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# Inhibition of DHCR24 activates LXR $\alpha$ to ameliorate hepatic steatosis and inflammation

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DATA

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# Abstract

Liver X receptor (LXR) agonism has theoretical potential for treating NAFLD/NASH, but synthetic agonists induce hyperlipidemia in preclinical models. Desmosterol, which is converted by  $\Delta$ 24-dehydrocholesterol reductase (DHCR24) into cholesterol, is a potent endogenous LXR agonist with anti-inflammatory properties. We aimed to investigate the effects of DHCR24 inhibition on NAFLD/NASH development. Here, by using APOE\*3-Leiden. CETP mice, a well-established translational model that develops diet-induced human-like NAFLD/NASH characteristics, we report that SH42, a published DHCR24 inhibitor, markedly increases desmosterol levels in liver and plasma, reduces hepatic lipid content and the steatosis score, and decreases plasma fatty acid and cholesteryl ester concentrations. Flow cytometry showed that SH42 decreases liver inflammation by preventing Kupffer cell activation and monocyte infiltration. LXRa deficiency completely abolishes these beneficial effects of SH42. Together, the inhibition of DHCR24 by SH42 prevents diet-induced hepatic steatosis and inflammation in a strictly LXR<sub>α</sub>-dependent manner without causing hyperlipidemia. Finally, we also showed that SH42 treatment decreased liver collagen content and plasma alanine transaminase levels in an established NAFLD model. In conclusion, we anticipate that pharmacological DHCR24 inhibition may represent a novel therapeutic strategy for treatment of NAFLD/NASH.

Keywords desmosterol; Kupffer cell; liver X receptor; nonalcoholic steatohepatitis;  $\Delta$ 24-dehydrocholesterol reductase Subject Categories Digestive System; Metabolism

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# Introduction

Nonalcoholic fatty liver disease (NAFLD) comprises a spectrum of diseases ranging from simple hepatic steatosis to nonalcoholic steatohepatitis (NASH). The latter is characterized by steatosis, hepatocellular damage, and inflammatory cell infiltration with or without fibrosis. Hepatic steatosis is defined as the accumulation of primarily neutral lipids such as triacylglycerols (TAG) and cholesteryl esters (CE) in the form of lipid droplets within hepatocytes, as well as nonparenchymal liver cells, including Kupffer cells (KCs; Berlanga et al, 2014). Hepatic steatosis results from an imbalance between lipid uptake, synthesis, secretion, and lipolysis and is usually associated with dyslipidemia, insulin resistance, hypertension, type 2 diabetes, and obesity (Chalasani et al, 2018). The essential trigger for the transition of simple steatosis to NASH is not yet completely elucidated. Despite the high global prevalence of NAFLD (32.4%) in the general population (Riazi et al, 2022), no FDA-approved medication is available yet for the treatment of NAFLD/NASH. In fact, lifestyle adjustment is still the main clinical intervention.

Inflammation contributes to the transition of simple steatosis to NASH and liver-resident KCs play a pivotal role in NAFLD/NASH pathogenesis (Baffy, 2009). Once activated during NAFLD progression, KCs produce proinflammatory factors, such as monocyte chemoattractant protein 1 (MCP-1) and tumor necrosis factor

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(TNF), leading to increased hepatic monocyte recruitment and inhibition of canonical insulin signaling, further aggravating liver injury and steatosis (Aparicio-Vergara et al, 2013; Wandrer et al, 2020). In addition, KC-derived pro-fibrinogenic factors increase collagen production by hepatic stellate cells, generating a vicious circle that exacerbates NAFLD and inflammation, and derive progression towards NASH (Tosello-Trampont et al, 2012; Schuster et al, 2018). Thereby, selective depletion of KCs from the liver alleviates hepatocellular damage and prevents diet-induced hepatic steatosis and insulin resistance (Huang et al, 2010; Lanthier et al, 2011). Furthermore, consumption of a cholesterol-rich diet causes cholesterol accumulation in KCs to yield foamy inflammatory KCs, which directly contribute to liver inflammation (Bieghs et al, 2013). Recently, we and others showed that NAFLD/NASH impaired KC self-renewal and induce KC death, thus reducing embryonicallyderived liver-resident KCs (Remmerie et al, 2020; Seidman et al, 2020; Tran et al, 2020). This KC niche is replenished through recruitment, differentiation, and proliferation of monocyte-derived macrophages, which are, however, reported to be more inflammatory than KCs and to contribute to the aggravation of liver inflammation and damage (Remmerie *et al*, 2020; Tran *et al*, 2020).

Liver X receptors (LXRs), that is, the LXR $\alpha$  and LXR $\beta$  isoform, are essential (oxy)sterol-activated transcription factors involved in lipid metabolism and immune responses (Bensinger & Tontonoz, 2008; Ito *et al*, 2015). LXR $\alpha$  is abundantly expressed in liver, adipose tissue, and macrophages, while  $LXR\beta$  is ubiquitously expressed (Prufer & Boudreaux, 2007). In macrophages, LXRa directly promotes reverse cholesterol transport via up-regulating ATP binding cassette (ABC) A1 (Venkateswaran et al, 2000). Thus, LXRadeficiency impairs cholesterol efflux and is associated with increased atherosclerosis (Bischoff et al, 2010; Ishibashi et al, 2013). Simultaneously, LXRs exert anti-inflammatory effects in immune cells (Joseph et al, 2004), suppress KC activation, and protect against hepatic injury (Wang et al, 2006). In addition, LXRs promote the formation of long-chain polyunsaturated fatty acids (PUFAs), for example, eicosapentaenoic acid and docosahexaenoic acid, which have anti-inflammatory activities (Li et al, 2013; Korner et al, 2019). The simultaneous regulation of lipid metabolism and inflammation serves LXRs as a potential drug target for NAFLD/NASH treatment (Bensinger & Tontonoz, 2008; Ito et al, 2015). However, at present no selective LXR agonists exist for the clinical treatment of NAFLD/ NASH. This is mainly due to the unfavorable effects of pharmacological LXR activation on sterol regulatory element-binding proteins (SREBPs)-induced lipogenesis, resulting in elevated atherogenic low-density lipoprotein (LDL) cholesterol and triglycerides. Therefore, rather than alleviating liver lipid levels, most synthetic LXR agonists actually cause hepatic steatosis and hypertriglyceridemia (Schultz et al, 2000; Grefhorst et al, 2002). Moreover, some synthetic LXR agonists have been reported to cause neutropenia due to the downregulation of neutrophil production in bone marrow and stimulation of their clearance by macrophages within peripheral tissues (Kirchgessner et al, 2016). Taken together, a successful drug candidate for the treatment of NASH through LXR activation, should be LXR selective, not induce sterol regulatory element-binding transcription factor 1 (Srebf1), and not cause neutropenia.

Previously, we showed that desmosterol increases LXR target genes while inhibiting SREBP target genes in macrophages (Spann *et al*, 2012; Muse *et al*, 2018). In addition, we reported that synthetic

SH42 inhibits  $\Delta$ 24-dehydrocholesterol reductase (DHCR24), an important enzyme intertwining the Bloch and Kandutsch-Russell pathways of distal cholesterol biosynthesis, and causes accumulation of desmosterol (Nes, 2011). Accordingly, we have recently reported that inhibition of DHCR24 induces LXR activation through the accumulation of desmosterol, promoting the resolution of inflammation without affecting Srebf1c expression in macrophages (Muller et al, 2017; Korner et al, 2019). Taken together, we hypothesized that inhibiting DHCR24 increases desmosterol levels and subsequently induces LXR activation in KCs, thereby suppressing inflammation, without inducing lipid synthesis via Srebf1. Using our published DHCR24 inhibitor SH42 (Muller et al, 2017), we aimed to investigate the potential therapeutic effects of DHCR24 inhibition on diet-induced NAFLD development in APOE\*3-Leiden.CETP mice, a well-established humanized mouse model for the study of (cardio) metabolic diseases.

# Results

# Inhibition of DHCR24 by SH42 markedly increases liver desmosterol levels and ameliorates hepatic steatosis

To assess the effects of DHCR24 inhibition on hepatic steatosis, *E3L.CETP* mice were fed a HFCD while being treated with vehicle or the synthetic DCHR24 inhibitor SH42 (Muller *et al*, 2017; Korner *et al*, 2019) for a period of 8 weeks (Fig EV1A). SH42 treatment did not affect food intake (Fig EV1B), while temporarily preventing HFCD-induced body weight gain as compared to the control group (Fig EV1C). After 8 weeks of treatment, body weight and body composition, that is, lean body mass (Fig EV1D) and fat body mass (Fig EV1E) of SH42-treated mice, were comparable to that of the control group. In addition, the weight of various tissues (i.e., liver, white adipose tissue, kidney, heart, lung, spleen, and brown adipose tissue) was unchanged by SH42 treatment (Fig EV1F). As anticipated, SH42 markedly increased hepatic desmosterol levels (10-fold, Figs 1A and EV1G).

Hepatic steatosis was evaluated by HE staining and scored as described previously (Liang et al, 2014). As compared to the control treatment, SH42 treatment ameliorated diet-induced hepatic steatosis (Fig 1B), as evident by a clear reduction of the hepatic steatosis score (-58%, Fig 1C), and liver lipid area stained by Oil Red O (-53%, Fig 1B and D). We next analyzed liver lipid profiles by comprehensive lipidomic analysis. Firstly, we observed a clear alteration in the lipid class composition after SH42 treatment (Fig 1E). Specifically, SH42 treatment caused a relative reduction of TAG (-21%) and DAG (-22%), accompanied by a relative increase of the other lipid classes, including CER, PC, PE, and SM (Fig 1E and Appendix Fig S1). Volcano plot analysis of the lipid species concentrations of different lipidomes revealed the majority of significantly altered lipid species by SH42 treatment were down-regulated, and most of them were TAGs (Fig 1F). Consistently, SH42 tended to decrease hepatic concentrations of triacylglycerides (TAG; -39%; P = 0.06,  $P_{\text{adj}} = 0.08$ , Fig 1G) and diacylglycerides (DAG; -20%; P = 0.08,  $P_{\rm adj} = 0.08$ , Fig 1H). In addition, SH42 treatment reduced hepatic free fatty acids (FFA; -16%; P = 0.02,  $P_adj = 0.08$ , Fig 1I) and cholesterol ester (CE) levels  $(-25\%; P = 0.08, P_adj = 0.08, Fig 1J)$ . Despite these strong effects on hepatic lipid content, SH42 did not



Figure 1.

### Figure 1. Inhibition of DHCR24 by SH42 markedly increases liver desmosterol and ameliorates hepatic steatosis.

E3LCETP mice fed a high-fat high-cholesterol diet (HFCD) were treated with vehicle (Ctrl) or DHCR24 inhibitor SH42 (SH42) (n = 8 mice per group).

A, B After 8 weeks of treatment, mice were sacrificed and livers were collected (A) to measure desmosterol levels, and (B) to stain with hematoxylin and eosin (HE) and Oil Red O.

- C, D Hepatic steatosis was scored using the HE-stained slides and (D) lipid-positive area was quantified using the Oil Red O stained slides (*n* = 7 mice per group; two values were identified as outliers based on a Grubbs' test and removed from statistical analysis).
- E Liver lipid classes were analyzed (Appendix Fig S1), and the relative average abundance of lipid classes in each group are depicted as pie charts.
- F Volcano plot analysis of liver lipid species concentrations is shown.
- G–J Hepatic (G) TAG, (H) DAG, (I) FFA and (J) CE lipid class concentrations were summarized. Benjamini–Hochberg correction was used for multiple hypothesis testing and Benjamini–Hochberg adjusted *P*-values are presented (*P*\_adj).

Data information: Values are mean  $\pm$  SEM. Differences between two groups (SH42/Ctrl) were determined using a Mann–Whitney test. \*P < 0.05, \*\*\*P < 0.001 vs. control (ctrl). CE, cholesteryl esters; CER, ceramides; DAG, diacylglycerols; DCER, dihydroceramides; FFA, free fatty acids; HCER, hexosylceramides; LPC, lysophosphatidylcholines; LPE, lysophosphatidylethanolamines; SM, sphingomyelins; TAG, triacylglycerols. Scale bar: 100 or 200  $\mu$ m as indicated.

Source data are available online for this figure.

affect fasting plasma glucose, insulin, or homeostatic Model Assessment for Insulin Resistance (HOMA-IR) scores (Fig EV1H–J). Taken together, inhibition of DHCR24 by SH42 markedly increases liver desmosterol levels, accompanied by amelioration of diet-induced hepatic steatosis without marked effects on body composition and glucose homeostasis.

## Inhibition of DHCR24 by SH42 prevents Kupffer cell activation and reduces immune cell infiltration into the liver

We next evaluated the effect of DHCR24 inhibition on the hepatic inflammation in mice treated with SH42 or vehicle for 8 weeks by immunohistochemistry. SH42 treatment significantly reduced the hepatic F4/80 content (-29%, Fig 2A and B), as well as the number of hepatic crown-like structures of macrophages surrounding dying hepatocytes (-79%, Fig 2C), which is a prominent feature of NASH development (Itoh *et al*, 2013).

Flow cytometry on MACS-purified hepatic leukocytes revealed that 4 weeks of SH42 treatment tended to decrease total hepatic leukocytes (-37%; P = 0.09, Fig 2D). Given the critical role of KCs in the progression of NASH (Huang et al, 2010; Lanthier et al, 2011), we next explored the effects of DHCR24 inhibition on KCs. We observed a significant increase in total KCs upon SH42 treatment (+44%, Fig 2E and F). Although SH42 did not significantly reduce MHCII<sup>+</sup>/CD11c<sup>+</sup>-activated KCs (Fig 2G and H), MHCII<sup>-</sup>/CD11c<sup>-</sup> resting KCs were significantly increased (+21%, Fig 2G and I), indicative of prevented KC activation. In addition, SH42 treatment reduced monocytes in both liver (-79%, Fig 2J) and blood (-43%; P = 0.06, Fig EV2A and B), and decreased hepatic neutrophils (-50%, Fig 2K) without affecting circulating neutrophils (Fig EV2A and C). Together, these data indicate that SH42 treatment prevents KC activation, limits hepatic immune cell recruitment, and dampens hepatic inflammation.

# Inhibition of DHCR24 by SH42 does not increase circulating lipids

Consistent with the potent increase in hepatic desmosterol levels, SH42 also markedly increased plasma desmosterol levels from undetectable levels (< 0.5  $\mu$ g ml<sup>-1</sup>) to 3.1  $\pm$  0.4  $\mu$ g ml<sup>-1</sup> (Fig 3A). Since synthetic LXR agonists usually induce lipogenesis and hypertriglyceridemia as unwanted effects (Grefhorst *et al*, 2002;

Kirchgessner *et al*, 2016), we next determined the effect of 8 weeks of SH42 treatment on circulating lipid levels using quantitative comprehensive lipidomic analysis. Analysis of the plasma lipidome revealed that SH42 treatment relatively decreased circulating CE while relatively increasing lactosylceramides (LCER), phosphatidyl-choline (PC), and phosphatidylethanolamine (PE; Appendix Fig S2). With respect to absolute lipid concentrations, SH42 did not affect plasma levels of total TAG (Fig 3B) and DAG (Fig 3C), while significantly decreasing plasma levels of FFA (-16%, Fig 3D) and CE (-24%; P = 0.08, Fig 3E). These data imply that inhibition of DHCR24 by SH42 increases plasma desmosterol levels and decreases FFA and CE levels, importantly without inducing hypertriglyceridemia.

# The therapeutic effects of DHCR24 inhibition by SH42 on hepatic steatosis are strictly dependent on LXR $\alpha$

Based on previous reports (Spann et al, 2012; Muse et al, 2018; Korner et al, 2019), we hypothesized that the therapeutic effects of SH42 were attributed to LXR activation triggered by increased desmosterol levels as a result of DHCR24 inhibition. Since LXRa plays a crucial role in both lipid metabolism (Peet et al, 1998) and macrophage/KC homeostasis (Bischoff et al, 2010; Ishibashi et al, 2013; Endo-Umeda et al, 2018), we next evaluated the effects of 4 weeks of SH42 treatment in HFCD-fed LXRa-deficient mice to evaluate the importance of  $LXR\alpha$  in the observed protective effects of SH42 in ameliorating hepatic steatosis and inflammation. SH42 treatment had no effect on body weight (Fig EV3A), body composition (Fig EV3B and C), or liver weight (Fig EV3D) in LXRα-deficient mice. Consistent with previous findings that  $LXR\alpha$ -deficient mice are more susceptible to high-fat diet-induced hepatic steatosis and inflammation (Endo-Umeda et al, 2018; Endo-Umeda & Makishima, 2019), we observed significant hepatic steatosis after 4 weeks of HFCD (Fig 4A and B). Although SH42 treatment largely increased plasma desmosterol levels also in LXRa-deficient mice (+80%, Fig EV3E), it failed to improve hepatic steatosis (Fig 4A) as evidenced by an unchanged steatosis score (Fig 4B) and similar lipid-positive areas stained by Oil Red O in SH42 treated mice compared with control (Fig 4A and C). Liver lipidomic analysis further revealed that almost no lipid species was significantly changed upon SH42 treatment as shown by volcano plot analysis (Fig 4D). Accordingly, SH42 treatment showed no effects on hepatic concentrations



Figure 2.

#### Figure 2. Inhibition of DHCR24 by SH42 prevents Kupffer cell activation and reduces immune cell infiltration into the liver.

E3LCETP mice fed a HFCD were treated with vehicle (Ctrl) or DHCR24 inhibitor SH42 (SH42) (n = 8 mice per group). After 8 weeks of treatment, mice were killed and livers were collected to stain with F4/80.

- A, B Representative pictures of F4/80 staining are shown and (B) F4/80 positive area was quantified.
- C, D Hepatic crown-like structures as indicated by the arrows in Fig 2A were counted (n = 7 and 8 mice, respectively; one value was identified as an outlier based on a Grubbs' test and removed from statistical analysis). In another experiment after 4 weeks of treatment (n = 6 mice per group), fresh livers were collected to isolate and count (D) CD45<sup>+</sup> cells.
- E-K (E and F) Total Kupffer cells (KCs), (G and H) activated KCs, (G and I) resting KCs, (J) recruited monocytes, and (K) neutrophils in the liver were measured via flow cytometry.

Data information: Values are mean  $\pm$  SEM. Differences between two groups (SH42/Ctrl) were determined using a Mann–Whitney test. \*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001 vs. ctrl. Scale bar: 100  $\mu$ m.

Source data are available online for this figure.



## Figure 3. Inhibition of DHCR24 by SH42 does not increase circulating lipids.

E3LCETP mice fed a HFCD were treated with vehicle (Ctrl) or DHCR24 inhibitor SH42 (SH42) (n = 8 mice per group).

A After 8 weeks of treatment, blood was collected to measure desmosterol levels and for quantitative lipidomic analysis.

B-E Plasma (B) TAG, (C) DAG, (D) FFA and (E) CE lipid class concentrations were summarized.

Data information: Values are mean  $\pm$  SEM. Differences between two groups (SH42/Ctrl) were determined using a Mann–Whitney test. \*\*P < 0.01 vs. ctrl. CE, cholesteryl esters; DAG, diacylglycerols; FFA, free fatty acids; TAG, triacylglycerols.

Source data are available online for this figure.

of TAG (Fig 4E), DAG (Fig 4F), FFA (Fig 4G), or CE (Fig 4H) in the absence of LXR $\alpha$ .

# The therapeutic effects of DHCR24 inhibition by SH42 on Kupffer cell activation and hepatic monocyte infiltration are strictly dependent on LXR $\alpha$

Subsequently, we analyzed hepatic immune cells by flow cytometry after 4 weeks of treatment with SH42 in HFCD-fed LXR $\alpha$ deficient mice. In contrast to *E3L.CETP* mice, SH42 did not affect hepatic leukocytes (Fig 5A) in LXR $\alpha$ -deficient mice. Strikingly, SH42 did not affect KC numbers (Fig 5B and C), and did not prevent KC activation as evidenced by unchanged high levels of NASH-associated MHCII<sup>+</sup>CD11c<sup>+</sup>-activated KCs (Fig 5D and E) and MHCII<sup>-</sup>CD11c<sup>-</sup> resting KCs (Fig 5D and F). SH42 did not reduce recruited monocytes in liver (Fig 5G) but reduced hepatic neutrophils in LXR $\alpha$ -deficient mice (-42%; Fig 5H). Congruent with these data, circulating monocytes and neutrophils were not affected by SH42 treatment (Fig EV4A–C). Taken together, these data show that inhibition of DHCR24 by SH42 does not prevent diet-induced hepatic steatosis and inflammation in LXR $\alpha$ -deficient mice.



Figure 4.

#### Figure 4. The therapeutic effects of DHCR24 inhibition on hepatic steatosis are strictly dependent on LXRx.

LXR $\alpha$ -deficient mice fed HFCD were treated with vehicle (Ctrl) or DHCR24 inhibitor SH42 (SH42) (n = 10 and 11 mice, respectively).

After 4 weeks of treatment, mice were killed and liver samples were collected to stain with HE and Oil Red O.

- B, C Hepatic steatosis was scored using the HE-stained slides and (C) lipid-positive area was quantified using the Oil Red O stained slides.
- Volcano plot analysis of liver lipid species concentrations is shown.
- E-H Hepatic (E) TAG, (F) DAG, (G) FFA and (H) CE lipid class concentrations were summarized. Benjamini-Hochberg correction was used for multiple hypothesis testing and Benjamini-Hochberg adjusted P-values are presented (P\_adj). One outlier was identified and excluded in the hepatic CE analysis.

Data information: Values are mean  $\pm$  SEM. Differences between two groups (SH42/Ctrl) were determined using a Mann–Whitney test if not indicated otherwise. Scale bar: 100 µm or 200 µm as indicated. CE, cholesteryl esters; CER, ceramides; DAG, diacylglycerols; DCER, dihydroceramides; FFA, free fatty acids; HCER, hexosylceramides; LPC, lysophosphatidylcholines; LPE, lysophosphatidylethanolamines; PC, phosphatidylcholines; PE, phosphatidylethanolamines; SM, sphingomyelins; TAG, triacylglycerols. Source data are available online for this figure.

# Treatment with SH42 reduces hepatic crown-like structures, liver collagen content, and plasma alanine transaminase levels in an established NAFLD model

Translationally, we next investigated whether the inhibition of DHCR24 could rescue NASH progression. To this end, E3L.CETP mice were fed with a HFCD for 10 weeks first to establish NAFLD and then treated with vehicle or SH42 while still on HFCD for an additional 8 weeks (Fig 6A). SH42 treatment did not significantly influence either total body weight (Appendix Fig S7A) or composition (Fig EV5A-C). Although we did not observe any difference in liver weight (Fig EV5D) and hepatic steatosis (Fig 6B-D), SH42 treatment reduced hepatic crown-like structures (-89%; Fig 6E and F) without significant influence on the F4/80 positive area (Fig 6E and G), and ameliorated liver collagen content (-50%; Fig 6H and I). In addition, SH42 treatment reduced plasma levels of the liver injury marker plasma alanine transaminase (ALT; -42%; Fig 6J). These effects were accompanied by a robust increase in plasma desmosterol levels (> 400-fold; Fig EV5E).

# Discussion

As key regulators of metabolic and inflammatory signaling (Bensinger & Tontonoz, 2008; Ito et al, 2015), LXRs have taken center stage as potential therapeutic targets for the treatment of cardiometabolic diseases. However, undesirable side effects of pharmacological LXR activation, including hyperlipidemia via induction of Srebf1c expression and neutropenia (Grefhorst et al, 2002; Kirchgessner et al, 2016), have prevented clinical application. Interestingly, desmosterol has been reported to be an endogenous ligand of LXR while inhibiting SREBP activity (Muse et al, 2018). Therefore, enhancing endogenous desmosterol levels by targeting DHCR24 is already considered as a promising strategy for activating LXR transcription programs to combat atherosclerotic cardiovascular disease (Rodriguez-Acebes et al, 2009; Spann et al, 2012; Muse et al, 2018). In this study, we exploited the synthetic DHCR24 inhibitor SH42, developed by our group (Muller et al, 2017), for the treatment of hepatic steatosis and inflammation, that is, the two hallmarks of NAFLD/NASH development. We show that inhibition of DHCR24 induces marked increases in desmosterol levels in both liver and circulation, which is accompanied by decreased hepatic steatosis and inflammation. Experiments performed in LXRα-deficient mice demonstrated that the therapeutic effects of SH42 are strictly dependent on LXR $\alpha$  activation, most

KCs, based on our previous study showing that desmosterol acti-

vates LXR in macrophages but not in hepatocytes (Muse et al, 2018; Snodgrass et al, 2020). In addition, we previously showed that inhibition of DHCR24 by SH42 increased desmosterol and induced the expression of LXR target genes, including Abca1 and Abcg1 in RAW 264.7 macrophages (Korner et al, 2019). LXRα is highly expressed in KCs and the region of enhancers unique to KCs are enriched in LXRregulating sequence motifs (Lavin et al, 2014), indicating a role of LXRs in KC homeostasis. Accordingly, we here observed that about 95% of KCs in LXRa-deficient mice displayed markers indicative of activated KCs (i.e., MHCII<sup>+</sup>/CD11c<sup>+</sup> KCs) versus less than 25% of KCs in wild-type mice upon feeding the same HFCD, which is in agreement with our previous study in LXRa-deficient mice showing a robust increase in KC activation and hepatic inflammation (Endo-Umeda et al, 2018). Taken together, our study suggests that pharmacological inhibition of DHCR24 is a feasible way to prevent KC activation and alleviate liver inflammation through desmosterolinduced LXRa activation.

likely due to increased desmosterol, as we have previously shown

while SH42 prevented circulating monocyte recruitment and

increased MHCII<sup>-</sup>/CD11c<sup>-</sup> resting KCs in E3L.CETP mice only

(Fig 2G-J). This effect on monocytes/KCs is likely attributed to the

direct effects of desmosterol-induced LXRa activation in monocytes/

We found that inhibition of DHCR24 by SH42 markedly increased desmosterol levels in both E3L.CETP mice and LXRa-deficient mice,

that SH42 does not have intrinsic LXR affinity (Korner et al, 2019).

Importantly, we observed that inhibition of DHCR24 by SH42 prevented high-fat diet-induced hepatic steatosis in E3L.CETP mice (Fig 1) but not in LXR $\alpha$ -deficient mice (Fig 4). We excluded the potential contribution of altered systemic glucose homeostasis in this process as evidenced by unchanged body composition, circulating glucose and insulin levels, as well as HOMA-IR scores in both groups (Fig EV1). We reasoned that the alleviated liver inflammation may largely contribute to the reduction of hepatic steatosis. The feed-forward loop between lipotoxic hepatocytes and proinflammatory immune cells, especially KCs, promotes NAFLD/NASH progression (Hirsova & Gores, 2015). Our study strongly suggests that this vicious circle can be abrogated by desmosterol-induced LXRa activation in liver-resident KCs. Consistently, both prevention of KC activation and depletion of KCs reduces liver inflammation (Bieghs et al, 2012) and alleviates hepatic steatosis (Huang et al, 2010; Lanthier et al, 2011). Therefore, the alleviated liver inflammation could be the direct effect of desmosterol in macrophages/KCs, which subsequently prevents hepatic steatosis. Additionally, the alleviation of liver inflammation has been shown to have multiple



Figure 5. The therapeutic effects of DHCR24 inhibition on liver inflammation are strictly dependent on LXRa.

LXRa-deficient mice fed HFCD were treated with vehicle (Ctrl) or DHCR24 inhibitor SH42 (SH42) (n = 10 and 11 mice, respectively).

A After 4 weeks of treatment, mice were killed and fresh liver samples were collected to isolate and count CD45<sup>+</sup> cells.

B-H (B and C) Total Kupffer cells (KCs), (D and E) activated KCs, (D and F) resting KCs, (G) recruited monocytes, and (H) neutrophils in the liver were measured via flow cytometry.

Data information: Values are mean  $\pm$  SEM. Differences between two groups (SH42/Ctrl) were determined using a Mann–Whitney test. \*\*P < 0.01 vs. ctrl. Source data are available online for this figure.

effects, including reduced fatty acid uptake, VLDL secretion, and lipid oxidation (Liu *et al*, 2014), which could be the cause of a reduction in liver lipids by SH42 treatment, and needs to be further investigated.

Pharmaceutical development of LXR agonists has been challenged by the fact that these compounds usually induce *de novo* lipogenesis (Grefhorst *et al*, 2002; Kirchgessner *et al*, 2016), which is mainly attributable to hepatocyte-specific LXR $\alpha$  activation (Bradley *et al*, 2007; Zhang *et al*, 2012). Importantly, DHCR24 inhibition by SH42 decreases plasma FFA and CE levels, without increasing plasma total TAG and DAG levels (Fig 3), which implies that LXR $\alpha$ activation by desmosterol, in contrast to synthetic agonists, does not lead to hyperlipidemia. This can be easily understood from the fact that desmosterol activates LXR $\alpha$  in macrophages/KCs only, whereas



Figure 6.

Figure 6. Treatment with SH42 reduces hepatic crown-like structures, liver collagen content, and plasma alanine transaminase levels in an established NAFLD mode.

A E3LCETP mice were fed with a HFCD for 10 weeks first and then treated with vehicle (Ctrl) or DHCR24 inhibitor SH42 (SH42) (n = 10 and 9 mice, respectively) for additional 8 weeks. Mice were killed and livers were collected.

- B Liver sections were stained with hematoxylin and eosin (HE) and Oil Red O.
- C, D Hepatic steatosis was scored using the HE-stained slides and (D) lipid-positive area was quantified using the Oil Red O stained slides.
- E-G Liver sections were stained with F4/80 (E) to quantify (F) hepatic crown-like structures as indicated by the arrows in Fig 6E and (G) F4/80 positive area.
- H, I To quantify the collagen content, liver sections were stained with (H) Sirius red/Fast green and (I) quantified accordingly.
- J Blood samples were collected after 18 weeks of treatment and plasma alanine transaminase (ATL) levels were measured.

Data information: Values are mean  $\pm$  SEM. Differences between groups (SH42/Ctrl) were determined using a Mann–Whitney test. \*P < 0.05, \*\*P < 0.01, vs. ctrl. Scale bar: 100 or 200  $\mu$ m as indicated.

Source data are available online for this figure.

LXR-induced hyperlipidemia is mainly due to increased *de novo* lipogenesis within hepatocytes (Muse *et al*, 2018). In fact, we have shown that desmosterol even suppresses SREBPs in macrophages via an Insig/SREBP cleavage-activating protein (SCAP)-dependent mechanism (Rodriguez-Acebes *et al*, 2009; Spann *et al*, 2012; Muse *et al*, 2018). Taken together, SH42-induced desmosterol does not evoke an undesirable increase in plasma and hepatic lipid levels that are typical for synthetic LXR agonists, likely by selective LXR $\alpha$  activation in macrophages/KC rather than hepatocytes.

A human study has demonstrated that desmosterol levels in circulation and liver are associated with NASH development (Simonen et al, 2013). However, whether desmosterol has a specific role in the pathophysiology of NAFLD/NASH is still unknown. To the best of our knowledge, our present study for the first time shows the potential application of pharmacological DHCR24 inhibition to enhance endogenous desmosterol levels for the treatment of NAFLD/NASH. Interestingly, a recent genetic study demonstrated that the rs588709 variant near the DHCR24 locus is associated with lower circulating TAG-rich VLDL particles (Sliz et al, 2020), which potentially extends the role of DHCR24 to human lipid metabolism. Triparanol is the most widely used (nonspecific) DHCR24 inhibitor (Muller et al, 2019); however, it was withdrawn from clinical application due to severe adverse side effects, such as nausea and vomiting, cataracts, and skin disorders (Kirby, 1967). In contrast to Triparanol with an IC50 value of 14 µM, SH42 is a much more potent DHCR24 inhibitor, having an IC<sub>50</sub> value of less than 10 nM in a cellular assay (Moebius et al, 1998; Muller et al, 2017) as judged by desmosterol accumulation and <sup>13</sup>C labelling of cholesterol. In our preclinical mouse model, we did not observe adverse effects of SH42 treatment on food intake, body weight, and body composition. Additionally, desmosterol-induced LXR activation via inhibition of DHCR24 did not cause a reduction of circulating neutrophils or neutropenia, another adverse effect of systemic LXR activation (Kirchgessner et al, 2016). Importantly, our study showed that DHCR24 inhibition by SH42 not only prevented but also affected NASH progression in mice with established hepatic steatosis, as evidenced by reducing the number of hepatic crown-like structures, collagen content, and plasma alanine transaminase levels. These advantages of targeting DHCR24 over synthetic LXR agonists will possibly lead to a promising application of DHCR24 inhibitors to combat NAFLD/NASH, in addition to other cardiometabolic and inflammatory diseases. Of note, our study is limited by our inability to provide direct evidence of SH42 binding to DHCR24, due to a lack of available pure enzyme and the fact that DHCR24 is

membrane-bound making its isolation and co-crystallization very difficult. Therefore, we cannot exclude that SH42 exerts indirect inhibitory actions responsible for desmosterol accumulation or interaction with yet undiscovered targets contributing to the improvement of the NAFLD phenotype in our study.

In conclusion, we show that pharmacological inhibition of DHCR24 increases desmosterol to prevent diet-induced hepatic steatosis and inflammation, two main hallmarks of NAFLD/NASH development, without inducing hyperlipidemia. As such, our study paves the way for developing a new therapeutic strategy for the treatment of NAFLD/NASH.

# Materials and Methods

#### Animals and treatments

Hemizygous APOE\*3-Leiden (E3L) mice were crossbred with homozygous human cholesteryl ester transfer protein (CETP) transgenic mice to generate heterozygous E3L.CETP mice on a C57BL/6J background (Westerterp et al, 2006). Because of the phenotypical heterogeneity of E3L.CETP mice, nonresponder mice were identified by 4h fasting plasma lipid levels, that is, total cholesterol levels < 2 mM and triglyceride levels < 2 mM, and excluded before high-fat highcholesterol diet (HFCD) treatment (Tarasco et al, 2018). LXRadeficient mice (also on C57BL/6J background), generated by Deltagen using gene-targeting methods as described (Plosch et al, 2006), were kindly provided by Tularik (San Francisco, CA, USA). Mice were group-housed in individually ventilated cages in standard conditions at room temperature (22°C) with 40  $\pm$  5% relative humidity and a 12-h light/dark (7 am lights on; 7 pm lights off) cycle. Water and standard laboratory diet (801203, Special Diets Services, UK) were available ad libitum, unless indicated otherwise. This study was approved by the Animal Ethical Committee of Leiden University Medical Center, Leiden, The Netherlands (AVD1160020173305, PE.18.034.007) and the Animal Ethical Committee of University Groningen, Groningen, The Netherlands (PE.18.034.030, 173305-02-001). All animals received humane care according to the criteria outlined in the NIH "Guide for the Care and Use of Laboratory Animals." All animal procedures were performed conform the guidelines from Directive 2010/63/EU of the European Parliament on the protection of animals used for scientific purposes.

At the age of 10-12 weeks, male *E3L.CETP* mice and LXR $\alpha$ -deficient mice were fed a HFCD (Altromin, Germany) containing

60% (energy) fat and 1% (wt/wt) cholesterol, and were randomized into two groups treated with either the DHCR24 inhibitor SH42 (0.5 mg·mouse<sup>-1</sup>; Muller *et al*, 2017; Korner *et al*, 2019) or vehicle (saline containing 3.3% ethanol and 3.3% Cremophor EL) 3 times per week by intraperitoneal injection. *E3L.CETP* mice were treated for 4 weeks (n = 6 mice per group) and 8 weeks (n = 8 mice per group) to evaluate effects on hepatic immune cells via flow cytometry analysis and hepatic steatosis via quantitative lipidomic analysis, respectively. LXR $\alpha$ -deficient mice were treated with either SH42 (n = 11 mice) or vehicle (n = 10 mice) for 4 weeks to evaluate effects on hepatic steatosis and immune cells. In a rescue experiment, male *E3L.CETP* mice were fed with HFCD for 10 weeks first and cotreated with SH42 (0.5 mg·mouse<sup>-1</sup>) or vehicle (saline containing 3.3% ethanol and 3.3% Cremophor EL) 3 times per week by intraperitoneal injection for additional 8 weeks.

Body weight was measured weekly. Body composition (i.e., fat body and lean body mass; EchoMRI-100; EchoMRI, Houston, TX, USA) was evaluated before and after intervention. Food intake was determined during the treatment period.

## Plasma glucose, insulin, and ALT assay

Four-hour-fasted (9 am till 1 pm) blood samples were collected via tail vein of *E3L.CETP* mice bleeding into heparin-coated capillaries after 8 weeks of treatment with vehicle or SH42. Blood samples were then centrifuged at 15,000 *g* for 5 min at 4°C to collect plasma. 2.5  $\mu$ l plasma was used to measure glucose (INstruchemie, The Netherlands) and 10  $\mu$ l to measure insulin (Mercodia AB, Sweden) according to the manufacturers' protocols. The homeostatic Model Assessment for Insulin Resistance (HOMA-IR) scores were calculated using the equation: HOMA-IR = fasting insulin (mU l<sup>-1</sup>) × fasting glucose (mmol l<sup>-1</sup>)/22.5 (Matthews *et al*, 1985).

To measure ALT levels, 4 h-fasted (9 am till 1 pm) blood samples were collected via tail vein of *E3L.CETP* mice after 16 weeks of HFCD treatment. Blood samples were then centrifuged at 15,000 g for 5 min at 4°C to collect plasma. Two microliter plasma was diluted 200 times and used for plasma ALT assay (ab282882, Abcam, UK) according to the manufacturer's protocol.

## Liver histology and hepatic steatosis scoring

After 8 or 18 weeks of treatment in E3L.CETP mice or 4 weeks in LXRa-deficient mice, mice were killed and transcardially perfused with ice-cold PBS, and a small piece of the right liver lobe was collected and fixed in phosphate-buffered 4% formaldehyde. Fixed tissues were embedded in paraffin and cut into sections of 5 µm thickness for staining with hematoxylin and eosin (HE) and/or the macrophage marker F4/80 (1  $\mu$ g ml<sup>-1</sup>, MCA497, Serotec, Oxford, UK), which was detected using ImmPRESS HRP goat anti-rat IgG detection kit (MP-744-15, Vector Laboratories, CA, USA). Hepatic steatosis was evaluated from HE-stained slides by two blinded experts, respectively, with the criteria proposed by Liang et al (2014). Hepatocellular vesicular steatosis, that is, microvesicular steatosis and macrovesicular steatosis, and hepatocellular hypertrophy were scored (grade 0-3) based on the percentage of the total area affected, and the grades were summed into one standard as "steatosis score" (0-9). Liver Oil Red O staining was performed on frozen liver tissue sections to visualize and quantify lipid droplets using ImageJ software (version 1.50i). F4/80 positive area using F4/80 stained slides were quantified using ImageJ software (version 1.50i). Hepatic crown-like structures formed by macrophages aggregating around dead hepatocytes were counted from F4/80 stained slides, which were blinded before analysis and expressed as number per area (Itoh *et al*, 2013). To quantify the collagen content, liver sections were stained with Sirius red/Fast green and quantified using ImageJ software (version 1.50i). Values in figures for each staining present means of nine randomly selected image fields (0.468 mm<sup>2</sup> per field) per mouse. The antibody used is listed in Appendix Table S1.

#### GC–MS analysis of desmosterol

After 8 weeks of treatment in E3L.CETP mice or 4 weeks in LXRadeficient mice, desmosterol levels were measured using GC-MS after alkaline hydrolysis, as described previously for plasma (Muller et al, 2017) and liver (Korner et al, 2019). In brief, a Bruker Scion GC-MS system (used for E3L.CETP mice) or an Agilent 7890B coupled to a 5977B MS (used for LXR KO mice) were utilized and operated in selected ion monitoring mode (SIM). An Agilent VF-5 ms column, 25 m  $\times$  0.25 mm  $\times$  0.25  $\mu m$  was used. Helium 99.9990% was used as carrier gas at a constant flow rate of 1.4 ml min<sup>-1</sup>. Injector and transfer line were operated at 280°C. The oven program started at 90°C held for 0.5 min, then ramped to 270 at 30°C min<sup>-1</sup>, and ramped to 310 at 10°C min<sup>-1</sup>. The following masses were used to quantify desmosterol levels against an external calibration line (0–10 ppm) m/z 445 + 355 for cholestanol (IS), m/z357 and 271 for cholestan (IS) and m/z 343 + 253 or 441.2 for desmosterol. Desmosterol was identified by matching characteristic ions and retention times with an authentic standard (Sigma Aldrich).

#### Quantitative lipidomic analysis

Quantitative lipidomic analysis was carried out in liver and plasma samples in E3L.CETP mice and in LXRa-deficient mice as described elsewhere (Korner et al, 2019). Briefly, a small piece of the right liver lobe was homogenized with 600 µl LCMS grade water in a bullet blender for 30 s and an aliquot corresponding to 5 mg liver tissue was used. For plasma samples, 25 µl was used. 25 µl internal standard mix (100 µl for liver samples; Lipidyzer<sup>™</sup> internal standard kit, containing > 50 labeled internal standards for 13 lipid classes, Sciex cat# 504156), 500 µl methyl tert-butyl ether (MTBE), and 160 µl methanol were subsequently added and the mixture was shaken for 30 min at room temperature. After adding 200 µl water, samples were centrifuged at 16,000 g for 3 min and the upper organic layer was collected in a glass vial. The remaining sample was extracted again by adding 300 µl MTBE, 100 µl methanol, and 100  $\mu$ l water for 30 min. The organic extracts were then pooled and dried under a gentle stream of nitrogen. The dry extract was subsequently dissolved in 250 µl Lipidyzer running buffer and analyzed according to the manufacturer's protocol. For data analysis, 13 lipid classes, including cholesteryl esters (CE), ceramides (CER), diacylglycerols (DAG), dihydroceramides (DCER), free fatty acids (FFA), hexosylceramides (HCER), lysophosphatidylcholines (LPC), lysophosphatidylethanolamines (LPE), phosphatidylcholines (PC), phosphatidylethanolamines (PE), sphingomyelins (SM) and triacylglycerols (TAG) were compared.

# Isolation of liver immune cells and peripheral blood mononuclear cells

After 4 weeks of SH42 treatment, blood and liver were collected and processed for flow cytometry as described previously (Hussaarts et al, 2015). In short, blood was collected from the retro-orbital sinus of anesthetized mice into heparin capillaries, after which mice were killed and transcardially perfused with PBS for 5 min to remove circulating cells. A piece of right liver lobe was cut off for other liver assays and the remainder of liver tissues were minced with a scalpel to form a paste, digested with collagenase type IV (Sigma Aldrich, USA), and hepatocytes were removed by low gravity centrifugation at 50 g for 3 min. CD45<sup>+</sup> leukocytes were isolated using CD45 microbeads (35 µl beads per liver; Miltenyi Biotec, USA) and an LS column, according to the manufacturer's instructions. In addition, peripheral blood mononuclear cells were isolated. To this end, 450 µl blood was mixed with red blood cell lysis/fix buffer (Becton Dickinson, USA) (3:20; v/v) for 15 min at room temperature. Subsequently, white blood cells were pelleted by centrifugation at 600 g for 5 min. The supernatant was discarded and cells were resuspended and washed with PBS. The cells isolated from the liver and blood were counted and characterized by flow cytometry.

## Flow cytometry analysis

All cell suspensions were stained with the fixable viability dye Zombie-UV (BioLegend, USA) and fixed with 1.9% paraformaldehyde. Within 24 h of fixation, the cells were preincubated with an Fc $\gamma$  blocking antibody (Thermo Fisher Scientific) for 15 min. The cells were then stained with fluorescently labeled antibodies listed in the Appendix Table S2 for 30 min at 4°C in the dark. Fluorescently labeled cells were measured on an LSRII (Becton Dickinson) and gates were set according to Fluorescence Minus One (FMO) controls using FlowJo<sup>TM</sup> Software (Becton Dickinson). Representative gating schemes are shown in Appendix Figs S3 and S4.

#### Statistical analysis

Group size was determined by power analysis of data from our previous studies to achieve a statistical power of 80% and *P*-value of 0.05. Mice were randomized into different groups before the treatments. Investigators were blinded for hepatic steatosis scoring and were not blinded for other assays. Outliers were identified using the Grubbs' test (https://www.graphpad.com/quickcalcs/Grubbs1.cfm) and removed from statistical analysis, which is clearly stated in figure legends if applicable. Because some datasets did not pass a statistical test for Gaussian distribution, differences between two groups were compared using a nonparametric Mann–Whitney test performed in GraphPad Prism 8.1 (GraphPad Software). For quantification of different lipid classes, Benjamini–Hochberg correction was used for multiple hypothesis testing (Figs 1G–J and 4E–H). P < 0.05 was considered significant (\*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001).

# Data availability

The authors declare that all data supporting the findings of this study are available within the paper and its Supplementary

### The paper explained

## Problem

Although more than 30% of the general population have nonalcoholic fatty liver disease (NAFLD), no medication is available yet for the treatment of NAFLD. As the liver X receptor (LXR) is a key regulator of metabolic and inflammatory signaling, LXR agonism has a theoretical potential for treating NAFLD, but synthetic LXR agonists induce hyperlipidemia in preclinical models and humans.

#### Results

In this study, we show the potential application of pharmacological  $\Delta$ 24-dehydrocholesterol reductase (DHCR24) inhibition by SH42 to prevent diet-induced hepatic steatosis and liver inflammation in a well-established translational model with human-like NAFLD characteristics. Inhibition of DHCR24 increased the endogenous LXR agonist desmosterol to exert metabolic and immune benefits in a strictly LXR $\alpha$ -dependent manner without causing hyperlipidemia.

#### Impact

Pharmacological DHCR24 inhibition by SH42 increases desmosterol to prevent diet-induced hepatic steatosis and inflammation, two main hallmarks of NAFLD/NASH development, without inducing hyperlipidemia. Pharmacological DHCR24 inhibition may represent a novel therapeutic strategy to activate LXR for the treatment of NAFLD and potentially other cardiometabolic diseases.

Information Appendix. This study includes no data deposited in external repositories.

Expanded View for this article is available online.

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## Author contributions

Enchen Zhou: Conceptualization; data curation; formal analysis; investigation; visualization; methodology; writing – original draft; project administration; writing – review and editing. Xiaoke Ge: Formal analysis; investigation; methodology; writing – review and editing. Hiroyuki Nakashima: Conceptualization; data curation; formal analysis; investigation; visualization; methodology; writing – review and editing. Rumei Li: Formal analysis; investigation; methodology; writing – review and editing. Hendrik J P van der Zande: Formal analysis; investigation; methodology; writing – review and editing. Cong Liu: Investigation; writing – review and editing. Zhuang Li: Investigation; writing – review and editing. Christoph Müller: Resources; writing – review and editing. Franz Bracher: Resources; writing – review and editing. Yassene Mohammed: Data curation; formal analysis; visualization; writing – review and editing. Jan Freark de Boer: Investigation; project administration; writing – review and editing. Folkert Kuipers: Resources; project administration; writing – review and editing. Bruno Guigas: Resources; project administration; writing – review and editing. Christopher K Glass: Resources; supervision; project administration; writing – review and editing. Patrick C N Rensen: Conceptualization; resources; data curation; supervision; funding acquisition; project administration; writing – review and editing. Martin Giera: Conceptualization; resources; data curation; supervision; funding acquisition; project administration; writing – review and editing. Martin Giera: Conceptualization; resources; data curation; supervision; funding acquisition; project administration; writing – review and editing. Yanan Wang: Conceptualization; resources; data curation; supervision; funding acquisition; project administration; writing – review and editing. Yanan Wang: Conceptualization; resources; data curation;

# Disclosure and competing interests statement

The authors declare that they have no conflict of interest.

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