under median value). Response was defined as ≥30% relative reduction or ≥5% absolute decline in MRI-PDFF from baseline to end of treatment [17].

All statistical analyses were performed using IBM SPSS Statistics version 26.0 (IBM Corp., Armonk, NY, USA).

Results
Baseline characteristics of study participants
A total of 70 participants with NAFLD were randomly assigned to receive either combination therapy with ezetimibe plus rosvastatin ($n = 34$) or rosvastatin monotherapy ($n = 36$). Three subjects in the combination group and 3 patients in the monotherapy group were excluded from the final analysis (Fig. 1). Baseline demographic and biochemical profiles are summarized in Table 1, showing that both groups had comparable characteristics at study entry. The study population included 42.9% women and 74.3% of participants had type 2 diabetes. All participants were Asian with an average BMI of 28.5±3.5 kg/m². The mean PDFF value before treatment was 16.5±7.8% and the mean CAP value was 313.7 dB/m±33.2 dB/m for the whole study population.

Effect of rosvastatin with/without ezetimibe on liver fat as assessed by MRI-PDFF
Compared to baseline, there were significant reductions in end-of-treatment MRI-PDFF for both the combination group (18.1 to 12.3%; $p < 0.001$) and the monotherapy group (15.0 to 12.4%; $p = 0.003$) (Table 2 and Additional file 1: Fig. S1A). In the per-protocol analysis, ezetimibe combined with rosvastatin was significantly better than rosvastatin monotherapy in reducing liver fat measured by MRI-PDFF (mean difference 3.2%; $p = 0.020$) (Fig. 2). The degree to which hepatic steatosis improved was similar among individual segments within the treatment group regardless of treatment assignment.

Effect of rosvastatin with/without ezetimibe on liver fibrosis as assessed by MRE
Compared to baseline, there were no significant differences at end of treatment by MRE in either the combination group or monotherapy group (2.0 to 2.1 kPa; $p = 0.507$ and 2.2 to 2.2 kPa; $p = 0.539$, respectively) (Table 2 and Additional file 1: Fig. S1B). MRE-derived liver fibrosis was not significantly different between groups ($p = 0.898$).

Effect of rosvastatin with/without ezetimibe on liver fat and fibrosis as assessed by transient elastography
We also evaluated changes in liver fat and fibrosis by TE. A significant decrease in CAP at end of treatment
was observed for ezetimibe plus rosuvastatin (321 to 287 dB/m; \( p=0.018 \)), but not for the rosuvastatin alone group (323 to 311 dB/m; \( p=0.104 \), Additional file 1: Table S2). However, the between-group difference was not statistically significant (\( p=0.253 \)). In terms of hepatic fibrosis, there was no significant difference in LSM from baseline to end of treatment in the combination group (6.0 to 6.1 kPa; \( p=0.339 \)) or the monotherapy group (6.7 to 6.0 kPa; \( p=0.881 \)).

**Effect of rosuvastatin with/without ezetimibe on biochemical indices**

Changes in anthropometric and biochemical variables are summarized in Table 3. There were significant decreases in BMI, waist circumference, triglyceride, LDL cholesterol, and C-reactive protein (CRP) in both groups (all \( p<0.05 \)). However, changes in all variables including BMI, alanine aminotransferase, aspartate aminotransferase, fasting glucose, HbA1c, and homeostatic model assessment insulin resistance were not significantly different between the two groups (all \( p > 0.05 \)).

To explore drug mechanisms, additional tests were performed on several values known to be inflammation markers of NASH progression. Significant improvement was reported in IL-18 in the combination group and in tPAI-1 and IL-18 in the monotherapy group, but there was no significant difference in treatment effect between groups (all \( p > 0.05 \)).

**Prediction of response by ezetimibe combination treatment**

Defining response as a \( \geq 30\% \) relative reduction or \( \geq 5\% \) absolute decline in MRI-PDFF from baseline to end of treatment, ezetimibe combination treatment more effectively improved hepatic steatosis in participants with higher BMI (\( \geq 30 \text{ kg/m}^2 \)), presence of T2DM, presence of sarcopenia, increased HOMA-IR (\( \geq 3.7 \), the median value of the study population), and increased baseline MRE (\( \geq 2.1 \), the median value in this study) (Fig. 3).

**Adverse events**

Three participants in each treatment arm dropped out of the study, but this was not associated with adverse events. One subject in each group had no documented hepatic steatosis (<5% on MRI-PDFF) at baseline. Two participants in each group were discontinued due to protocol violations. There was no significant adverse event reported in this study.

**Discussion**

This randomized controlled clinical trial showed ezetimibe with rosuvastatin was significantly better than rosuvastatin monotherapy for reducing liver fat as measured by MRI-PDFF. Consistent with this finding, a significant reduction of liver fat was also confirmed in end-of-treatment CAP measured by TE, but only for the combination group. Ezetimibe combination treatment
Table 1  Baseline demographic and biochemical characteristics

|                          | Ezetimibe + rosuvastatin (n = 34) | Rosuvastatin alone (n = 36) | p-value |
|--------------------------|----------------------------------|-----------------------------|---------|
| Demographics             |                                  |                             |         |
| Age, years               | 50.3 (12.9)                      | 52.5 (19.1)                 | 0.647   |
| Female; n (%)            | 12 (35.3)                        | 18 (50.0)                   | 0.214   |
| Weight, kg               | 78.2 (19.3)                      | 77.2 (15.5)                 | 0.958   |
| BMI, kg/m²               | 27.7 (6.6)                       | 28.6 (3.6)                  | 0.196   |
| Waist circumference, cm  | 95.0 (15.3)                      | 96.5 (11.1)                 | 0.685   |
| Presence of diabetes, n (%) | 26 (72.2)                     | 26 (76.5)                   | 0.684   |
| Seated SBP, mmHg         | 126.0 (11.0)                     | 123.0 (13.0)                | 0.676   |
| Seated DBP, mmHg         | 81.0 (11.0)                      | 80.5 (13.0)                 | 0.851   |
| Biochemical profile      |                                  |                             |         |
| ALT, IU/l                | 37.5 (30.0)                      | 30.0 (22.0)                 | 0.099   |
| AST, IU/l                | 24.5 (13.0)                      | 24.5 (13.0)                 | 0.706   |
| Platelets, x 10^9/μl     | 263.0 (89.0)                     | 258.5 (77.0)                | 0.307   |
| Alk Phos, U/l            | 64.0 (22.0)                      | 68.5 (36.0)                 | 0.121   |
| GGT, U/l                 | 42.5 (36.0)                      | 35.0 (33.0)                 | 0.347   |
| Total bilirubin, mg/dl   | 0.6 (0.3)                        | 0.7 (0.4)                   | 0.557   |
| Glucose, mg/dl           | 1160.0 (26.0)                    | 1140.0 (35.0)               | 0.958   |
| Insulin, uU/ml           | 12.4 (10.3)                      | 14.7 (9.1)                  | 0.171   |
| HbA1c, %                 | 6.4 (0.5)                        | 6.4 (0.9)                   | 0.548   |
| FFA, mmol/l              | 5090.0 (307.0)                   | 5325.0 (260.0)              | 0.541   |
| HOMA-IR                  | 3.5 (3.5)                        | 3.9 (2.7)                   | 0.166   |
| HOMA-beta                | 85.5 (70.3)                      | 99.6 (60.4)                 | 0.347   |
| Triglycerides, mg/dl     | 176.5 (143.0)                    | 217.5 (159.0)               | 0.177   |
| HDL, mg/dl               | 45.0 (16.0)                      | 42.0 (8.0)                  | 0.266   |
| LDL, mg/dl               | 115.3 (64.7)                     | 111.2 (39.9)                | 0.742   |
| CRP, mg/l                | 1.5 (1.8)                        | 1.5 (1.5)                   | 0.897   |
| eGFR, CKD-EPI equation   | 102.0 (23.0)                     | 95.0 (28.0)                 | 0.347   |

Data are expressed as median (IQR) unless indicated as number (%) with p-values from the Mann-Whitney U test or χ² analysis

Abbreviations: BMI Body mass index, SBP Systolic blood pressure, DBP Diastolic blood pressure, ALT Alanine aminotransferase, AST aspartate aminotransferase, GGT Gamma-glutamyl transferase, FFA Free fatty acids, HOMA-IR Homeostatic model assessment of insulin resistance, HDL High-density lipoprotein, LDL Low-density lipoprotein, CRP C-reactive protein, eGFR Estimated glomerular filtration rate

Ezetimibe did not significantly decrease fibrosis as measured by MRE or TE. In subgroup analyses, ezetimibe combination treatment was more effective in subjects with higher BMI, presence of T2DM, presence of sarcopenia, increased HOMA-IR, and increased baseline MRE.

Ezetimibe inhibits the absorption of dietary fat, including cholesterol, via NPC1-L1 [18]. Several previous animal models confirmed improvement in hepatic steatosis by ezetimibe. Improvement in triglyceride and free fatty acid by reduction of chylomicron synthesis in enterocytes [9], downregulation of SREBP-1c [19], or autophagy induction through AMPK activation and TFEB nuclear translocation [11] has been suggested as mechanisms by which ezetimibe contributes to improved hepatic steatosis and inflammation. However, previous studies on the effects of ezetimibe on hepatic steatosis in humans were inconsistent. Some showed that long-term treatment with ezetimibe improved histological indications of NAFLD (including steatosis) [20] whereas others, including Loomba et al., reported that while it reduced liver fat by 15.3% compared to placebo, it produced no significant improvement in hepatic steatosis by MRI-PDFF [12].

Our study demonstrated that significant improvement in hepatic steatosis occurred after treatment with the ezetimibe combination, and there may be several reasons for this result. First, this study was conducted in combination with a statin. Unlike statins, which inhibit cholesterol synthesis in the liver, ezetimibe targets gastrointestinal cholesterol absorption within the small intestine. In this process, ezetimibe can inhibit an increase in dietary cholesterol absorption, which can arise from the use of statins [21]. NPC1-L1 inactivation causes multiple lipid transport defects and had a protective effect against diet-induced hyperlipidemia in a mouse model [22]. Dietary cholesterol exacerbates hepatic steatosis and inflammation in an LDL receptor-deficient mice model of metabolic syndrome [23]. In addition, hepatic NPC1-L1 induces hepatic steatosis by participating in cholesterol reabsorption from bile to the liver, which can be inhibited by ezetimibe [24]. This suggests that ezetimibe may have a greater effect on NAFLD in combination with a statin. Similarly, ezetimibe showed a prominent effect in the protection of cardiovascular diseases in a large randomized controlled trial (IMPROVE-IT study) in combination with a statin [13].

Second, differences in the characteristics of the current study population may have the potential to influence the ezetimibe effect. Subgroup analysis showed that combination treatment was more effective in improving hepatic steatosis in individuals with higher BMI, presence of T2DM or sarcopenia, higher HOMA-IR, and increased baseline MRE (≥2.1, the median value in this study). Inhibiting absorption of dietary cholesterol by ezetimibe can contribute to improvement in inflammation and insulin resistance [25]. Considering this mechanism, ezetimibe efficacy may be greater in subjects with obesity, higher insulin resistance, and T2DM, who have poor dietary habits and who may have elevated dietary cholesterol. Similarly, ezetimibe was more effective in protecting against cardiovascular disease in subjects with T2DM in our meta-analysis [26] and the results were consistent with a separate propensity score-matched cohort study [27]. The ability of ezetimibe to improve inflammation by modulating autophagy and inflammasomes [11] may also
be greater in individuals with more advanced NASH or hepatic fibrosis. Therefore, a higher proportion of participants with insulin resistance in the presence of T2DM, which aggravates the inflammatory condition [28], could potentiate the effects of ezetimibe. Compared to the previous trial by Loomba et al. [12], our study treated all participants with the same dose of statin and recruited a higher proportion (>70%) of subjects with T2DM. These differences may be responsible for the apparently positive ezetimibe effect on NAFLD.

Although this study cannot fully explain its contributory mechanisms, there is evidence that ezetimibe changes the bile acid pool or microbiome [29]. Bile acid and microbiome are currently being studied as potentially important therapeutic targets for NASH or hepatic steatosis [30, 31]. The target protein of ezetimibe, NPC1-L1, known to be present in the intestine, was recently shown to be present in hepatocyte and Kupffer cells, affecting bile cholesterol reabsorption, which is expected to have a pleiotropic effect on hepatic steatosis [30, 32].

---

**Table 2** Ezetimibe plus rosuvastatin versus rosuvastatin monotherapy: longitudinal changes in liver fat and liver fibrosis

| Liver segment | Baseline | Post-treatment | p-value | Baseline | Post-treatment | p-value | p-value |
|---------------|----------|---------------|---------|----------|---------------|---------|---------|
| I             | 16.6 (8.2) | 11.3 (6.4) | < 0.001 | 13.2 (7.3) | 11.5 (7.0) | 0.042 | 0.012 |
| II            | 16.7 (8.4) | 10.8 (6.0) | < 0.001 | 14.2 (7.4) | 11.8 (7.4) | 0.006 | 0.012 |
| III           | 17.1 (8.3) | 11.2 (6.6) | < 0.001 | 14.2 (7.6) | 11.9 (7.2) | 0.012 | 0.011 |
| IV (A)        | 18.0 (8.4) | 12.4 (6.7) | < 0.001 | 15.3 (7.9) | 12.5 (7.5) | 0.004 | 0.038 |
| IV (B)        | 18.9 (8.8) | 13.4 (7.5) | < 0.001 | 15.9 (7.3) | 13.2 (7.8) | 0.003 | 0.051 |
| V             | 18.9 (8.7) | 12.9 (6.7) | < 0.001 | 16.1 (7.6) | 13.2 (7.8) | 0.002 | 0.029 |
| VI            | 18.1 (8.3) | 12.4 (6.7) | < 0.001 | 14.5 (7.5) | 11.7 (7.6) | 0.001 | 0.039 |
| VII           | 18.6 (7.9) | 12.5 (6.5) | < 0.001 | 15.2 (7.2) | 12.4 (7.3) | 0.001 | 0.018 |
| VIII          | 19.6 (8.5) | 13.5 (7.2) | < 0.001 | 16.6 (7.4) | 13.6 (7.9) | 0.002 | 0.028 |
| MRI-PDFF average, % | 18.1 (8.2) | 12.3 (6.4) | < 0.001 | 15.0 (7.3) | 12.4 (7.4) | 0.003 | 0.020 |
| MRE, kPa      | 2.0 (0.5) | 2.1 (0.5) | 0.507 | 2.2 (0.4) | 2.2 (0.7) | 0.539 | 0.898 |

Magnetic resonance imaging proton density fat fraction (MRI-PDFF) with colocalized MRI measurements was used in the longitudinal liver fat mapping. Longitudinal changes in liver fibrosis were measured using magnetic resonance elastography (MRE). Data are expressed as mean (SD) or difference with p-values from paired t-test or per-protocol analysis. Abbreviations: MRI-PDFF magnetic resonance imaging proton density fat fraction, MRE magnetic resonance elastography.

---

![Fig. 2](image-url)  
**Fig. 2** Liver fat (%) by MRI-PDFF at baseline and end of treatment by regimen.

Absolute mean difference; 3.2%, p=0.020
Progression from simple steatosis to NASH or fibrosis is of clinical importance. It is known that oxidative stress plays an important role in the progression of inflammation and fibrosis [33]. Ezetimibe has been shown to inhibit these actions through increased Nrf2 activity [34] or induction of autophagy flux [11]. In the markers related to inflammation that were evaluated in this study, only IL-18, an inflammasome-related cytokine, was significantly reduced in the ezetimibe combination group. IL-18 is involved in the inflammatory process and plays an important role in NAFLD [35]. This suggests that ezetimibe treatment may contribute to improvement in hepatic steatosis and the inflammatory process by reducing IL-18. Furthermore, fibrosis progression was evaluated by pre- and post-MRE and TE, and no significant improvement in fibrosis was found in either group. It may be that a 6-month treatment period is not long enough to show a difference. Considering the long progression from steatosis to fibrosis, further study is needed to understand the long-term effect of ezetimibe on hepatic fibrosis. On the other hand, there are clinical studies suggesting that statins are associated with a lower risk of hepatic fibrosis. Ciardullo et al. reported the lower odds of advanced fibrosis in statin users [36]. Statins may play an important role in NASH management in terms of reducing inflammation and oxidative stress [37]. Combination of statin and ezetimibe may contribute to the improvement of NAFLD by various mechanisms.

In this study, we also report improvement of hepatic steatosis in the monotherapy group. Previous studies of statins have shown inconsistent results regarding their potential contribution. Dongiovanni et al. reported the protective role of statins in individuals at risk of non-alcoholic steatohepatitis [38]. However, in a study by Nelson et al. [7], statin use was not associated with improvement in hepatic steatosis. In the current study,
participants received counseling and education regarding healthy lifestyles and behavior modification. It is possible that this led to improvement of hepatic steatosis even in the statin alone group. A second possibility is that increased public interest in NAFLD contributed to better self-care by study participants. Since the study did not employ a placebo group, it is not possible to state definitively whether statin monotherapy improved hepatic steatosis. We observed significant weight loss in both study arms, and this may have accounted for some of the changes we observed in liver fat. All patients with NAFLD should receive lifestyle education; however, individual response to this may be different outside of a trial. Therefore, our results can be attenuated in the real-world setting. However, the main results of the current study that the combination of ezetimibe and rosuvastatin was significantly superior to rosuvastatin monotherapy in improving steatosis measured...
by MRI-PDFF suggest a beneficial role of ezetimibe in improving NAFLD.

There are several limitations in this study. Baseline hepatic steatosis was marginally lower in the monotherapy group compared to the combination group. Liver biopsy, the gold standard to measure hepatic steatosis or fibrosis, was not performed. In this study, it was confirmed that co-administration of ezetimibe with statin contributed to the improvement of hepatic steatosis. However, further evidence should be implemented to confirm clinically relevant effects of ezetimibe on hepatic fibrosis or clinical outcomes in NASH patients. The small sample size and short observation period of 6 months were not sufficient to determine whether concomitant use of ezetimibe could lead to improvement in hepatic fibrosis. We enrolled patients with hepatic steatosis documented by abdominal ultrasound, and one subject in each group excluded having no documented hepatic steatosis (<5% on MRI-PDFF) at baseline. We performed the final analysis as a per-protocol analysis, which may lead to an increased risk of selection bias. The open-label design implies that the interpretation of the results of this study should be made with caution. On balance, there are several notable strengths. First, we used the MRI-PDFF method, recognized as a highly reliable method of measuring hepatic steatosis [12], to evaluate the primary outcome. PDFF methods are reported to have higher diagnostic performance in noninvasive detection of hepatic steatosis in patients with NAFLD than CAP techniques [39], which might explain the discrepant findings regarding the degree of improvement in hepatic steatosis measured by PDFF or CAP. Second, we confirmed the effect of ezetimibe in combination with a statin, which is widely prescribed in the clinical environment. Third, we utilized MRE to confirm the effect on progression of hepatic fibrosis. We also measured by TE, another method to evaluate hepatic steatosis and fibrosis. Although the difference between groups was unclear, significant improvement in hepatic steatosis was confirmed only in the ezetimibe combination group. Fourth, in subgroup analyses, it was possible to identify characteristics of participants in whom ezetimibe acted more effectively on hepatic steatosis.

Conclusions
In conclusion, the use of ezetimibe in combination with rosuvastatin significantly improved hepatic steatosis in patients with NAFLD. Individuals with higher BMI, T2DM, insulin resistance, and severe liver fibrosis were likely to be good responders to ezetimibe treatment. These data indicate that ezetimibe plus rosuvastatin is a safe and effective therapeutic option to treat patients with NAFLD and dyslipidemia.

Abbreviations
ALT: Alanine aminotransferase; AST: Aspartate aminotransferase; BMI: Body mass index; CAP: Controlled attenuation parameter; CRP: C-reactive protein, DM: Diabetes mellitus; ESSENTIAL: Effects of Statin Monotherapy and Statin/ Ezetimibe Combination Therapy on Non-alcoholic Steatohepatitis in patients with Hyperlipidemia and Fatty Liver; HDL-C: High-density lipoprotein cholesterol; HOMA-IR: Homeostatic model assessment insulin resistance; IL: Interleukin; LDL-C: Low-density lipoprotein cholesterol; LSM: Liver stiffness measurement; MRE: Magnetic resonance elastography; MRI-PDFF: Magnetic resonance imaging-derived proton density fat fraction; NAFLD: Non-alcoholic fatty liver disease; NASH: Non-alcoholic steatohepatitis; NPC-1 L1: Niemann-Pick C1-like 1; ROI: Regions of interest; SMI: Skeletal muscle mass index; SREBP-1c: Sterol regulatory element-binding protein-1c; T2DM: Type 2 diabetes; TE: Transient elastography; WC: Waist circumference.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12916-022-02288-2.

Acknowledgements
We acknowledge the clinical research coordinator, Jihyun Cheon. Caron Modes, Evolved Editing, LLC, provided editorial assistance. She had no role in the study design, implementation, or analysis.

Authors’ contributions
YC and Y-H.L conceived and designed the study. Y.C, M.L, B-W.L, E.S.K, B-S.C, and Y-H.L recruited the patients and carried out data collection in human studies. H.R and J-Y.C measured liver images. Y.C, Y.K, and Y-H.L analyzed and interpreted the data. All authors critically discussed the study design, data, and manuscript for the scientific content. Y.C, H.R, and Y-H.L wrote the manuscript, with contribution from all authors. The authors read and approved the final manuscript.

Funding
This research was supported by grants of Yuhan Corporation and the Korea Health Technology R&D Project through the Korea Health Industry Development Institute (KHIDI), funded by the Ministry of Health & Welfare, Republic of Korea (HI14C1324). Funding agencies did not affect the collection, analysis, or presentation of the data.

Availability of data and materials
The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

Declarations
Ethics approval and consent to participate
All participants provided written informed consent and the Ethics Committee of the Yonsei University College of Medicine approved this study (4-2017-1168), which conforms to the ethical principles of the 1975 Declaration of Helsinki.

Consent for publication
All participants granted consent for the publication of the de-identified result.

Competing interests
The authors declare that they have no competing interests.

Author details
1Department of Endocrinology and Metabolism, Inha University School of Medicine, Incheon, Republic of Korea. 2Graduate School, Yonsei University College
References

1. Angulo P, Kleiner DE, Dam-Larsen S, Adams LA, Bjornsson ES, Charatcharoenwitthaya P, et al. Liver fibrosis, but no other histologic features, is associated with long-term outcomes of patients with nonalcoholic fatty liver disease. Gastroenterology. 2015;149(2):389–397.e310.

2. Wagenknecht LE, Zaccaro D, Espeland MA, Karter AJ, O’Leary DH, Haﬀner SM. Diabetes and progression of carotid atherosclerosis: the insulin resistance atherosclerosis study. Arterioscler Thromb Vasc Biol. 2003;23(8):1035–41.

3. Mach F, Baigent C, Catapano AL, Koskinas KC, Casula M, Badimon L, et al. 2019 ESC/EAS Guidelines for the management of dyslipidaemias: lipid modification to reduce cardiovascular risk. Eur Heart J. 2019;41(11):111–88.

4. Ramkumar S, Raghu Nathan A, Raghu Nathan S. Statin therapy: review of safety and potential side effects. Acta Cardiol Sin. 2016;32(6):631.

5. Ioannou GN. The role of cholesterol in the pathogenesis of NASH. Trends Endocrinol Metab. 2016;27(2):84–95.

6. Ekstedt M, Franzén LE, Mathiesen UL, Holmqvist M, Boderman G, Rechagias S. Nonalcoholic fatty liver disease and chronically elevated liver enzymes: a histopathological follow-up study. J Hepatol. 2007;47(1):135–41.

7. Nelson A, Torres DM, Morgan AE, Fincke B, Harrison SA. A pilot study using simvastatin in the treatment of nonalcoholic steatohepatitis: a randomized placebo-controlled trial. J Clin Gastroenterol. 2009;43(10):990–4.

8. Takez H, Doi H, Okado T, Hashimoto T, Goto Y, Kimura G. Eﬀects of ezetimibe on visceral fat in the metabolic syndrome: a randomized controlled study. Eur J Clin Investig. 2012;42(12):1287–94.

9. Deushi M, Nomura M, Kawakami A, Haraguchi M, Ito M, Okazaki M, et al. Ezetimibe ameliorates steatohepatitis via AMP activated protein kinase–TFEB-mediated activation of autophagy and NLRP3 inﬂammasome inhibition. Autophagy. 2017;13(10):1767–81.

10. Ciardullo S, Perseghin G. Statin use is associated with lower prevalence of bile cholesterol reabsorption: hepatic NPC1L1-exacerbated steatosis and decreased VLDL-TG secretion in mice fed a high-fat diet. Lipids Health Dis. 2015;14(12):239–50.

11. Cannon CP, Blazing MA, Giugliano RP, McCagg A, White JA, Theroux P, et al. Ezetimibe added to statin therapy after acute coronary syndromes. N Engl J Med. 2015;372(25):2387–97.

12. Korean guidelines for the management of dyslipidemia 4th ed. In: Committee of clinical practice guideline of the Korean Society of Lipid and Atherosclerosis (KOSLA). 2018.

13. Gu J, Philo L, Nguyen P, Hofflich H, Hernandez C, Bettencourt R, et al. Sitagliptin vs. placebo for non-alcoholic fatty liver disease: a randomized controlled trial. J Hepatol. 2016;65(2):369–76.

14. Kim KM, Lim S, Choi SH, Kim JH, Shin CS, Park KS, et al. Cardiometabolic implication of sarcopenia: The Korean National Health and Nutrition Examination Study (KHANES) 2008–2010. JC Metab Endocr. 2014;4;63–9.

15. Loomba R. NpR-PPIF treatment response criteria in nonalcoholic steatohepatitis. Hepatology. 2020;73:881–3.

16. Labonté ED, Camarota LM, Rojas KC, Jandacek RJ, Gilham DE, Davies JP, et al. Reduced absorption of saturated fatty acids and resistance to diet-induced obesity and diabetes by ezetimibe-treated and Npc1l1−/− mice. Am J Physiol Gastrointest Liver Physiol. 2008;294(4):G776–83.

17. Murakata T, Aski K, Iwasaki T, Shinoda K, Nakamura A, Aburatani H, et al. Ezetimibe decreases SREBP-1c expression in liver and reverses hepatic insulin resistance in mice fed a high-fat diet. Metabolism. 2011;60(5):617–28.

18. Park H, Shima T, Yamaguchi K, Mitsuyoshi H, Minami M, Yasui K, et al. Eﬃcacy of long-term ezetimibe therapy in patients with nonalcoholic fatty liver disease. J Gastroenterol. 2011;46(1):101–7.

19. Miettinen TA, Kylling H. Synthesis and absorption markers of cholesterol in serum and lipoproteins during a large dose of statin treatment. Eur J Clin Invest. 2003;33(11):976–82.

20. Davies JP, Scott C, Oishi K, Liapas A, Ioannou YA. Inactivation of NPC1L1 causes multiple lipid transport defects and protects against diet-induced hypercholesterolemia. J Biol Chem. 2005;280(13):12710–20.

21. Subramanian S, Goodspeed L, Wang S, Kim J, Zeng L, Ioannou GN, et al. Dietary cholesterol exacerbates hepatic steatosis and inﬂammation in obese LDL receptor-deﬁcient mice. J Lipid Res. 2011;52(9):1626–35.

22. Toyoda Y, Takada T, Umezawa M, Tomura F, Yamashita Y, Takeda K, et al. Identiﬁcation of hepatic NPC1L1 as an NAFLD risk factor evidenced by ezetimibe-mediated steatosis prevention and recovery. FASEB J. 2019;33(2):283–95.

23. Cho Y, Kim R-H, Park H, Wang JH, Lee H, Kang E. Eﬀect of ezetimibe on glucose metabolism and inﬂammatory markers in adipose tissue. Biomedicines. 2020;8(11):512.

24. Hong N, Lee Y-H, Tsujita K, Gonzalez JA, Kramer CM, Kovarik T, et al. Comparison of the eﬀects of ezetimibe-statin combination therapy on major adverse cardiovascular events in patients with and without diabetes: a meta-analysis. Endocrinol Metab. 2018;33(2):219–27.

25. Lee Y-H, Hong N, Lee CI, Park SH, Lee B-W, Cha B-S, et al. Diﬀerential association between ezetimibe-simvastatin combination with major adverse cardiovascular events in patients with or without diabetes: a propensity score-matched cohort study. Sci Rep. 2018;8(1):1–7.

26. Lee B-W, Lee Y-H, Park C-Y, Rhee E-J, Lee W-Y, Kim N-H, et al. Non-alcoholic fatty liver disease in patients with type 2 diabetes mellitus: a position statement of the Fatty Liver Research Group of the Korean Diabetes Association. Diabetes Metab J. 2020;44(3):382–401.

27. Caty E, Pachikian BD, Salazar N, Neyrnck AM, Cani PD, Delzenne NM. Ezetimibe and simvastatin modulate gut microbiota and expression of genes related to cholesterol metabolism. Life Sci. 2015;132:77–84.

28. Wiest R, Albillos A, Trauner M, Bajaj JS, Jalan R. Targeting the gut-liver axis in nonalcoholic steatohepatitis via AMP activated protein kinase-TEFβ-mediated activation of autophagy and NLRP3 inﬂammasome inhibition. Autophagy. 2012;6(12):1287–94.

29. Deushi M, Nomura M, Kawakami A, Haraguchi M, Ito M, Okazaki M, et al. Ezetimibe ameliorates steatohepatitis via AMP-activated protein kinase-TFEB-mediated inhibition of bile cholesterol reabsorption: hepatic NPC1L1-exacerbated steatosis and decreasing VLDL-TG secretion in mice fed a high-fat diet. Lipids Health Dis. 2019;18(1):1–10.

30. Albano E, Ottolanc E, Occhino G, Reale E, Vidal Mi. Role of oxidative stress in the progression of non-alcoholic steatohepatitis: Aliment Pharmac Ther. 2005;22:71–3.

31. Vuppalanchi R, Noureddin M, Alkhoum N, Sanjay Al Therapeutic pipeline in nonalcoholic steatohepatitis. Nat Rev Gastroenterol Hepatol. 2021;18(6):373–92.

32. Toyoda Y, Takada T, Yamashita Y, Suzuki H. Pathophysiological importance of bile cholesterol reabsorption: hepatic NPC1L1-exacerbated steatosis and decreasing VLDL-TG secretion in mice fed a high-fat diet. Lipids Health Dis. 2019;18(1):1–10.

33. Albano E, Mottaran E, Occhino G, Reale E, Vidal Mi. Role of oxidative stress in the progression of non-alcoholic steatohepatitis: Aliment Pharmac Ther. 2005;22:71–3.

34. Han DH, Nam KT, Park JS, Kim SH, Lee M, Kim G, et al. Ezetimibe, an NPC1L1 inhibitor, is a potent Nrf2 activator that protects mice from diet-induced nonalcoholic steatohepatitis. Free Radic Biol Med. 2016;99:520–32.

35. Gueret-Kabiber M, Yang S, Lin HZ, Schwenz K, Krzych U, Devill AL. Altered hepatic lymphocyte subpopulations in obesity-related murine fatty livers: potential mechanism for sensitization to liver damage. Hepatology. 2000;31(3):633–40.

36. Ciardullo S, Perseghin G. Statin use is associated with lower prevalence of advanced liver fibrosis in patients with type 2 diabetes. Metabolism. 2022;131:154752.

37. El-Din SHS, El-Lakkany NM, El-Naggar AA, Hammam OA, Abd El-Latif HA, Ain-Shekaa AA, et al. Ezetimibe and/or β-carotene on non-alcoholic fatty liver in rats. Res Pharm Sci. 2015;10(4):275.

38. Dongiovanni P, Petta S, Mannisto V, Mancino VM, Pimpone R, Karja V, et al. Ezetimibe and simvastatin modulate gut microbiota and expression of genes related to cholesterol metabolism. Life Sci. 2015;132:77–84.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.