Loss of hepatic \textit{Flcn} protects against fibrosis and inflammation by activating autophagy pathways

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Non-alcoholic fatty liver disease (NAFLD) is the most frequent liver disease worldwide and can progress to non-alcoholic steatohepatitis (NASH), which is characterized by triglyceride accumulation, inflammation, and fibrosis. No pharmacological agents are currently approved to treat these conditions, but it is clear now that modulation of lipid synthesis and autophagy are key biological mechanisms that could help reduce or prevent these liver diseases. The folliculin (FLCN) protein has been recently identified as a central regulatory node governing whole body energy homeostasis, and we hypothesized that FLCN regulates highly metabolic tissues like the liver. We thus generated a liver specific \textit{Flcn} knockout mouse model to study its role in liver disease progression. Using the methionine- and choline-deficient diet to mimic liver fibrosis, we demonstrate that loss of \textit{Flcn} reduced triglyceride accumulation, fibrosis, and inflammation in mice. In this aggressive liver disease setting, loss of \textit{Flcn} led to activation of transcription factors TFEB and TFE3 to promote autophagy, promoting the degradation of intracellular lipid stores, ultimately resulting in reduced hepatocellular damage and inflammation. Hence, the activity of FLCN could be a promising target for small molecule drugs to treat liver fibrosis by specifically activating autophagy. Collectively, these results show an unexpected role for \textit{Flcn} in fatty liver disease progression and highlight new potential treatment strategies.

\textbf{Abbreviations}

FLCN  Folliculin  
HFD  High fat diet  
H&E  Hematoxylin and eosin  
MCD  Methionine/Choline Deficient diet  
mTORC1  Mammalian Target of Rapamycin Complex 1  
NAFLD  Non-alcoholic fatty liver disease  
NASH  Non-alcoholic steatohepatitis  
TFEB  Transcription Factor EB  
TFE3  Transcription Factor Binding To IGHM Enhancer 3

Non-alcoholic fatty liver disease (NAFLD) is a disease spectrum characterized by triglyceride accumulation in hepatocytes and is associated with obesity, insulin resistance, type 2 diabetes, and dyslipidemia\textsuperscript{1}. Initially exhibiting NAFLD, approximately one-third of patients will progress toward non-alcoholic steatohepatitis (NASH)\textsuperscript{2}. Furthermore, approximately one-third of patients with NASH will progress to cirrhosis and final liver dysfunction\textsuperscript{3,4}. There is currently no pharmacological agent approved to treat NASH or to slow its progression\textsuperscript{5,6}. The only proven treatments are weight loss and increased physical activity, with fibrosis improvement in only 10–20\% of patients\textsuperscript{5,6}.

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NALDH is characterized by hepato-cellular injury, immune cell-mediated inflammation, and progressive liver fibrosis. The accumulation of lipid intermediates within hepatocytes causes lipotoxicity, cellular stress, and eventually cell death via distinct mechanisms. Fatty acid modulates cell signaling, disrupts metabolic homeostasis, and induces ER stress. Chronic liver injury eventually leads to fibrosis as a consequence of chronic wound-healing responses initiated by dying hepatocytes. In a healthy state, wound-healing restores liver function and structure by recruiting liver progenitor populations and inflammatory cells, which can release angiogenic and matrix remodeling factors. In NASH, wound-healing is perpetual, and the persistent inflammation prevents the termination of the process thus perpetuating fibrogenesis and inflammation. Therefore, regulation of the lipid content is crucial to maintain liver homeostasis.

AMP-activated protein kinase (AMPK) is a master regulator of cellular energy metabolism and key transducer of metabolic inputs that can stimulate the switch from lipogenesis, an anabolic ATP-consuming process, to the catabolism of fatty acids and generation of energy. In the recent years, a causal role of the mechanistic target of rapamycin (mTOR) signaling pathway has also been linked to hepatic steatosis. mTOR is a serine/threonine kinase found in two distinct complexes, mTORC1 and mTORC2, that controls various biosynthetic pathways. These include autophagy and lipid metabolism, two biological pathways that are known to be tightly linked to the transition from a healthy liver to NAFLD and NASH. Consequently, there was hope that inhibition of mTOR with rapalogs such as rapamycin could reduce steatosis, notably by restoring autophagy. However, more recent studies indicate that long term treatment with these inhibitors induce liver damage and enhance tumorigenesis.

In that settings, we searched for a new therapeutic target linked to the AMPK and mTOR signaling pathways but for which its modulation would not cause long-term side effects such as rapalogs.

Folluculin (Flen) loss of function has been studied in nematodes and various tissues of knockout mice, showing that Flen loss leads to a chronic activation of AMPK with surprisingly beneficial effects. For example, mice with Flen knockout specifically in skeletal muscle developed a pronounced metabolic shift towards oxidative phosphorylation, increased mitochondrial biogenesis, and red colored muscles. Similarly, mice with conditional Flen knockout in adipocytes developed resistance to high-fat diet (HFD)-induced obesity as well as browning of white adipocytes and increased mitochondrial activity in brown fat. In addition, FLCN is also a crucial mTORC1 regulator. FLCN is a GTPase activating protein (GAP) toward the Rags GTPase C and D, two important proteins required for mTORC1 localization on the lysosomes. Therefore, Flen loss prevents the activation of mTORC1 on the lysosomal membrane and activates autophagy, without affecting the mRNA translation machinery. Still, little is known about how FLCN regulates a highly metabolic tissue such as the liver and, more specifically, the potential role of FLCN in liver fibrosis progression has never been investigated.

In this study, we generated a liver specific Flen knockout mouse model challenged with a methionine/choline deficient, high fat, high sucrose (MCD) diet, mimicking transition to liver fibrosis in vivo. We demonstrate that liver-Flen KO mice were protected from MCD-induced triglyceride accumulation and liver damage. Importantly, loss of Flen in liver tissue resulted in increased autophagic activity and nuclear localization of TFE3 and TFEB, reduced fibrosis, and reduced inflammation, underlying potential mechanisms for liver fibrosis protection. Altogether, these results identified FLCN as a novel potential therapeutic target for NAFLD and NASH.

**Results**

**Hepatic loss of Flen protects against high fat diet stimulated NAFLD.** To study the role of FLCN in the liver, we generated a liver-specific Flen KO (liver-Flcn−/−) mouse model by crossing Flenfl/flox mice with albumin-cre+− mice (Fig. 1A and supplemental Fig. 1A,B). Liver-Flcn−/− mice were born at the expected Mendelian frequency, displayed no developmental defects, survived without difficulty, were fertile, and had similar weights compared with wild-type mice at weaning (4 weeks of age).

We initially established whether Flcn loss in the liver affects whole-body metabolism by feeding them either normal chow or a high fat diet (HFD) containing 60% kcal from fat for 6 weeks. Control mice (Cre mice) gained significantly more weight than liver-KO mice when challenged with HFD compared to Cre mice (Fig. 1C). To understand how liver-specific loss of Flcn protected against HFD-induced obesity, we then examined if it improved systemic energy metabolism. First, glucose and insulin tolerance tests showed that loss of Flcn in hepatocytes significantly increased glucose uptake and enhanced insulin responsiveness, respectively, under both chow and a HFD regimen (Fig. 1D and supplemental Fig. 1C). Secondly, we employed the use of metabolic cages capable of simultaneously measuring animal food intake, physical activity and energy expenditure. Mice fed either chow or HFD were examined individually in metabolic cages. Liver-Flcn−/− mice were found to be more active during the dark phase of the circadian cycle concomitant with increased maximal oxygen consumption (VO2) compared to Cre controls independent of diet (Fig. 1E and supplemental Fig. 1D). Together, the data suggest that the increased energy expenditure and metabolic activity of liver-Flcn−/− mice are underlying factors in the resistance of these animals to HFD-induced obesity. Hence, loss of Flcn specifically in the liver improved metabolic homeostasis and prevented weight gain in a diet-induced obesity model.

**Flcn loss protects against MCD diet-induced fibrosis and inflammation.** Since liver phenotypes associated with an HFD are highly associated with obesity and whole-body insulin resistance, the effects observed in this experimental setting may have been an indirect consequence of reduced adiposity. To test whether protection against steatosis and NAFLD was due to cell-autonomous effects of Flcn KO in hepatocytes, we investigated whether loss of hepatic Flcn was also protective in a model of liver fibrosis that does not cause weight gain or insulin resistance. To this end, we subjected Cre and liver-Flcn−/− mice to an MCD diet for six weeks. The MCD diet is a classical dietary model of liver fibrosis. Although the diet comprises high sucrose (46%) and fat (10%),
it lacks methionine and choline, which are indispensable for hepatic mitochondrial β-oxidation and very low-density lipoprotein (VLDL) synthesis. This diet is a model of a metabolic challenge and results in a significant rapid onset of a NASH-like phenotype with fibrosis, inflammation, oxidative stress, and liver cell death. As seen for mice under HFD (Fig. 1), hepatic loss of Flcn protected against hepatic lipid accumulation in mice under the MCD diet, as shown with the H&E staining (Fig. 2A). Indeed, specific quantification of the total triglycerides in the liver tissue also showed a significant reduction in fat accumulation in liver-Flcn−/− mice (Fig. 2C and Supplemental Fig. 2B–D). These results suggest that liver-specific loss of Flcn reduces fat accumulation not through increased fat export into the blood, but rather by protecting against liver damage (Fig. 2D).

One hallmark of NASH is fibrosis, which generates liver damage and eventually leads to hepatocyte death. In our mouse model, relative mRNA transcript levels of genes related to fibrosis were induced in Cre mice fed the MCD diet, but generally to a lower extent in liver-Flcn−/− mice (Fig. 3A). Sirius Red staining also revealed that fibrosis was aggravated in Cre mice challenged on the MCD diet, but this effect was diminished in liver-Flcn−/− mice (Fig. 3B, quantified in C). Immunohistochemistry staining of α-smooth muscle actin (α-SMA), a fibrosis-related marker, was significantly increased in Cre mice challenged with the MCD diet and again loss of Flcn was protective (Fig. 3D, quantified in E). Finally, immunohistochemistry staining of 4-Hydroxynonenal (4-HNE), a lipid peroxidation marker, was significantly increased in Cre mice challenged with the MCD diet but diminished in liver-Flcn−/− mice (Fig. 3E, quantified in G). Taken together, these results demonstrate that liver-specific loss of Flcn protects against fibrosis.

NASH development is conjointly characterized by increased inflammation that contributes to liver damage and disease progression. In our mouse model, relative mRNA transcript levels of genes involved in inflammation...
were lower in liver-Flcn−/− mice compared to Cre mice when fed the MCD diet (Fig. 3H). Using a mouse protein cytokine array, we determined the cytokine and chemokine secretion profiles in mouse livers. Cytokine protein levels were higher in Cre mice challenged with the MCD diet compared to liver-Flcn−/− mice (Fig. 3I,J). More specifically, some macrophage chemo-attractants (RANTES, MCP-1, IP-10) were induced in Cre mice fed the MCD diet but to a lesser extent in liver-Flcn−/− mice (Fig. 3J). Normal livers are infiltrated by specialized macrophages that contribute to inflammation when activated. Notably, immunohistochemistry staining of F4/80, a marker of murine macrophages, was increased in Cre mice fed the MCD diet, but were not increased in liver-Flcn−/− mice (Fig. 3K, quantified in L). Collectively, our data indicates that liver-specific loss of Flcn reduces fibrosis and inflammation in mice challenged with the MCD diet.

**Flcn loss activates autophagy in a mouse model of liver fibrosis.** This protective impact on inflammation and fibrosis following loss of Flcn prompted us to investigate the molecular signature of liver-Flcn−/− mice responsible for the protective phenotype using high-throughput whole-transcriptome sequencing (mRNA-seq). Unsupervised hierarchical clustering highlighted three distinct patterns of gene expression changes (Fig. 4A, Supplemental Table 1), clusters 1 and 3 not being affected by the Flcn status. Interestingly, cluster 2 highlighted genes whose expression was induced in Cre mice receiving the MCD diet but that were not increased in liver-Flcn−/− mice (Fig. 4A,B). Gene enrichment analysis using GO Enrichr revealed the major pathways differentially affected in this cluster by the diet and the genotypes (Fig. 4C). The most significantly affected biological pathway was extracellular matrix organization, which represents genes involved in fibrosis development typically observed in NASH, such as Acta2, Mmp2, Serpinel1, and Timp1 (as validated by qRT-PCR and consistent with histology experiments in Fig. 3). Other important pathways affected include mitotic sister chromatid segregatio-
Figure 3. Flcn loss reduces MCD diet-induced fibrosis and inflammation. (A) Relative quantitative real-time PCR analysis of fibrosis-related genes mRNA transcript levels in livers of mice fed either on chow or methionine/choline deficient diet (MCD) over a period of 6 weeks (mean ± SEM of the RNA fold change of indicated mRNAs; two-way ANOVA, *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001; n = 8 mice per condition). (B) Sirius red staining in liver sections of mice fed as described in (A). Images are representative of 6 mice per condition. (C) Quantification of the sirius staining in liver sections of mice fed as described in (A) (mean ± SEM, two-way ANOVA, **P < 0.01; ****P < 0.0001; n = 6 mice per condition). (D) α-SMA Immunohistochemistry (IHC) staining in liver sections of mice fed as described in (A). Data are representative of 4 mice per condition. (E) Quantification of α-SMA IHC staining in liver sections of mice fed as described in (A) (mean ± SEM, two-way ANOVA, *P < 0.01; ***P < 0.001; n = 4 mice per condition). (F) 4-HNE Immunohistochemistry (IHC) staining in liver sections of mice fed as described in (A). Data are representative of 4 mice per condition. (G) Quantification of 4-HNE IHC staining in liver sections of mice fed as described in (A) (mean ± SEM, two-way ANOVA, **P < 0.01; ***P < 0.001, ****p < 0.0001; n = 4 mice per condition). (H) Relative quantitative real-time PCR analysis of inflammation-related genes mRNA transcript levels in livers of mice fed as described in (A) (mean ± SEM of the RNA fold change of indicated mRNAs; two-way ANOVA, *P < 0.05, **P < 0.01, ****P < 0.0001; n = 8 mice per condition). (I-J) Mouse cytokines and chemokines quantification in liver homogenates of mice fed as described in (A) (mean ± SEM, two-way ANOVA, *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001; n = 3 mice per condition). (K) F4/80 Immunohistochemistry (IHC) staining in liver sections of mice fed as described in (A). Data are representative of 4 mice per condition. (L) Quantification of F4/80 IHC staining in liver sections of mice fed as described in (A) (mean ± SEM, two-way ANOVA, *P < 0.05; **P < 0.01; n = 4 mice per condition).
tion, sister chromatic segregation, and mitotic nuclear division. These genes are generally induced in regenerating livers following injury. Moreover, pathways involved in inflammation such as neutrophil activation and cellular response to cytokine stimulus were differentially expressed. Overall, genes involved in fibrosis, inflammation, and mitotic cell cycle were induced in mice fed the MCD diet but to a lesser extent in Flcn KO mice (Fig. 4D–F). Concurrently, no significant changes in expression between Cre and Flcn KO mice were detected for pathways involved in lipolysis (GO:0016042) and lipogenesis (GO:0006954) (Supplemental Fig. 3A–C).

Emerging data supports a role of autophagy in the liver to regulate intracellular lipid stores and energy homeostasis. Autophagy is an important process in which the cells degrade its own content by sequestering cargoes inside the lysosomes. Degradation of these cargoes inside the lysosome is a multistep process that can

![RNA-seq analysis](https://example.com/rna-seq.png)

**Figure 4.** RNA-seq reveals major pathways affected by Flcn loss. (A,B) Unsupervised hierarchical clustering following RNA-seq analysis in livers of mice fed either chow or methionine/choline deficient diet (MCD) over a period of 6 weeks. (C) Gene enrichment analysis using GO Enrichr. (D) Unsupervised hierarchical clustering following RNA-seq analysis of fibrosis-related genes (GO:0030198). (E) Unsupervised hierarchical clustering following RNA-seq analysis of inflammation-related genes (GO:0006954). (F) Unsupervised hierarchical clustering following RNA-seq analysis of mitotic cell cycle-related genes (GO:0000278).

| Rank | Term                                           | Gene Count | P-Value  | FDR   |
|------|-----------------------------------------------|------------|----------|-------|
| 1    | extracellular matrix organization              | 43         | 8.8e-17  | 4.5e-13 |
| 2    | mitotic sister chromatid segregation           | 21         | 1.5e-11  | 3.8e-08 |
| 3    | sister chromatid segregation                   | 12         | 2.8e-09  | 4.8e-06 |
| 4    | mitotic nuclear division                       | 16         | 4.4e-08  | 8.9e-05 |
| 5    | neutrophil activation involved in immune response | 48  | 7.7e-08  | 7.8e-05 |
| 6    | cellular response to cytokine stimulus         | 45         | 9.7e-08  | 8.2e-05 |
be regulated by master transcription factors such as Transcription Factor EB (TFEB) and Transcription Factor Binding To IGHM Enhancer 3 (TFE3)\(^\text{15,30}\). The best described regulator of TFEB and TFE3 is the mTOR Complex 1 (mTORC1). Under non-stress conditions, mTORC1 phosphorylates TFE3 and TFEB, resulting in cytoplasmic sequestration and inhibition of their activity\(^\text{15}\). Conversely, nutrient starvation induced inhibition of mTORC1 removes the repressive phosphorylation on TFEB and TFE3, resulting in their nuclear translocation and activation of a panel of genes involved in autophagy\(^\text{15}\). We and others have shown that FLCN is a negative regulator of TFEB and TFE3 and positive regulator of mTORC1\(^\text{23,31-33}\). Moreover, FLCN exhibits GTPase-activating protein (GAP) activity upon nutrient replenishment targeting the Ras-related GTase C and Ras-related GTase D (RagC/D) resulting in mTORC1 activation, inhibition of TFEB and TFE3 nuclear translocation, and impaired autophagy and lysosomal function\(^\text{24,25}\).

Unsupervised hierarchical clustering of autophagy-related genes from our RNA-seq data revealed an important effect of the diet (Autophagy clusters A1 and A3, Fig. 5A), but also highlighted a subset of genes induced in liver-Flcn\(^{-/-}\) mice (Autophagy cluster A2, Fig. 5A-B). Relative mRNA transcript levels of autophagy-related genes mRNA, such as Ctsa, Depp1, and Ctsf, were significantly increased in liver-Flcn\(^{-/-}\) mice compared to Cre mice, while the diet had no effect (Fig. 5C). These genes are generally involved in lysosome/autophagosome maturation and activation, which can reduce hepatocellular injury and inflammation by clearing damaged or misfolded proteins and suppressing transcription or maturation of pro-inflammatory cytokines\(^\text{34}\). Immunoblot analysis additionally revealed higher protein levels of p62 in Cre mice challenged with the MCD diet, suggesting impaired autophagy, while Fln KO reduced protein levels (Fig. 5D, quantified in E). p62 mRNA transcript levels were induced in liver-Flcn KO but were unaffected by the diet (Fig. 5F). Therefore, the change in p62 protein levels is unlikely to be due to regulation at the mRNA level of p62. To further assess autophagy induction in a Flcn KO model, we measured autophagy flux in mouse embryonic fibroblasts (MEFs) in the absence and presence of BafA1 and Chloroquine, two compounds known to pharmacologically block autophagosome-lysosome fusion. Flcn loss in this cell line increased autophagy flux upon autophagy blockade, as measured by higher LC3-II accumulation in Flcn KO after treatment with BafA1 and Chloroquine (Fig. 5G, quantified in H). We also measured LC3 levels in liver protein lysates from mice fed either the chow or the MCD diet. We show that Flcn KO mice had higher LC3-I and LC3-II levels in both diets, suggesting functional and active autophagy (Supplemental Fig. 4A, quantified in B and C). Furthermore, the major transcription factors regulating autophagy and lysosome biogenesis, TFE3 and TFEB, were localized in the nuclei of hepatocytes, which correlated with increased autophagy in Flcn KO mice (Fig. 5I-L). Taken together, our data suggest that targeting Flcn improves resistance to fibrosis and inflammation, possibly by inducing autophagy when challenged with a liver fibrosis-inducing diet.

**Discussion**

In this study, we have engineered a mouse model to specifically ablate Flcn in the liver to elucidate the potential key role of this factor in the control of hepatic metabolism. Interestingly, liver specific loss of Flcn markedly enhanced whole-body energy metabolism at the baseline level as shown by the increased energy expenditure as well as improved glucose and insulin tolerance in these mice under HFD. Remarkably, deletion of Flcn in mouse liver tissue reduced fibrosis and inflammation when fed the fibrosis inducing MCD diet. Overall, our results establish a critical role of Flcn in liver-fibrosis progression.

Our work has highlighted for the first time a protective role for Flcn loss in the liver. Similarly, Flcn deletion in other tissues such as adipocytes and muscles led to improved capabilities and whole-body metabolism homeostasis\(^\text{21,23}\). The protective phenotype observed in the liver is comparable to previous attempts to activate autophagy and inhibit mTORC1 with rapalogs and rapamycin\(^\text{21,44}\), but was not accompanied by the deleterious long-term effects of these compounds on liver damage and tumorigenesis\(^\text{37,38}\). Indeed, we did not observe any harmful effect of Flcn deletion in livers. The difference is possibly the result of the more specific regulation of mTORC1 by FLCN on the lysosomal surface that is not affecting the mRNA translation machinery\(^\text{23,46}\). On the other hand, rapalogs regulate all mTOR complexes in the cell, reinforcing the potential therapeutic benefit of targeting more precisely FLCN.

Under the fibrosis-inducing diet, which mimics fibrosis as observed with NASH, RNA-seq data revealed that major pathways affected by Flcn loss specifically in the liver include inflammation, fibrosis, and mitotic cell cycle gene expression. The reduction in cell cycle gene expression is possibly the consequence of reduced liver damage and therefore of the reduced necessity to regenerate liver cells. We have previously shown that loss of Flcn in C. elegans and mammalian cells increased innate immune response in a process dependent on TFEB and TFE3\(^\text{35}\). In this study, we observed a decrease in inflammation gene expression in liver-Flcn KO mice fed the MCD diet. It is possible that FLCN modulates the immune microenvironment and polarizes the macrophages toward an anti-inflammatory state. Indeed, it was reported that FLCN loss in hematopoietic stem cells induced CD11c + CD206 + activated phagocytic macrophages, corresponding to the anti-inflammatory M2 macrophage subtype\(^\text{35}\). However, another study found that bone marrow-derived macrophages from Flcn myeloid KO mice rather promotes spontaneous M1-type polarization and enhanced baseline activation status\(^\text{35}\). Also, RAW 264.7 macrophages targeted with shRNA for Flcn had increased phagocytic activity toward dying/dead cells\(^\text{96}\). The previous effects of Flcn loss on macrophage activation were all dependent on TFEB/TFE3 activation status, highlighting the important role of autophagy in FLCN’s mechanism of regulation. Therefore, the importance and significance of the inflammation observed in liver-Flcn KO remains to be investigated.

FLCN seemed to be involved in regulating autophagy in a severe liver disease setting and it is possibly the primary mechanism of action by which loss of Flcn reduces liver fibrosis. Several compounds known to activate autophagy pathways such as acetylsphikinin and resveratrol also reduced MCD-induced fibrosis and inflammation\(^\text{37,38}\). Acetylsphikinin contributed to the removal of cellular lipid droplets through lipophagy in a process dependent on the AMPK/mTOR pathways\(^\text{37}\). Indeed, acute inhibition of mTOR and activation of autophagy...
Figure 5. Flcn loss activates autophagy in a mouse model of liver fibrosis. (A–B) Unsupervised hierarchical clustering following RNA-seq analysis in livers of mice fed either on chow or methionine/choline deficient diet (MCD) over a period of 6 weeks. (C) Relative quantitative real-time PCR analysis of autophagy-related genes mRNA transcript levels in livers of mice fed as described in (A) (mean ± SEM of the RNA fold change of indicated mRNAs; two-way ANOVA, *P < 0.05, **P < 0.01, ***P < 0.001; n = 8 mice per condition). (D) Immunoblot of liver protein lysates extracted from mice fed as described in (A). Data are representative of 6 mice per condition. Full-length blots are presented in Supplementary Fig. 5. (E) Quantification of the relative amount of p62 immunoblot. Data normalized to tubulin levels (mean ± SEM of the relative fold change; two-way ANOVA, *P < 0.05, **P < 0.01, ***P < 0.001; n = 6 mice per condition). (F) Relative RNAseq analysis of p62 mRNA transcript levels in livers of mice fed as described in (A) (mean ± SEM of p62 RNA fold change; one-way ANOVA, ***P < 0.001; n = 5 mice per condition). (G) Immunoblot of protein lysates of WT and Flcn KO MEFs incubated in presence of DMSO, BafA1 (100 nM), or Chloroquine (CQ; 100 μM) for 6 h. Data are representative of three independent experiments. Full-length blots are presented in Supplementary Fig. 5. (H) Quantification of the relative amount of LC3-II/LC3-1 immunoblot of MEFs treated as described in (G) (mean ± SEM of three independent experiment, one-way ANOVA, *P < 0.05). (I) TFE3 Immunohistochemistry (IHC) staining in liver sections of mice fed as described in (A) Data are representative of 6 mice per condition. (J) Quantification of TFE3 IHC staining in liver sections of mice fed as described in (A) (mean ± SEM, two-way ANOVA, ***P < 0.001; n = 4 mice per condition). (K) TFE3 Immunohistochemistry (IHC) staining in liver sections of mice fed as described in (A) Data are representative of 4 mice per condition. (L) Quantification of TFE3 IHC staining in liver sections of mice fed as described in (A) (mean ± SEM, two-way ANOVA, ***P < 0.001; n = 4 mice per condition).

In conclusion, we revealed that loss of Flcn in the liver is involved in autophagy, resulting in reduction of inflammation, fibrosis, and enhanced whole-body energy metabolism. These results show an unexpected role for Flcn in NAFLD and NASH progression. In light of the recently published cryo-EM structures of FLCN41,42, it is envisioned that pharmacological agents targeting the pocket of the FLCN GAP enzyme could be developed to achieve the loss of function phenotype in the liver shown in this study. Such molecules would represent new possibilities for treatment of obesity and NASH through the role of FLCN in hepatocyte homeostasis.

Material and methods

Antibodies. The FLCN rabbit polyclonal antibody was generated by the McGill Animal Resource Centre using recombinant GST-FLCN. TFE3 (Bethyl Laboratories, cat# A303-673A), TFE3 (Cell Signaling Technology, cat# 14779S), a-SMA (Abcam, cat# 5694), F4/80 (Cell Signaling Technology, cat# 70,076), 4-HNE (Millipore, cat# 393,207), p62 (Abcam, cat# ab56416), LC3B (Abcam, cat# ab48934), and Tubulin (Sigma-Aldrich, Cat# T9026) antibodies are commercially available.

Animals. All procedures for generating the Flcn knockout mouse model were performed at McGill University. Maintenance and experimental manipulation of mice were performed according to the guidelines and regulations of the McGill University Research and Ethic Animal Committee and the Canadian Council on Animal Care. The study was carried out in compliance with the ARRIVE guidelines and all experimental protocols were approved by the McGill University Research and Ethic Animal Committee. All studies were carried out using C57BL/6 mice housed on a 12 h light:12 h dark cycle at 22 °C. Flcn+/− (BDH6) mice were generated as previously described22. To generate liver-specific Flcn knockout mice, Flcn+/− mice were crossed with Albumin-Cre transgenic mice (kind gift from Dr. André Marette, Laval University, Quebec, Canada). Liver-Flcn+/− knockout mice were generated by crossing Flcn+/− mice homozygous for the floxed allele with Flcn+/−, Albumin-Cre+/− mice. Mice were fed on a low-fat diet containing 10 kcal percent fat (Research Diet, cat# D12450B) or a low fat diet containing 60 kcal percent fat (Research Diet, cat# D12492), or a Methionine/Choline Deficient (MCD) diet (Envigo, cat# TD.90262) beginning at 6 weeks of age and fed ad libidum. Controls used were Flcn+/+/-/+/−, Albumin-Cre+/−/+/− mice raised in identical conditions. Body weights were measured weekly.

Cell lines culture and treatments. Primary MEFs were isolated from C57BL/6 E12.5 Flcn floxed mice (generously provided by Dr. L.S. Schmidt, NCI, Bethesda, MD, USA) and cultured as described previously19. FLCN KO MEFs were generated after immortalization of primary Flx/Flox MEFs with SV40 large T and retroviral infection with a Cre recombinase, followed by puromycin selection. Cell lines were maintained in Dulbecco’s Modified Eagle’s Medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 100 U/m cillin, 100 μg/mL streptomycin, and 50 μg/mL gentamycin (Wisent) in 5% CO2 at 37 °C. For drug treatment experiments, cells were incubated for 6 h in medium containing one of the following reagents: DMSO (0.4%, Bioshop Canada), Chloroquine (100 μM, Sigma-Aldrich), and BafA1 (100 nM, Sigma-Aldrich).

Protein extraction and immunoblotting. Liver tissues were quickly snap frozen using a BioSpec BioSqueezer (Fisher Scientific, cat# NC1033496) cooled in liquid nitrogen following sacrifice. Approximately 100 mg of tissue were solubilized in RIPA buffer using a micropestle and sonicated 10 s. The mixture was then centrifuged at 16,000 × g for 15 min at 4 °C and the supernatant collected. Proteins were separated on SDS-PAGE gels and revealed by western blot as previously described22 using the antibodies listed above and IRDye.
800CW Goat anti-Rabbit IgG Secondary Antibody / IRDye 680RD Goat anti-Mouse IgG Secondary Antibody (LI-COR Biosciences). Membranes were scanned using the Odyssey imaging system (LI-COR Biosciences) and quantification performed using ImageStudio software (version 5.2.5, https://www.licor.com/bio/image-studio-lite/).

**Histology.** All histological procedures were performed using standard procedures at the histology core facility of the Goodman Cancer Research Centre (McGill University). Liver tissues were fixed in 4% formaldehyde for 2 days at room temperature immediately after sacrifice, embedded in paraffin, and cut to 4-μm sections on slides using a Leica Microtome (Leica Biosystems). The slides were stained with hematoxylin and eosin (Leica Biosystems, cat# 3802098), or Sirius Red (Abcam, cat# ab150681) using a Leica ST5020 Multistainer (Leica Biosystems) according to the standard protocol. Stained images were acquired using an Aperio ScanScope XT digital scanner (Leica Biosystems) and analyzed with ImageScope software (version 12.3.3.5048, https://www.leicabiosystems.com/fr/imagerie-pathologique/integre/aperio-imagescope/).

**Metabolic cages.** Mice were housed individually in metabolic cages for a total 3 days on a 12 h light: 12 h dark cycle with free access to water and food using a comprehensive laboratory animal monitoring system (CLAMS, Columbus Instruments). Food intake, energy expenditure, physical activity, maximal carbon dio-

| Gene     | Forward primers | Reverse primers          |
|----------|-----------------|--------------------------|
| ACTA2    | GACTCACAACGTCGCCCTATC | GCAGTAGCTCAGGAGGAATTAG |
| AKR1A1   | CTCGAAGATTGACGACCTATTTA | ACCCTTCAGGCCATCTT |
| ALDOA    | TGCTGTTTGGGCTGATCA | TGCGCTCCATCTTCTT |
| APOA4    | GACCCACGTCAAGGAGAAATG | CACTCTCGTACCTTATAATC |
| APOE     | GGAAAGCCACCGACCATTAAT | CTGTATCTGCTCAGGT |
| ASAH1    | ACCGGCAAGAAGTCTATAA | GGCTGGGCAATCTGGAGG |
| ATP5D    | GACCGAGAGTGGCCTTGAG | CTTCGCTGGAACCAATCC |
| B2M      | CTGGACGGCGGTGATGCTAT | CGGTCTGTCGATGATG |
| BAX      | GAGATGACTAGGGAGAACAATA | AGTATAAGAGGGCAACCA |
| CCL1     | GGATGGGTCAGAGCACATACTA | TACGGGTCAACCTTCACAT |
| CCL5     | CCGAGAAGAAGTGGTTGAGAAGG | AGCAATGACAGGGAAGCTATAC |
| CD36     | GGATGTTTCTAAGCCTTTTC | TGCCGCGGTCTCATAAT |
| CTS6     | GAGCGAAGACGACATCTT | TGCCCAATACCTGAGACT |
| CXYL1    | CAAACCAGGTGAAGAAAGAAGA | CATGTCCACCTCTTAAAA |
| FAS      | CTCCAGCTGTTGAAACATAC | CTGTCGCTCCTAGGT |
| FGF21    | CCAGAAGCAAGAGGAGAACT | AGCGCTGAGTACAGGAAG |
| G6PC     | CAGCTTCTGCGTCTAAATAAA | GCAAGATGAAAGTGGCAATAA |
| IL-1β    | GGCGAACCCTACATCATTT | GCGATGTTGACCAATTCCATC |
| LCN2     | GTGTCGCACTCTCCTTC | GAGATGTCGATCCAAATAA |
| LGALS3   | CTAACCCAGGCTGTACTTAA | ACAAGAGGATAAGGAGGAG |
| MCOLN1   | CCACCACGGAGAATGAGCATAC | GCTGTTATCTGATGTC |
| MMP2     | GAGGACTATAGGACCCGAAATAA | TGGTGAGCCTCTCATACT |
| NEU1     | CAGGAAGTTGTCGCCCTGAGA | GACAGAAAGCCACCTCTG |
| PEPCK    | GATGGTCGAAGAGGACCTTGTG | CGATTGCTGATGCTTAAAA |
| SDHA     | TTAAGCTTGCTTCCCCTCAT | AGATGTCGCCACACTCAA |
| SREBNE1  | AGAGAGAGAGAATTGAGAAAGA | GGAGAAATGTTGCTCTGTTAG |
| SOCS3    | GGGTGGCGAAAAGAAAAAGG | GTGAGGCTCAAGGCACCAGT |
| TBP      | ACCCTATGCTAGGCTTGG | GCCATAAGGCTATTGG |
| TGFBI    | TCACGCTCCAGAGAAGAA | GTGTCGCCGCTCAAATAAT |
| TIMP1    | AATCAGAGAACCAACCTTATAC | CATATCCAGAGGGCTTTC |
| TNFa     | GGGTGTTTCACTTCCATTCTC | TGGAACCTGAGGCATAAT |
| UQCRC2   | TTGCTTAAAAGCAGAGCATG | CTCAATCTCCAGGGTTTATC |
| VLCAD    | CATCCTCACAACGAGGAATT | CCAAACCTGGTACGAGT |

**Table 1.** List of mouse primers used in RT-qPCR experiments.
ide (CO₂) production (VCO₂) and oxygen (O₂) consumption (VO₂) rates were measured simultaneously using recorded values from a 72 h period.

**GTTs and ITTs.** For the GTTs, mice on either chow or a HFD were subjected to a 16 h fast with free access to water and then injected intraperitoneally with 2 g of 20% d-glucose per kilogram of body weight. The ITTs were performed similarly, with an initial 16 h of fasting and subsequent intraperitoneal injection of 0.75 U of insulin per kilogram of body weight. Blood glucose levels were measured at 0, 15, 30, 60, and 120 min with a OneTouch ultramini glucose meter (OneTouch).

**Serum content quantification.** Blood was extracted from cardiac puncture just before sacrifice and incubated with Aprotinin from bovine lung (0.03 TIU, Sigma-Aldrich, cat# A6279) for 2 h at 4 °C. The coagulated blood was spun 10 min at 800 x g at 4 °C and the supernatant (serum) stored at -80 °C. Serum total cholesterol, serum free cholesterol, and serum cholesterol esters were quantified using Cholesterol/ Cholesteryl Ester Assay Kit (Abcam, cat# ab65359) according to manufacturer's instructions. Serum Triglyceride and serum free glycerol were quantified using Serum Triglyceride Determination Kit (Sigma-Aldrich, cat# TR0100) according to manufacturer's instructions. Serum alanine aminotransferase was quantified using ALT determination kit (Pointe Scientific, cat# A7526) according to manufacturer's instructions.

**Liver triglyceride quantification.** Approximately 10 mg of frozen liver tissues were washed with PBS, resuspended in a 2:1 chloroform:methanol solution, and solubilized with a micropestle. The mixture was spun for 10 min at 15,000 x g at 4 °C and the supernatant incubated at 50 °C until the chloroform was completely evaporated. The dried pellet was dissolved in a solution containing 60% Butanol, 25% Triton X-100, and 15% Methanol and quantified using Serum Triglyceride Determination Kit (Sigma-Aldrich, cat# TR0100) according to manufacturer's instructions.

**Quantitative real-time PCR.** Total RNA was isolated and purified from approximately 25 mg of frozen liver tissue or cells using Total RNA Mini Kit (Geneaid, cat# RT100) according to the manufacturer's instructions. For quantitative real-time PCR analysis, 0.5 μg of total RNA was reverse-transcribed using the iScript Reverse Transcription Supermix for RT-qPCR (BioRad, cat# 1708841). SYBR Green reactions using the SYBR Green qPCR Supermix (BioRad, cat# 1,725,125) and specific primers (Table 1) were performed using a CFX Connect Real-Time PCR Detection System (BioRad). Relative expression of mRNAs was determined using the software CFX Maestro (BioRad, version 4.1.2434.0124, https://www.biorad.com/en-ca/product/cfx-maestro-software-for-cfx-real-time-pcr-instruments?ID=OKZP7E15) after normalization against housekeeping genes B2M, TBP, RPLP0, and PTP4a1.

**Immunohistochemistry.** Liver tissues embedded in paraffin were cut to 4-μm sections on slides and stained for α-SMA, 4-HNE, F4/80, TFE3 and TFEB using routine immunohistochemical protocols provided by the GCRC Histology Core using Ventana BenchMark ULTRA system (Roche). Images were acquired using Aperio Scanscope XT (Leica Biosystems) and staining quantification was performed using image analysis algorithms in Aperio ImageScope software (version 12.3.3.5048, https://www.leicabiosystems.com/fr/imagerie-pathologique/integre/aperio-imagescope/).

**Mouse protein cytokine array.** Approximately 100 mg of liver tissue were solubilized in RIPA buffer using a micropestle and sonicated 10 s. The mixture was then centrifuged at 16,000 x g for 15 min at 4 °C and the supernatant collected. 31 mouse cytokine/chemokine biomarkers were simultaneously quantified by using a Discovery Assay called the Mouse Cytokine Array / Chemokine Array 31-Plex (Eve Technologies Corp, cat# MD31).

**RNA-seq analysis.** Total RNA was isolated and purified from approximately 25 mg of frozen liver tissue using Total RNA Mini Kit (Geneaid, cat# RT100) according to the manufacturer's instructions. Library preparation and sequencing was made at the Institute for Research in Immunology and Cancer's Genomics Platform (IRIC). 500 ng of total RNA was used for library preparation. RNA quality control was assessed with the Bioanalyzer RNA 6000 Nano assay on the 2100 Bioanalyzer system (Agilent Technologies). Library preparation was done with the KAPA mRNAseq Hyperprep kit (KAPA, cat# KK8581). Ligation was made with Illumina dual-index UMI (IDT) and 10 PCR cycles was required to amplify cDNA libraries. Libraries were quantified by QuBit and BioAnalyzer DNA1000. All libraries were diluted to 10 nM and normalized by qPCR using the KAPA library quantification kit (KAPA, cat# KK4973). Libraries were pooled to equimolar concentration. Sequencing was performed with the Illumina Nextseq500 using the Nextseq High Output 75 (1 × 75 bp) cycles kit using 2.6 pM of the pooled libraries. Around 25 M single-end PF reads were generated per sample.

Following data acquisition, adaptor sequences and low quality score bases (Phred score < 30) were first trimmed using Trimmomatic41. The resulting reads were aligned to the GRCh38 mouse reference genome assembly, using STAR44, and read counts were obtained using HTSeq46. For all downstream analyses, we excluded lowly-expressed genes with an average read count lower than 10 across all samples, resulting in 14,057 expressed genes in total. The R package limma46 was used to identify differences in gene expression levels between the different conditions. Nominal p-values were corrected for multiple testing using the Benjamini–Hochberg method. To assess the effect liver-Flcn KO in the response to MCD-diet, we first obtained differentially expressed genes (FDR < 0.05 and |log2FC|> 1) in Cre.MCD_diet vs Cre.Chow and Flcn-KO.MCD_diet vs Flcn-KO.Chow, and
then filtered for those that show \(\text{difference in log2FC}>1\) (differentially responsive (DR) genes). Unsupervised hierarchical clustering of the 1,110 DR genes shows three distinct patterns of changes in expression. Pathway enrichment analyses were performed using Enrichr۵.۷

### Statistical analyses

Data are expressed as mean ± SEM. All experiments and measurements were performed on at least 3 mice as indicated. Statistical analyses for all data were performed using student’s t-test, one-way ANOVA or two-way ANOVA as indicated using GraphPad Prism 7 software (version 7.0a, https://www.graphpad.com/scientific-software/prism/). Statistical significance is indicated in figures (* \(P < 0.05\), ** \(P < 0.01\), *** \(P < 0.001\), **** \(P < 0.0001\)).

### Data availability

RNA-sequencing data has been deposited in the Gene Expression Omnibus under the accession GSE156918.

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Competing interests
The authors declare no competing interests.

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