

# Lactate transporter MCT1 in hepatic stellate cells promotes fibrotic collagen expression in nonalcoholic steatohepatitis

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## Abstract

Circulating lactate is a fuel source for liver metabolism but may exacerbate metabolic diseases such as nonalcoholic steatohepatitis (NASH). Indeed, haploinsufficiency of lactate transporter monocarboxylate transporter 1 (MCT1) in mice reportedly promotes resistance to hepatic steatosis and inflammation. Here, we used adeno-associated virus (AAV) vectors to deliver thyroxin binding globulin (TBG)-Cre or lecithin-retinol acyltransferase (Lrat)-Cre to MCT1<sup>f/f</sup> mice on a choline deficient, high fat NASH diet to deplete hepatocyte or stellate cell MCT1, respectively. Stellate cell MCT1KO (AAV-Lrat-Cre) attenuated liver type 1 collagen protein expression and caused a downward trend in trichrome staining. MCT1 depletion in cultured human LX2 stellate cells also diminished collagen 1 protein expression. Tetra-ethylenglycol-cholesterol (Chol)-conjugated siRNAs, which enter all hepatic cell types, and hepatocyte-selective tri-N-acetyl galactosamine (GN)-conjugated siRNAs were then used to evaluate MCT1 function in a genetically obese NASH mouse model. MCT1 silencing by Chol-siRNA decreased liver collagen 1 levels, while hepatocyte-selective MCT1 depletion by AAV-TBG-Cre or by GN-siRNA unexpectedly increased collagen 1 and total fibrosis without effect on triglyceride accumulation. These findings demonstrate that stellate cell lactate transporter MCT1 significantly contributes to liver fibrosis through increased collagen 1 protein expression *in vitro* and *in vivo*, while hepatocyte MCT1 appears not to be an attractive therapeutic target for NASH.

## Introduction

Nonalcoholic fatty liver disease (NAFLD) is the most common chronic liver disease, afflicting over a quarter of the world's population. It describes a spectrum of liver diseases ranging from simple steatosis to nonalcoholic steatohepatitis (NASH)<sup>1,2</sup>. Steatosis is considered relatively benign as lifestyle modifications can reverse fatty liver to a healthy condition. On the other hand, NASH is characterized by severe steatosis, inflammation, and fibrosis. Severe fibrotic stages of NASH can develop into permanent liver damage, and the disease can progress to cirrhosis and hepatoma. Currently, NASH is a leading cause for liver transplantation, and there is no FDA-approved therapeutic for NASH<sup>2-5</sup>. Since NASH is closely associated with the hallmarks of type 2 diabetes and obesity, including chronic overnutrition, insulin resistance and dyslipidemia, it is likely that substrate overload to the liver contributes to its cause<sup>5</sup>. Accordingly, NAFLD and NASH patients display hyperactive liver tricarboxylic acid (TCA) cycle flux due to an overabundance of upstream metabolites<sup>6,7</sup>. These findings suggest that lowering substrate influx to the liver is a promising strategy to prevent and possibly alleviate steatosis and NASH.

Human subjects with type 2 diabetes and obesity reportedly have increased plasma lactate levels<sup>8-14</sup>. Lactate is produced and released into the circulation when cellular glycolytic flux surpasses mitochondrial oxidative capacity. Once considered to be simply a metabolic waste product, lactate is now recognized as a primary fuel for the TCA cycle in liver and thus an essential energy source<sup>15,16</sup>. Additionally, it is a critical regulator that contributes to whole-body energy homeostasis<sup>17,18</sup>. Under physiological conditions,

cellular lactate levels are tightly controlled by monocarboxylate transporters (MCTs). MCTs are members of the solute carrier 16A (SLC16A) family, which are proton-coupled transmembrane protein transporters. Among 14 MCT isoforms, only MCTs 1-4 have been shown to transport monocarboxylate molecules such as lactate, pyruvate, short-chain fatty acids, and ketone bodies<sup>19</sup>. Notably, MCT1/SLC16A1 is denoted as a primary lactate transporter as it is the most widely distributed MCT isoform in various metabolic tissues and has a high affinity for lactate, maintaining basal cellular homeostasis according to transmembrane lactate gradients<sup>18,20</sup>. Reportedly, MCT1 haploinsufficiency in mice reduces MCT1 protein levels to nearly half in major metabolic tissues such as liver, brain, and white adipose tissues, and these mice are resistant to diet-induced obesity and liver steatosis and inflammation<sup>21-23</sup>. The role of MCT1 in hypothalamus and adipose tissues in these phenotypes was ruled out, as selective MCT1 depletion in those tissues either increased food intake and body weight<sup>24</sup> or enhanced systemic inflammation and insulin resistance<sup>25</sup>. Thus, the question of which tissue or tissues are responsible for the phenotype of whole body MCT1 haploinsufficiency is not solved.

The above considerations suggest the possibility that MCT1KO in one or more liver cell types may explain the effects of MCT1 haploinsufficiency in mice. Since hepatocytes account for the majority of liver cells and have high rates of lipogenesis and triglyceride (TG) accumulation, lactate levels governed by hepatocyte MCT1 could be involved in regulating steatosis. On the other hand, while hepatic stellate cells account for only 5-10% of the hepatic cell population, they are the major cell type contributing to hepatic fibrogenesis<sup>26, 27</sup>. Fate tracing studies have revealed that 82-96% of myofibroblasts are

derived from hepatic stellate cells, which are liver-specific pericytes<sup>27</sup>. During NASH progression, multiple liver injury signals stimulate the transition of vitamin A-storing quiescent hepatic stellate cells into fibrogenic, proliferative myofibroblasts that produce and secrete collagen fibers<sup>28-31</sup>. As a result, healthy hepatic parenchyma is replaced with a collagen-rich extracellular matrix, turning the liver into a hardened and scarred tissue<sup>32</sup>. In general, major organ fibrosis is directly correlated with morbidity and mortality, contributing up to 45 % of deaths in developed countries<sup>33</sup>. Thus, targeting activated hepatic stellate cells has become a major strategy in NASH therapeutics development<sup>5,31</sup>. However, the role of MCT1 in hepatic stellate cells activation or fibrogenesis has not been investigated.

The aim of the present studies was to investigate the role of hepatic lactate transport via MCT1 in lipid metabolism and fibrogenesis in NASH, and to determine its potential suitability as a therapeutic target. Two key unanswered questions were of particular interest: 1) is it hepatocyte-specific MCT1 depletion that protects mice with MCT1 haploinsufficiency from liver lactate overload and NAFLD and 2) does liver stellate cell MCT1 promote hepatic fibrogenesis that occurs in NASH? We tested the possible enhancement of lipogenesis and fat accumulation via MCT1 function specifically in hepatocytes using adeno-associated virus (AAV) mediated thyroxin binding globulin (TBG)-Cre MCT1KO in MCT1<sup>f/f</sup> mice, and in other experiments by silencing hepatocyte MCT1 with tri-N-acetyl galactosamine (GN)-conjugated siRNA. These experiments showed hepatocyte MCT1 loss decreased expression of enzymes in the *de novo* lipogenesis (DNL) pathway, but did not diminish overall steatosis. Surprisingly,

hepatocyte MCT1KO increased liver fibrosis in two mouse models of NASH. In contrast, hepatic stellate cell-selective MCT1KO, achieved by injection of AAV9-lecithin-retinol acyltransferase (*Lrat*)-Cre into *MCT1<sup>f/f</sup>* mice, did attenuate collagen production and fibrosis. Our findings underscore the critical importance of implementing cell type-specific targeting strategies to diminish NASH fibrogenesis.

## Results

### **MCT1 depletion prevents transforming growth factor 1β (Tgf1β)-stimulated type 1 collagen production in cultured human LX2 stellate cells**

As fate tracing studies have revealed that 82-96% of myofibroblasts are derived from hepatic stellate cells<sup>27</sup>, we employed a simple *in vitro* system utilizing LX2 human hepatic stellate cells to investigate effects of MCT1 silencing on expression of type 1 collagen, a major component of fibrosis. Cells were transfected with Lipofectamine and native MCT1-targeting siRNA (MCT1-siRNA), which diminished *MCT1* expression by about 80% (Figure 1A), or nontargeted control (NTC)-siRNA, and then treated with Tgf1β (10ug/ml) for 48 hours. As expected, Tgf1β stimulated expression of ACTA2 and collagen 1 isoform, COL1A1, by several folds (Figure 1B, 1C). *MCT1* silencing significantly inhibited Tgf1β-stimulated ACTA2 mRNA expression as well as collagen 1 protein production (Figure 1B, 1C), indicating cell-autonomous functions of MCT1 in hepatic stellate cells.

### **Identification of a potent, chemically modified siRNA candidate targeting MCT1**

Given the therapeutic potential of *MCT1* silencing in preventing fibrogenesis (Figure 1), we aimed to develop MCT1-siRNA compounds, chemically modified for stability, potency and delivery *in vivo* for use in this research and potentially for therapeutic advancement.

Asymmetrical siRNA compounds used here are composed of 15 double-strand nucleotides with a short overhanging single-strand that promotes cellular uptake<sup>34,35</sup>. To enhance the stability of the constructs, the 2'-OH of each ribose was modified to either 2'-O-methyl or 2'-fluoro. In addition, phosphorothioate linkage backbone modifications were applied to avoid exonuclease degradation. Tetra-ethylenglycol-cholesterol linker (Chol) was conjugated to the 3' end-sense strand to enhance stability and cellular uptake of candidate compounds. Each Chol-conjugated, fully chemically modified MCT1-siRNA (Chol-MCT1-siRNA) candidate construct's sequence and targeting region on the *Mct1* transcript is described in Table 1 and Figure 2A.

We performed *in vitro* screening to select the most potent Chol-MCT1-siRNA compounds that were initially synthesized (Figure 2A, 2B). Each Chol-MCT1-siRNA compound candidate was transfected into mouse hepatocyte FL83B cells. As opposed to native siRNA, our Chol-MCT1-siRNA does not require transfection reagents as it is fully chemically modified. The silencing effect on *Mct1* mRNA was monitored after 72 hours (Figure 2B). Several compounds elicited a silencing effect greater than 80% compared to the NTC-siRNA. The two most potent Chol-MCT1-siRNA, Chol-MCT1-2060 (IC50: 59.6nM, KD%: 87.2), and Chol-MCT1-3160 (IC50: 32.4nM, KD%: 87.7) were evaluated for their inhibitory effect on MCT1 protein levels (Figure 2D, 2E). Based on its IC50 value and silencing potency, Chol-MCT1-3160 construct was chosen for further studies *in vivo* (Table 2).

### **Distinct cellular biodistribution of Chol- vs GN-conjugated siRNAs**

For *in vivo* studies, further chemical modifications were applied (Figure 3A). MCT1-siRNAs utilized in *in vivo* studies are double-strand oligonucleotides comprised of 18

sense and 20 antisense nucleotides. At the 5'-end of the antisense strand, a 5'-(E)-Vinyl-phosphonate modification was added to prevent phosphatase-induced degradation, enhancing *in vivo* stability and promoting its accumulation in target cells. Either a hydrophobic Chol or a hepatocyte-targeting GN was attached to the 3'-end of sense strands of MCT1-siRNAs to direct different hepatic cellular biodistribution.

To validate the biodistribution of siRNAs with these two conjugates, 10mg/kg of each siRNA was subcutaneously injected into 16-18 weeks-old male C57BL/6 wild-type mice twice within a 15 day period. On day 15, mice were sacrificed and the livers were perfused to isolate multiple hepatic cell types, including hepatocytes, stellate cells, and Kupffer cells. Isolation of each hepatic cell type was validated for enrichment (Supplement Figure 1A-1F). As expected, GN-conjugated, fully chemically modified MCT1-siRNA (GN-MCT1-siRNA) silenced *Mct1* mRNA only in the hepatocyte fraction (Figure 3B-3D), as GN binds to the asialoglycoprotein receptor primarily expressed in hepatocytes. On the other hand, Chol-MCT1-siRNA silenced *Mct1* mRNA levels in all hepatic cell types (Figure 3E-3G), as its cellular uptake is highly dependent on the non-specific, hydrophobic interaction between cholesterol and plasma membranes. Notably, the hepatic stellate cell fraction distinguishes GN-MCT1-siRNA from Chol-MCT1-siRNA in biodistribution, as only the latter silences *Mct1* in stellate cells (Figure 3C vs 3F). We also confirmed that both GN-MCT1-siRNA and Chol-MCT1-siRNA do not affect MCT1 levels in other major metabolic tissues (Supplement Figure 1G-1J).

**Subcutaneous injection of Chol-MCT1-siRNA or GN-MCT1-siRNA silences hepatic MCT1 in a genetically obese NASH mouse model**

We next investigated the effect of Chol-MCT1-siRNA on reversing severe steatosis in the genetically obese ob/ob mouse on a NASH-inducing Gubra Amylin NASH (GAN)<sup>36</sup> diet. These mice normally develop severe steatosis from an early age but hardly develop fibrosis until fed the GAN diet. Each siRNA (10mg/kg) was subcutaneously injected once every 10 days and mice were fed a GAN diet for 3 weeks before sacrifice (Figure 4A). Hepatic MCT1 protein levels were visually monitored by MCT1-positive staining immunohistochemistry (Figure 4B, 4C), showing more than 70% MCT1 protein depletion (Chol-MCT1-siRNA: 77.99% & GN-MCT1-siRNA: 71.35% silencing). Similar silencing was observed when *Mct1* mRNA levels were measured by rt-qPCR (Figure 4D). Importantly, there was no surge in plasma lactate level (Figure 4E), addressing the concern of potential lactic acidosis after MCT1 depletion in the liver, the major lactate-consuming tissue. We also monitored food intake and body weight over time (Supplement Figure 2A, 2B), as there was a report of decreased food anticipation activity upon hepatic MCT1 deletion followed by reduced plasma β-hydroxybutyrate levels<sup>37</sup>. Intriguingly, the GN-MCT1-siRNA administration led to a decrease in both food intake and body weight, while the Chol-MCT1-siRNA did not. Neither Chol-MCT1-siRNA administration nor hepatocyte-specific MCT1KO improved glucose tolerance on the genetically obese NASH mouse model or a 12 week HFD-induced NAFLD model, respectively (Supplement Figure 3A, 3B).

**Hepatic MCT1 depletion downregulates lipogenic genes but not steatosis in the ob/ob NASH diet mouse model.**

In order to fully analyze steatosis in NASH, lipid droplet morphology and total hepatic TG was assessed (Figure 4F-4H). The results showed that neither GN-MCT1-siRNA nor

Chol-MCT1-siRNA decreased total hepatic TG levels (Figure 4H), although quantitative analysis of H&E images showed a small decrease in mean lipid droplet size and increased number of lipid droplets upon MCT1 silencing (Figure 4F, 4G). These data suggest the possibility that hepatic MCT1 depletion either 1) inhibits formation or fusion of lipid droplets, or 2) enhances lipolysis to diminish lipid droplet size.

To investigate the underlying mechanism by which lipid droplet morphological dynamics change, we monitored the effect of hepatic MCT1 depletion on DNL-related gene expression. Both GN-MCT1-siRNA and Chol-MCT1-siRNA strongly decreased the mRNA and protein levels related to representative DNL genes (Supplement Figure 4A-4D). Intriguingly, both modes of hepatic MCT1 depletion also inhibited expression of the upstream regulatory transcription factors SREBP1 and ChREBP. Because phosphorylated AMPK (pAMPK), an active form of AMPK, is known to inhibit SREBP nuclear translocation<sup>38</sup> as well as the DNA binding activity of ChREBP<sup>39</sup>, we evaluated pAMPK levels. As a result, there was a significant increase in pAMPK levels and pAMPK/AMPK ratio in both GN-MCT1-siRNA and Chol-MCT1-siRNA injected groups (Supplement Figure 4E, 4F).

### **Opposite effects of Chol-MCT1-siRNA versus GN-MCT1-siRNA on fibrotic collagen expression**

We next monitored fibrosis, a tissue damaging phenotype that is associated with NASH. Consistent with the results in LX2 stellate cells (Figure 1), Chol-MCT1-siRNA administration to ob/ob mice on GAN diet significantly reduced liver collagen 1 protein levels (Figure 5A, 5B). This result could be attributable to the fact that subcutaneous injection of the Chol-MCT1-siRNA compound is able to silence genes in hepatic stellate

cells, the predominant cell type that produces collagens. Interestingly, decreases in mRNA encoding collagen 1 isoforms were not detected by rt-qPCR analysis (Figure 5C), indicating possible effects at the level of translation or protein turnover. Surprisingly, an opposite phenotype on collagen 1 protein expression was observed in response to administration of GN-MCT1-siRNA compared to Chol-MCT1-siRNA (Figure 5D, 5E). Hepatocyte-specific GN-MCT1-siRNA actually enhanced the expression of type 1 collagen protein in these experiments (Figure 5D, 5E), and this effect was also apparent at the mRNA expression level (Figure 5F). Overall fibrosis as detected by Sirius Red was also analyzed in these experiments, as this staining detects all types of collagen fibers that are involved in hepatic fibrosis, such as III, IV, V, and VI. In line with collagen 1 mRNA and protein levels, GN-MCT1-siRNA significantly enhanced Sirius red positive areas in the images, as shown in Figure 5G, 5H. Despite its inhibitory effect on collagen 1 production levels, Chol-MCT1-siRNA did not reduce Sirius Red positive areas, suggesting that reduction of major type 1 collagen might not be the major contributor to fibrosis in this model or that compensatory effects are occurring.

### **MCT1KO by AAV-Lrat-Cre and AAV-TBG-Cre constructs confirm cell type specificity of MCT1KO effects**

The results presented above suggested that Chol-MCT1-siRNA downregulates type 1 collagen protein by depleting MCT1 in hepatic stellate cells, which hepatocyte-specific GN-MCT1-siRNA cannot target. To test this hypothesis, we developed and validated AAV9-Lrat-Cre constructs to generate hepatic stellate cell-specific MCT1 knockout mice. Male MCT1<sup>f/f</sup> mice<sup>40</sup> were intravenously injected with AAV9-Lrat-Cre (1X10<sup>11</sup>gc) and sacrificed 3 weeks later (Supplement Figure 5A). Isolation of hepatocytes and hepatic

stellate cells were validated with their representative marker genes encoding albumin, and desmin, respectively (Supplement Figure 5B, 5C). Successful depletion of hepatic stellate cell selective *Mct1* mRNA was confirmed in the MCT1<sup>f/f</sup> mice injected with the AAV9-Lrat-Cre construct. *Mct1* mRNA levels in the hepatocytes, which accounts for up to 70% of total liver cell types, were intact (Supplement Figure 5D, 5E). No change in MCT1 protein level was observed in other metabolic tissues (Supplement Figure 5F, 5G).

We employed the choline-deficient HFD-induced NASH mouse model (CDHFD) to test these AAV constructs on steatosis and fibrosis. CDHFD induces severe steatosis due to inhibited VLDL secretion and β-oxidation and thereby exacerbates NASH fibrosis in a relatively short time<sup>41,42</sup>. MCT1<sup>f/f</sup> mice were intravenously injected with 2X10<sup>11</sup>gc of AAV8-TBG-Cre or AAV9-Lrat-Cre or both (Figure 6A). A week after the injections, mice were fed with a CDHFD for 8 weeks to induce NASH. Depletion of hepatic *Mct1* mRNA in each group was confirmed (Figure 6B). There was no food intake or body weight difference between the groups (Figure 6C, 6D). Similar to results we obtained by MCT1 silencing with siRNAs (Figure 4), MCT1 deletion in either hepatocytes or in hepatic stellate cells did not resolve steatosis (Figure 6E, 6F).

Also consistent with results obtained by hepatocyte selective MCT1 silencing with GN-MCT1-siRNA (Figure 5), hepatocyte-specific knockout of MCT1 (Hep KO) enhanced the collagen 1 level compared to the control group (Figure 7A). In contrast, hepatic stellate cell-specific MCT1 knockout (HSC KO) prevented CDHFD-induced collagen 1 protein levels (Figure 7B). MCT1KO in combined hepatocytes and hepatic stellate cells blunted the effect shown in each single KO (Figure 7C). Overall liver fibrosis detected by trichrome staining again confirmed the acceleration of fibrosis in the Hep KO group and a downward

trend in the HSC KO group (Figure 7D, 7E). Dual MCT1KO in hepatocytes plus hepatic stellate cells showed no change in overall fibrosis, similar to the Chol-MCT1-siRNA results (Figure 5G, 5H). Additionally, liver stiffness was monitored via ultrasound-based shear wave elastography (SWE) in a noninvasive diagnostic mode for liver disease<sup>43,44</sup>. After 8 weeks of CDHFD, all groups had the same level of increased liver stiffness above what the control mice showed at 4 weeks, however, Hep KO mice exhibited elevated liver stiffness over all other groups at 4 weeks of the diet (Figure 7F, 7G). There was no change in plasma alanine transaminase (ALT) levels among the groups (Figure 7H).

## Discussion

The major finding of this study is that MCT1 function in hepatic stellate cells promotes collagen 1 expression, as MCT1 depletion in this cell type, either *in vitro* in cell culture (Figure 1B) or *in vivo* in mice (Figure 7B), attenuates fibrotic collagen 1 protein production. This finding in mice was made by generating novel AAV9-Lrat-Cre constructs that can be injected into MCT1<sup>f/f</sup> mice to elicit stellate cell selective MCT1 depletion, as verified by isolation and *Mct1* mRNA analysis in liver cell types (Supplement Figure 5). Previous use of Lrat-Cre for germline transmission had validated constitutive gene KO selectively in hepatic stellate cells in mice<sup>27</sup>, while our AAV-Lrat-Cre construct allows inducible gene KO, eliminating time-demanding mouse crossing and breeding. Surprisingly, MCT1KO in hepatocytes, both *in vivo* (Figure 7A) and *in vitro* (Supplement Figure 6B), evoked the opposite effect: a robust upregulation of collagen 1 expression and fibrosis. This may be in part a cell-autonomous effect in hepatocytes, as we observed increased collagen 1 expression in Hep2G cells in culture upon silencing *MCT1* (Supplement Figure 6). Interestingly, the inhibitory effect of stellate cell MCT1KO was observed only at the

collagen 1 protein level, while the hepatocyte MCT1KO affected both collagen mRNA and protein. In addition, the hepatocyte selective MCT1KO caused increased total fibrosis, evidenced by trichrome staining of liver (Figure 7D, 7E), while the stellate cell MCT1KO was associated with a trend towards diminished trichrome staining that did not reach statistical significance (Figure 7D, 7E). Nonetheless, taken together, our data suggest that lactate flux in hepatic stellate cells strongly promotes collagen 1 translation or inhibits protein turnover rates (Supplement Figure 7: Graphical abstract).

Strong support for the above conclusions was obtained in an alternative experimental model of NASH--genetically obese ob/ob mice on a GAN diet<sup>36</sup>. In these experiments we employed chemically stabilized siRNA, taking advantage of RNA modifications 2-Fluoro, 2-O-Methyl ribose, and phosphorothioate backbone replacement to block siRNA degradation and immune responses as well as enhance *in vivo* delivery effectiveness and silencing longevity<sup>34,35</sup>. Conjugation of such siRNA compounds with GN promotes binding to the asialoglycoprotein receptor that is primarily expressed in hepatocytes at high levels, and we confirmed hepatocyte selective gene silencing with such constructs (Figure 3B-3D). Moreover, injection of GN-MCT1-siRNA did not affect MCT1 expression in other major tissues including iWAT, gWAT, BAT, intestine, heart, lung, kidney, and spleen (Supplement Figure 1G, 1H). Hepatocyte selective MCT1 silencing strongly upregulated collagen 1 in livers of ob/ob mice on a GAN diet (Figure 5D-5F), as did hepatocyte selective MCT1KO in the CDHFD mouse model (Figure 7A). In contrast, cholesterol-conjugated, chemically modified siRNA targeting MCT1 silenced the gene in all three liver cell types tested, including hepatic stellate cells (Figure 3E-3G), and did

attenuate liver collagen 1 expression (Figure 5A, 5B). This result is consistent with the idea that depleting MCT1 in stellate cells decreases collagen 1 production, as shown by MCT1KO in stellate cells (Figure 7B). The data in Figure 3 and Supplement Figure 1 also show the effectiveness of RNA interference (RNAi) in interrogating cell specific processes in liver, and highlight the multiple advantages over traditional small molecule inhibitors in developing therapeutics<sup>45,46</sup>. To date, five RNAi-based therapeutic agents have received FDA approvals targeting multiple disease areas, and a great many clinical trials of oligonucleotide therapeutics are in progress<sup>47-50</sup>.

Analysis of gene expression profiles also showed that hepatic MCT1 positively regulates the levels of SREBP1 and ChREBP, major transcription factors regulating liver lipid metabolism, as well as their target DNL genes. These effects were apparently not sufficient to reverse severe steatosis in the genetically obese NASH mouse model (Figure 4), although a slight decrease in mean lipid droplet size was observed. The remaining steatosis may be attributed to the continuous supply of fatty acids from adipose tissue lipolysis, which accounts for up to 65% of hepatic fat accumulation as opposed to only 25% coming from DNL<sup>51</sup>. We also cannot rule out the possibility of compensatory effects from other MCT isoforms that are expressed. However, since MCT1 haploinsufficiency showed greatly reduced HFD-induced hepatic steatosis, the discrepancies with our data may be due to other tissues being involved or the different mouse models used. Overall, it is clear from our studies that steatosis is not much affected by hepatic MCT1 depletion in two models of NASH in mice. A possible future strategy for NASH therapeutics may be to combine the depletion of MCT1 in stellate cells to decrease fibrosis with a potent anti-

steatosis target such as DGAT2<sup>36,52,53</sup> which we and others have shown is effective in reducing steatosis when depleted. Although there are still remaining challenges in developing a successful hepatic stellate cell-selective delivery system<sup>54</sup>, ongoing studies are identifying promising candidates such as M6P-polyethylene glycol (M6P-PEG)<sup>55</sup>, IGF2R-specific peptide coupled nanocomplex<sup>56</sup>, and others<sup>57</sup>.

To understand the underlying mechanism by which hepatic MCT1 depletion drives the downregulation of DNL genes expression, we investigated AMPK activation, as phosphorylated AMPK has a negative regulatory effect on SREBP1 and ChREBP activation<sup>38,39</sup>. Indeed, hepatic MCT1 silencing enhanced AMPK phosphorylation, consistent with the previous MCT1 haploinsufficient mice study<sup>22</sup>. These data are also in line with another study in which MCT1 inhibition reduced ATP production and activated AMPK, thus deactivating SREBP1c and lowering levels its target SCD1<sup>58</sup>. It remains to be examined whether other mechanisms are also at play that connect MCT1 function in liver to DNL gene regulation.

In summary, the data presented here highlight hepatic stellate cell MCT1 as a potential therapeutic target to prevent NASH fibrogenesis related to collagen 1 production. Its utility as a therapeutic target is complicated by our finding that MCT1 depletion in hepatocytes actually increases fibrosis. This work highlights the importance of contemplating cell-type specificity when developing therapeutic strategies, especially in systems of complex cellular landscapes such as NASH.

## Materials and methods

### Oligonucleotide synthesis

The 15-20 or 18-20 sequence oligonucleotides were synthesized by phosphoramidite solid-phase synthesis on a Dr Oligo 48 (Bolytic, Fremont, CA), or MerMade12 (Biosearch Technologies, Novato, CA) using 2'-F and 2'-O-Me phosphoramidites with standard protecting groups (Chemgenes, Wilmington, MA). For the 5'-VP coupling, 5'-(E)-Vinyl tetraphosphonate (pivaloyloxymethyl) 2'-O-methyl-uridine 3'-CE phosphoramidite was used (Hongene Biotech, Union City, CA). Phosphoramidites were prepared at 0.1 M in anhydrous acetonitrile (ACN), except for 2'-O-methyl-uridine phosphoramidite dissolved in anhydrous ACN containing 15% dimethylformamide. 5-(Benzylthio)-1H-tetrazole (BTT) was used as the activator at 0.25 M, coupling time for all phosphoramidites was 4 min. Detritylations were performed using 3% trichloroacetic acid in dichloromethane. Capping reagents used were CAP A (20% n-methylimidazole in ACN) and CAP B (20% acetic anhydride and 30% 2,6-lutidine in ACN). Phosphite oxidation to convert to phosphate or phosphorothioate was performed with 0.05 M iodine in pyridine-H<sub>2</sub>O (9:1, v/v) or 0.1 M solution of 3-[(dimethylaminomethylene)amino]-3H-1,2,4-dithiazole-5-thione (DDTT) in pyridine respectively. All synthesis reagents were purchased from Chemgenes. Unconjugated oligonucleotides were synthesized on 500Å long-chain alkyl amine (LCAA) controlled pore glass (CPG) functionalized with Unylinker terminus (ChemGenes). Chol conjugated oligonucleotides were synthesized on a 500Å LCAA-CPG support, functionalized with a tetra-ethylenglycol cholesterol moiety bound through a succinate linker (Chemgenes). GN conjugated oligonucleotides were grown on a 500Å LCAA- CPG functionalized with an aminopropanediol-based trivalent GalNAc cluster (Hongene).

## **Deprotection and purification of oligonucleotides for screening of sequences**

Prior to the deprotection, synthesis columns containing oligonucleotides were treated with 10% diethylamine (DEA) in ACN to deprotect cyanoethyl groups. Synthesis columns containing the oligonucleotides covalently attached to the solid supports were cleaved and deprotected for 1 hour at room temperature with anhydrous Mono-Methylamine gas (Airgas). Columns with deprotected oligonucleotides were washed with 1mL of 0.1M sodium acetate in 85% ethanol aqueous solution, followed by rinse with an 85% ethanol aqueous solution. The excess ethanol was dried from the column on a vacuum manifold. Finally, the oligonucleotides were eluted off the columns with MilliQ water.

## **Deprotection and purification of oligonucleotides for *in vivo* experiments**

Chol- or GN-conjugated oligonucleotides were cleaved and deprotected with 28-30% ammonium hydroxide and 40% aq. Methylamine in a 1:1 ratio for 2h at room temperature. VP-containing oligonucleotides were cleaved and deprotected as described previously. Briefly, CPG with VP-oligonucleotides was treated with a solution of 3% diethylamine in 28–30% ammonium hydroxide for 20 hours at 35°C. The cleaved oligonucleotide solutions were filtered to remove CPG and dried under a vacuum. The pellets were resuspended in 5% ACN in water and purified on an Agilent 1290 Infinity II HPLC system. VP and GN-conjugated oligonucleotides were purified using a custom 20 x 150mm column packed with Source 15Q anion exchange resin (Cytiva, Marlborough, MA). Run conditions were the following. Eluent A: 20 mM sodium acetate in 10% ACN in water. Eluent B: 1 M sodium bromide in 10% ACN in water. Linear gradient 10–35% B 20 min at 40°C. Chol-conjugated oligonucleotides were purified using 21.2 x 150mm PRP-C18 column (Hamilton Co, Reno, NV). Run conditions were the following: Eluent A, 50 mM

sodium acetate in 5% ACN in water; Eluent B: 100% ACN. Linear gradient, 40–60% B 20 min at 60°C. Flow used was 40mL/min for both systems. Peaks were monitored at 260 nm. Fractions collected were analyzed by liquid chromatography-mass spectrometry (LC-MS). Pure fractions were combined and dried under a vacuum and resuspended in 5% ACN. Oligonucleotides were desalted by size exclusion on a 50 x 250mm custom column packed with Sephadex G-25 media (Cytiva, Marlborough, MA), and lyophilized. Reagents for deprotection and purification were purchased from Fisher Scientific, Sigma-Aldrich, and Oakwood Chemicals.

### **LC–MS analysis of oligonucleotides**

The identity of oligonucleotides is verified by LC–MS analysis on an Agilent 6530 accurate mass Q-TOF using the following conditions: buffer A: 100 mM 1,1,1,3,3,3-hexafluoroisopropanol (HFIP) (Oakwood Chemicals) and 9 mM triethylamine (TEA) (Fisher Scientific) in LC–MS grade water (Fisher Scientific); buffer B: 100 mM HFIP and 9 mM TEA in LC–MS grade methanol (Fisher Scientific); column, Agilent AdvanceBio oligonucleotides C18; linear gradient 0–35% B 5min was used for unconjugated and GN-conjugated oligonucleotides; linear gradient 50–100% B 5min was used for Cholesterol conjugated oligonucleotides; temperature, 60°C; flow rate, 0.85 ml/min. LC peaks are monitored at 260nm. MS parameters: Source, electrospray ionization; ion polarity, negative mode; range, 100–3,200 m/z; scan rate, 2 spectra/s; capillary voltage, 4,000; fragmentor, 200 V; gas temp, 325°C.

### **LX2 human hepatic stellate cell studies**

Human hepatic stellate cell line, LX2, was purchased from Millipore Sigma, cat SCC064. LX2 cells were cultured in DMEM/high glucose media (Gibco, cat 11995-065 and Fisher,

cat 11965092) with 10% FBS. To test the preventative effect of MCT1 depletion in Tgf1 $\beta$ -stimulated hepatic stellate cell conditions, LX2 cells were plated in 6 well plates (300k cells/well) or 12 well plates (150k cells/well) in DMEM/high glucose media with 2% FBS. The next day, cells were transfected with either NTC-siRNA or MCT1-siRNA (IDT, cat 308915476) using lipofectamine RNAi Max (Thermofisher, cat 13778075) for 6 hours in less serum optiMEM media (Thermofisher, cat31985062). Then, cells were maintained in serum-starved media with or without 10ng/ml of recombinant human Tgf1 $\beta$  (R&D Systems, cat 240-B/CF) for 48 hours and harvested. As a housekeeping gene,  $\beta$ -actin (ACTB) was used.

### **Human HepG2 hepatoma cell studies**

Human hepatoma cell line, HepG2, was purchased from ATCC, cat HB-8065. Cells were cultured in RPMI media (Gibco, cat 11875-093) with 10% FBS. To test the effect of MCT1 depletion, cells were plated in 6 well plates (300k cells/well) or 12 well plates (150k cells/well). The next day, cells were transfected with either NTC-siRNA or MCT1-siRNA using lipofectamine RNAi Max (Thermofisher, cat 13778075) for 6 hours in less serum optiMEM media (Thermofisher, cat31985062). After 48 hours, HepG2 cells were harvested. The media were saved to further test for secreted factors that may affect hepatic stellate cell activation. LX2 cells were incubated with the conditioned media (40% conditioned media + 60% fresh media), and cells were harvested after 48 hours.

### ***In vitro* screening of chemically modified siRNAs**

Mouse hepatocyte cell line FL83B was purchased from (ATCC, CRL-2390). FL83B cells were plated in 12 well plates (150k cells/well) in F-12K medium with 3% FBS. Then, 1.5uM

of each Chol-MCT1-siRNA candidate compound was added and Chol-NTC-siRNA was used as a control. Then, 72 hours after the treatment, cells were harvested, and the *Mct1* mRNA silencing potency was monitored. To further evaluate the half maximal inhibitory concentration (IC<sub>50</sub>) values, the dose-dependent silencing effect of the compounds was calculated upon six different concentrations (1.5uM, 0.75uM, 0.38uM, 0.19uM, 0.05uM, and 0uM). As a housekeeping gene, β-2-microglobulin (*B2m*) was used.

### **Generation and validation of hepatic stellate cell-specific AAV9-Lrat-Cre**

Lrat-Cre-mediated KO mice have been widely utilized in the field to delete genes in hepatic stellate cells<sup>27</sup>. We newly synthesized AAV9-Lrat-Cre to establish an inducible hepatic stellate cell KO system in collaboration with Vector Biolabs. Proximal mouse Lrat promoter region from -1166bp, including the putative transcriptional start site (TSS), to +262 bp downstream sequence was chosen<sup>59</sup>. A 1428b Lrat promoter was synthesized and cloned into Vector Biolabs' AAV-CMV-Cre vector to replace CMV promoter with Lrat promoter. The AAV-Lrat-Cre was then packaged into AAV9 virus. As a control, AAV-Lrat-null constructs were used.

### **Animal studies**

All animal procedures were performed in accordance with animal care ethics approval and guidelines of University of Massachusetts Chan Medical School Institutional Animal Care and Use Committee (IACUC, protocol number A-1600-19). All wild-type C57BL6/J male mice, and genetically obese ob/ob male mice were obtained from Jackson Laboratory. MCT1<sup>f/f</sup> mice were generated in the Rothstein lab<sup>40</sup>. Mice were group-housed on a 12-hour light/dark cycle and had *ad libitum* access to water and food. **For obese**

**NASH model studies**, 10 week old genetically obese ob/ob male mice (n=6) were subcutaneously injected with 10mg/kg of siRNAs accordingly (Chol-NTC-siRNA, Chol-MCT1-siRNA, GN-NTC-siRNA, and GN-MCT1-siRNA), every 10-12 days. Mice were fed the GAN diet (Research diets, cat D09100310) for 3 weeks. Food intake and body weight were monitored. Mice were sacrificed with CO<sub>2</sub>, and double-killed with cervical dislocation.

**For CDAHFD-induced NASH model studies**, 8 week old male MCT1<sup>f/f</sup> mice (n=10) were intravenously injected with 2X10<sup>11</sup>gc of AAV-TBG-Cre or AAV-Lrat-Cre or both. As a control, the same amount of AAV-TBG-null or AAV-Lrat-null control were used. A week after the injection, mice were fed a CDHFD (Research diets, cat A06071302i) for 8 weeks and sacrificed.

### **Primary mouse cell isolation**

Male C57BL/6 wild-type mice 16-18 week old (n=4) were subcutaneously injected with 10mg/kg of siRNAs accordingly (Chol-NTC-siRNA, Chol-MCT1-siRNA, GN-NTC-siRNA, and GN-MCT1-siRNA), twice within 15 days. Mice were put on a chow diet (LabDiet, cat 5P76) and sacrificed on day 15. Primary hepatocytes, hepatic stellate cells, and Kupffer cells were isolated from the livers using the modified perfusion method previously described<sup>60,61</sup>. Briefly, livers were digested in situ with 14mg pronase (Sigma-Aldrich, cat P5147) and 3.7U collagenase D (Roche, cat 11 088 882 001) via inferior vena cava. Digested livers were isolated and minced with 0.5mg/ml pronase, 0.088U/ml collagenase, and 0.02mg/ml DNase I (Roche, cat 10 104 159 001). After centrifuging cells for 3min at 50g at 4C, primary hepatocytes were obtained in the pellet. The remaining supernatant was collected and centrifuged for 10 min at 580g at 4C and the pellet was saved for further hepatic stellate cell separation using Nycodenz (Accurate Chemical, cat 1002424)

gradient solution. Lastly, Kupffer cells were isolated from the remaining cells using a Percoll (Sigma, cat P1644) gradient solution. Separation of hepatocytes, hepatic stellate cells, and Kupffer cells was validated using representative mRNA markers of each cell type such as *Alb*, *Des*, and *Clec4f*, respectively, by rt-qPCR.

### **Serum analysis**

Retro-orbital bleeding was performed prior to sacrificing mice. Blood was collected in heparinized capillary tubes and centrifuged 10min at 7000 rpm at 4C. Supernatant plasma was saved for further serum analysis. Plasma lactate level was measured using a specific apparatus, Lactate Plus meter (Nova biomedical, cat 62624). ALT level was determined using ALT Colorimetric Activity Assay Kit (Cayman, cat 700260). Absorbances were detected using a Tecan safire2 microplate reader.

### **Glucose tolerance test (GTT)**

GTT was performed after 16 hours of fasting. Basal glucose level was measured using a glucometer (Contour NEXT ONE glucose Meter), then mice were Intraperitoneally injected with 1g/kg body weight D-glucose dissolved in sterile saline. Blood glucose was measured with a single drop of tail blood at 15, 30, 45, 90, and 120 minutes after the glucose injection.

### **Shear wave elastography (SWE)**

Mouse liver stiffness was monitored by Vega robotic ultrasound imager, SonoEQ 1.14.0 (SonoVol), as described in a previous study<sup>43</sup>. Before SWE measurement, mice had their abdomen hair shaved and the residual hair was removed using chemical depilation cream (Nair). After being anesthetized with isoflurane, mice were located in prone position on

the fluid chamber through an acoustically transmissive membrane with ultrasound transducer imaging from below. During the imaging, wide-field B-mode was captured, a 3D volume was reconstructed, liver was visualized, and fiducial markers in 3D space indicating the position of the desired SWE capture were placed. Liver stiffness was monitored by Young's modulus.

### **RNA isolation and rt-qPCR**

Frozen mouse livers samples (25mg) or *in vitro* cell samples were homogenized in Trizol (Ambion) using Qiagen TissueLyser II. Chloroform was added and centrifuged for 15 min at maximum speed at 4C. The supernatant was collected and 100% isopropanol was added. After another 10 min centrifugation at maximum speed at 4C, the pellet was saved and washed with 70% ethanol with 5 min centrifugation at maximum speed at 4C. The pellet was dried briefly and resuspended with ultrapure distilled water (Invitrogen). cDNA was synthesized using 1ug of total RNA using iScript cDNA synthesis Kit (Biorad) on Bio-Rad T100 thermocycler. Real-time quantitative PCR was performed using iQ SybrGreen Supermix on CFX96 1000 thermocycler (Biorad) and analyzed as described<sup>62</sup>. Primer sequences used for rt-qPCR were listed in Table 3.

### **Immunoblotting**

Frozen mouse livers samples (50mg) or *in vitro* cell samples were homogenized in a sucrose lysis buffer (250mM sucrose, 50mM Tris-Cl pH7.4) with 1:100 phosphatase and protease inhibitor cocktail (Sigma-Aldrich) using Qiagen TissueLyser II. Protein concentration was determined by BCA assay. Immunoblotting loading samples were prepared after adjusting the protein concentration using 5X SDS (Sigma-Aldrich) and

denatured by boiling. Proteins were separated in 4–15% sodium dodecyl sulfate/polyacrylamide gel electrophoresis gel (Bio-Rad) and transferred to nitrocellulose membranes. The membranes were blocked with Tris-buffered saline with Tween (TBST) with 5% skim milk or 5% bovine serum albumin. Membranes were incubated with primary antibodies overnight at 4C, washed in TBST for 30 mins, then incubated with secondary antibodies for an hour at room temperature, and washed for 30 min in TBST. Antibodies used in the studies were listed in Table 4. ECL (Perkin Elmer) was added to the membranes and the protein signals were visualized with ChemiDox XRS+ image-forming system.

### **Immunofluorescence**

FL83B cells were seeded in coverslips placed in 12 well plates (150k cells/well). Then, 1.5uM of either Chol-NTC-siRNA or Chol-MCT1-siRNA at final concentration was added for 72 hours. To stain mitochondria membranes, cells were incubated with Mitotracker at 37C (Thermofisher, cat M7512) for 45 min in serum-free media. Then, cells were fixed with 4% paraformaldehyde at room temperature for 30 min. Fixed cells were blocked by fresh permeabilization buffer (0.5% Triton, 1% FBS in PBS) at room temperature for 30 min and incubated with 1:100 Anti-MCT1 (Proteintech, cat 20139-1-AP) overnight at 4C. As a secondary antibody, 1:1000 goat-anti-Rabbit-488 was used, while cells were protected from light. Coverslips were mounted on Prolong Gold antifade Mountant with Dapi (Invitrogen, cat P35934). Images were acquired using an Olympus IX81 microscope (Central Valley, PA) with dual Andor Zyla sCMOS 4.2 cameras (Belfast, UK). Images were quantified using ImageJ software.

## Histological analysis

For the immunohistochemistry (IHC), half of the biggest lobe of each mouse liver was fixed in 4% paraformaldehyde and embedded in paraffin. Sectioned slides were stained where indicated with H&E, Trichrome, Sirius Red, and anti-MCT1 (Proteintech, cat 20139-1-AP) at the University of Massachusetts Chan Medical School Morphology Core. The whole stained slides were scanned with ZEISS Axio Scan Z1. Images were analyzed by ZEN 3.0 and ImageJ software.

## H&E lipid droplet analysis

To quantify the mean size and mean number of lipid droplets, H&E images further underwent thorough image analysis. The 2D RGB images (8 bits per channel) were read into the Fiji version<sup>63</sup> based on ImageJ2<sup>64</sup>. An ImageJ macro language program was then written to analyze each image. First, the Labkit plugin<sup>65</sup> was used to classify pixels as either lipid or background. The classifier used was trained on a few short line segments drawn in either lipid or background regions. The binary objects created then had their holes filled and were culled using "Analyze Particles" to eliminate objects larger than 10,000 pixels (typically veins). Then the "watershed" algorithm was used to separate touching lipid droplets, and "Analyze Particles" was used again, this time to keep objects with a circularity of 0.5-1 and a size of 40-5000 pixels. Pixel size was converted to um using a 0.47um/pixel width conversion ratio (so a 0.22 um<sup>2</sup>/pixel conversion factor).

## Trichrome and Sirius Red Image analysis for fibrosis

To quantify the % of fibrotic regions, 2D RGB images (8 bits per channel) of Sirius Red and Trichrome (without the hematoxylin stain) were read into the Fiji version<sup>63</sup> based on

ImageJ2<sup>64</sup>. “Analyze Particles” was used to threshold the Sirius Red images (20X magnification) in the green channel, keeping pixels with an intensity < 100 and object size > 100 pixels as fibrotic regions. Pixels in the Trichrome images (2.5X magnification) in the red channel with intensity < 60 and object size > 0.0005 pixels were considered as fibrotic regions.

### **Quantification and statistical analysis**

All statistical analyses were calculated using GraphPad Prism 9 (GraphPad Software). A two-sided unpaired Student t-test was used for the analysis of the statistical significance between the two groups. For more than three groups, One-way ANOVA was used for the analysis of statistical significance. Data were presented as mean ± SD or otherwise noted. Differences were considered significant when  $p < 0.05$  (\*:  $p < 0.05$ , \*\*:  $p < 0.005$ , and \*\*\*:  $p < 0.0005$ ). Data were excluded only when a technical error occurred in sample preparation. Sample sizes were decided based on previous publications.

## Author contributions

K.M., and M.P.C. designed the study, analyzed the data, and wrote the manuscript. K.M., B.Y., M.K., M.E., N.R., E.T., S.M., and C.D. performed most of the experiments and analyzed the data. D.E., N.M., B.B., A.K. contributed to the synthesis of the oligonucleotides and provided guidance in oligonucleotide technology. L.L. performed lipid droplet image analysis. B.M., J.R. generously provided MCT1<sup>f/f</sup> mice.

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## Tables

**Table 1.**

Antisense strands:

Oligo ID	Chemically modified RNA sequence
MCT1-507	P(mU)#{(fG)#{(mU)(fU)(mA)(fC)(mA)(fG)(mA)(fA)(mA)(fG)(mA)#{(fA)}#(mG)#{(fC)#{(mU)#{(fG)#{(mC)#{(fG)}}
MCT1-1976	P(mU)#{(fA)#{(mA)(fA)(mC)(fU)(mU)(fA)(mA)(fG)(mG)(fC)(mA)#{(fC)}#(mA)#{(fU)#{(mA)#{(fU)#{(mU)#{(fA)}}
MCT1-2013	P(mU)#{(fU)#{(mU)(fA)(mA)(fA)(mA)(fG)(mU)(fU)(mA)(fA)(mG)#{(fG)}#(mC)#{(fU)#{(mC)#{(fU)#{(mC)#{(fU)}}
MCT1-2042	P(mU)#{(fU)#{(mU)(fA)(mA)(fA)(mA)(fC)(mA)(fA)(mA)(fU)(mG)#{(fA)}#(mA)#{(fU)#{(mU)#{(fC)#{(mA)#{(fG)}}
MCT1-2060	P(mU)#{(fU)#{(mU)(fC)(mC)(fU)(mU)(fU)(mU)(fA)(mA)(fA)(mA)#{(fU)}#(mG)#{(fA)#{(mC)#{(fA)#{(mU)#{(fU)}}
MCT1-2120	P(mU)#{(fU)#{(mU)(fA)(mC)(fA)(mA)(fA)(mC)(fA)(mA)(fC)(mA)#{(fA)}#(mC)#{(fA)#{(mA)#{(fA)#{(mA)#{(fC)}}
MCT1-3067	P(mU)#{(fU)#{(mU)(fU)(mC)(fU)(mG)(fC)(mC)(fU)(mC)(fU)(mA)#{(fU)}#(mU)#{(fC)#{(mA)#{(fG)#{(mA)#{(fA)}}
MCT1-3160	P(mU)#{(fU)#{(mC)(fU)(mU)(fA)(mC)(fA)(mC)(fA)(mA)(fG)(mG)#{(fU)}#(mU)#{(fU)#{(mU)#{(fA)#{(mA)#{(fA)}}
MCT1-3290	P(mU)#{(fA)#{(mU)(fA)(mU)(fU)(mA)(fG)(mA)(fA)(mA)(fG)(mG)#{(fU)}#(mU)#{(fA)#{(mA)#{(fA)#{(mA)#{(fU)}}
MCT1-4340	P(mU)#{(fU)#{(mG)(fA)(mA)(fU)(mU)(fU)(mG)(fU)(mA)(fU)(mG)#{(fA)}#(mG)#{(fA)#{(mA)#{(fU)#{(mA)#{(fA)}}

Sense strands:

Oligo ID	Chemically modified RNA sequence
Chol-MCT1-507	(fC)#{(mU)#{(fU)(mC)(fU)(mU)(fU)(mC)(fU)(mG)(fU)(mA)(fA)#{(mC)}#(fA)-Chol
Chol-MCT1-1976	(fU)#{(mG)#{(fU)(mG)(fC)(mC)(fU)(mU)(fA)(mA)(fG)(mU)(fU)#{(mU)}#(fA)-Chol
Chol-MCT1-2013	(fG)#{(mC)#{(fC)(mU)(fU)(mA)(fA)(mC)(fU)(mU)(fU)(mU)(fA)#{(mA)}#(fA)-Chol
Chol-MCT1-2042	(fU)#{(mU)#{(fC)(mA)(fU)(mU)(fU)(mG)(fU)(mU)(fU)(mU)(fA)#{(mA)}#(fA)-Chol
Chol-MCT1-2060	(fC)#{(mA)#{(fU)(mU)(fU)(mU)(fA)(mA)(fA)(mA)(fG)(mG)(fA)#{(mA)}#(fA)-Chol
Chol-MCT1-2120	(fG)#{(mU)#{(fU)(mG)(fU)(mU)(fG)(mU)(fU)(mU)(fG)(mU)(fA)#{(mA)}#(fA)-Chol
Chol-MCT1-3067	(fA)#{(mA)#{(fU)(mA)(fG)(mA)(fG)(mG)(fC)(mA)(fG)(mA)(fA)#{(mA)}#(fA)-Chol
Chol-MCT1-3160	(fA)#{(mA)#{(fC)(mC)(fU)(mU)(fG)(mU)(fG)(mU)(fA)(mA)(fG)#{(mA)}#(fA)-Chol

Chol-MCT1-3290	(fA)#{(mA)#{(fC)(mC)(fU)(mU)(fU)(mC)(fU)(mA)(fA)(mU)(fA)}#(mU) #(fA)-Chol
Chol-MCT1-4340	(fC)#{(mU)#{(fC)(mA)(fU)(mA)(fC)(mA)(fA)(mA)(fU)(mU)(fC)}#(mA) #(fA)-Chol

**Table 1. Sequences of chemically modified siRNA candidates targeting MCT1 used in *in vitro* screening.** siRNAs utilized in *in vitro* screening were a double-strand oligonucleotide comprised of 15 sense and 20 antisense nucleotides. The sequences of each candidate's antisense and sense strands were listed. (P: 5'-Phosphate, #: phosphorothioate, m:2'-O- methyl, f: 2'-fluoro, Chol: tetra-ethylenglycol-cholesterol conjugate)

**Table 2**

Antisense strands:

Oligo ID	Chemically modified RNA sequence
MCT1-3160	VP(mU)#{fU}#{mC}(mU)(mU)(fA)(mC)(mA)(mC)(mA)(mA)(mG)(mG) #{fU}#{mU}#{fU}#{mU}#{mA}#{mA}#{fA}

Sense strands:

Oligo ID	Chemically modified RNA sequence
Chol-MCT1-3160	(mU)#{mA}#{mA}(mA)(mA)(mC)(mC)(fU)(fU)(fG)(mU)(fG)(mU) (mA)(mA)(mG)#{mA}#{mA}-Chol
GN-MCT-3160	(mU)#{mA}#{mA}(mA)(mA)(mC)(mC)(fU)(fU)(fG)(mU)(fG)(mU) (mA)(mA)(mG)#{mA}#{mA}-GN

**Table 2. Sequences of the selected final chemically modified siRNA candidates targeting MCT1 used for *in vivo* studies.** MCT1-3160 was selected for the final construct for *in vivo* studies. MCT1-siRNAs utilized in *in vivo* study was a double-strand oligonucleotide comprised of 18 sense and 20 antisense nucleotides. To sense strands, either Chol- or GN- was attached. (VP: 5’-(E)-vinyl phosphonate, #: phosphorothioate, m:2’-O- methyl, f: 2’-fluoro, Chol: tetra-ethylenglycol-cholesterol conjugate, GN: tri-N-Acetyl-galactosamine)

**Table 3**

Mouse primers:

Gene	Forward	Reverse
<i>Mct1</i>	TGTTAGTCGGAGCCTTCATTTC	CACTGGTCGTTGCAC TGAAATA
<i>Mct1</i> (Exon 2,3 overlapping)	TGCAACGACCAGTGAAGTATC	GCTGCCGTATTATTACCAAG
<i>Srebf1</i>	GGAGCCATGGATTGCACATT	GGCCC GGAAAGTC ACTGT
<i>Mlxip1</i>	TCTGCAGATCGCGTGGAG	CTTGTCCC GGCA TAGCAAC
<i>Fasn</i>	GGAGGTGGTGATAGCCGGTAT	TGGGTAATCCATAGAGCCCAG
<i>Scd1</i>	CCGGAGACCCCTTAGATCGA	TAGCCTGTAAAAGATTCTGCAAACC
<i>Acly</i>	TGGT GGAATGCTGGACAA	GCCCTCATAGACACCATCTG
<i>Srebf2</i>	GCGTTCTGGAGACCATGGA	ACAAAGTTGCTCTGAAAACAATCA
<i>Hmgcr</i>	CTTGT GGAATGCCTTGATTG	AGCCGAAGCAGCACATGAT
<i>Hmgcs2</i>	GCCGTGAACTGGGTCGAA	GCATATATAGCAATGTCTCCTGCAA
<i>Tgf1β</i>	CTCCC GTGGCTCTAGTGC	GCCTTAGTTGGACAGGATCTG
<i>Ihh</i>	CTCTTG C CTACAAGCAGTTCA	CCGTGTTCTCCTCGTCCTT
<i>Acta2</i>	ATGCTCCCAGGGCTGTTTC	GTGGTGCCAGATCTTTCCATGTCG
<i>Gli2</i>	CAACGCCTACTCTCCCAGAC	GAGCCTTGATGTACTGTACAC
<i>Gli3</i>	CACAGCTCTACGGCGACTG	CTGCATAGTGATTGC GTTCTTC
<i>Col1a1</i>	GCTCCTCTAGGGGCCACT	CCACGTCTCACCATTGGGG
<i>Col1a2</i>	GTAACTTCGTGCCTAGCAACA	CCTTGTCAGAACATACTGAGCAGC
<i>Col3a1</i>	CTGTAACATGGAAACTGGGGAAA	CCATAGCTGA ACTGAAAACCACC
<i>Timp1</i>	CTCAAAGACCTATAGTGCTGGC	CAAAGTGACGGCTCTGGTAG
<i>Alb</i>	TGCTTTCCAGGGGTGTGTT	TTACTT CCTGC ACTAATTGGCA
<i>Des</i>	CTAAAGGATGAGATGGCCCG	GAAGGTCTGGATAGGAAGGTTG
<i>Clec4f</i>	GAGGCCGAGCTGAACAGAG	TGTGAAGCCACCACAAAAAGAG
<i>B2m</i>	CATGGCTCGCTCGGTGAC	CAGTCAGTATGTT CGGCTTCC

Human primers:

Gene	Forward	Reverse
<i>MCT1</i>	TGGAAGACACCCTAAACAAGAG	AAAGCCTCTGTGGGTGAATAG
<i>ACTA2</i>	AGCGTGGCTATTCCCTCGT	CTCATTTC CAAAGTCCAGAGCTACA
<i>TGF1β</i>	CAACGAAATCTATGACAAGTTCAA GCAG	CTTCTCGGAGCTCTGATGTG
<i>COL1A1</i>	ACGT CCTGGTGAAGTTGGTC	ACCAGGGAAAGCCTCTCTCTC
<i>TIMP1</i>	AATTCCGACCTCGTCATCAGG	ATCCCCTAAGGCTTGGAACCC
<i>ACTB</i>	GATGAGATTGGCATGGCTTT	GAGAAGTGGGTGGCTT

**Table 3. List of primers used for rt-qPCR**

**Table 4**

Reagent	Source	Identifier
Anti-MCT1	Proteintech	Cat # 20139-1-AP
Anti-FASN	Cell Signaling	Cat # 3180s
Anti-ACLY	Cell Signaling	Cat # 4332
Anti-SCD1	Cell Signaling	Cat # 2794s
Anti-ChREBP	Novus Bio	Cat # NB400-135
Anti-SREBP1	Millipore	Cat # MABS1987
Anti-GAPDH-HRP	Cell Signaling	Cat # 8884s
Anti-H3	Cell Signaling	Cat # 4499s
Anti-Tubulin	Sigma Aldrich	Cat # T5168
Anti-pAMPK (T172)	Cell Signaling	Cat # 2535s
Anti-AMPK $\alpha$	Cell Signaling	Cat # 2793s
Anti- $\alpha$ SMA	Cell Signaling	Cat # 19245s
Anti-Collagen 1	Southern Biotech	Cat # 1310-01
Anti-HSP90-HRP	Cell Signaling	Cat # 79631s
Goat Anti-Rabbit IgG-HRP	Invitrogen	Cat # 65-6120
Goat Anti-Mouse IgG-HRP	Invitrogen	Cat # 65-6520
Goat Anti-Mouse IgG-HRP	Thermofisher	Cat # G21040
Mouse Anti-Goat IgG-HRP	Santa Cruz	Cat # sc-2354
Goat-anti-Rabbit-488	Thermofisher	Cat # A11008
Prolong-gold Antifade Mountant	Thermofisher	Cat # P36931

**Table 4. List of antibodies used in this study**

**Figure 1. MCT1 depletion attenuates Tgf1 $\beta$ -stimulated collagen 1 production in human hepatic stellate cell lines, LX2.** Cells were transfected with either NTC-siRNA or MCT1-siRNA for 6 hours. Then, Tgf1 $\beta$  was treated to induce collagen production. 48 hours after the Tgf1 $\beta$  treatment, cells were harvested and processed for rt-qPCR or Western blotting. **(A)** MCT1 mRNA expression levels. **(B)** Collagen 1 protein levels. Quantification was added below. **(C)** Representative fibrogenic marker genes, ACTA2, and COL1A1 expression levels were monitored. (t-test, One way ANOVA, \*: p<0.05, \*\*: p<0.01, \*\*\*: p<0.001, \*\*\*\*: p<0.0001)

**Figure 2. Screening of chemically modified Chol-MCT1-siRNA *in vitro*.** **(A)** Targeted regions of multiple Chol-MCT1-siRNA candidates on *Mct1* transcript. **(B)** Silencing efficacy of each Chol-MCT1-siRNA candidate (1.5uM) on *Mct1* mRNA expression levels was monitored 72 hours after the transfection in mouse hepatocyte cell lines, FL83B *in vitro*. Chol-NTC-siRNA was used as a control. **(C)** Dose-response potency test was performed to identify the most potent Chol-MCT1-siRNA compound. IC50 values were determined using six serially diluted concentrations of each compound starting from 1.5uM. IC50 values and knockdown % of the two most potent compounds were shown in the table below. **(D)** 72 hours after the transfection of Chol-MCT1-2060 compounds (1.5uM), MCT1 protein expression levels were visually monitored by immunofluorescence. **(E)** 72 hours after the transfection of either Chol-MCT1-2060 or Chol-MCT1-3160 compounds (1.5uM), their silencing efficacy on MCT1 protein expression levels was examined by Western blotting.

**Figure 3. Biodistribution of Chol- and GN-MCT1-siRNA in the liver.** Male C57BL/6 wild-type mice (16-18 wk, n=4) were subcutaneously injected with 10mg/kg of each siRNA, twice within 15 days, while fed a chow diet. Mice were sacrificed on day 15. **(A)** Chemical structure of the fully chemically modified siRNA that was used for further *in vivo* studies; Chol-MCT1-siRNA and GN-MCT1-siRNA. **(B, E)** Primary hepatocytes, **(C, F)** stellate cells, and **(D, G)** Kupffer cells were isolated from each mouse using different gravity centrifugations and gradient solutions after the liver perfusion. *Mct1* mRNA expression levels in each cell type fraction were measured. (t-test, \*: p<0.05, \*\*: p<0.01)

**Figure 4. Hepatic MCT1 depletion did not resolve steatosis in a genetically obese NASH mouse model.** **(A)** Male ob/ob mice (10 wk, n=6) were subcutaneously injected with 10mg/kg of siRNA once every 10 days. Mice were fed a GAN diet for 3 weeks and sacrificed. **(B)** Livers were stained with MCT1 antibody and the representative images of each group are shown. **(C)** % of MCT1 positive area shown in immunohistochemistry images were quantified. **(D)** Hepatic *Mct1* mRNA level was measured by rt-qPCR upon each siRNA administration. **(E)** Plasma lactate levels were monitored. **(F)** Mean size of lipid droplets was quantified from H&E images (mean, sem). **(G)** Mean number of lipid droplets was quantified from H&E images (mean, sem). **(H)** Liver TG levels were examined in each group.

**Figure 5. Opposite effects of Chol-MCT1-siRNA versus GN-MCT1-siRNA on fibrotic type 1 collagen expression.** Representative fibrogenic gene expression levels were measured for **(A, B)** mRNA and **(C, D)** protein. **(E, F)** Protein expression levels were quantified. **(G)** Livers were stained with Sirius Red and the representative images of each

group are shown. **(H)** % of Sirius Red positive areas were quantified. (t-test, \*: p<0.05, \*\*: p<0.01, \*\*\*: p<0.001)

**Figure 6. MCT1 depletion did not resolve steatosis in the CDHFD-induced NASH model. (A)** Male MCT1<sup>f/f</sup> mice (8 wk, n=10) were intravenously injected with 2X10<sup>11</sup>gc of AAV-TBG-Cre or AAV-Lrat-Cre or both. The same amount of AAV-TBG-null or AAV-Lrat-null was used as a control. A week after the injection, mice were fed a CDHFD for 8 weeks and sacrificed. **(B)** *Mct1* mRNA expression levels in whole livers were examined. **(C)** Food intake and **(D)** body weights were monitored. **(E)** CDHFD-induced steatosis was monitored by H&E. **(F)** % of lipid droplet areas was quantified. (t-test, One way ANOVA, \*: p<0.05, \*\*: p<0.01, \*\*\*: p<0.001, \*\*\*\*: p<0.0001)

**Figure 7. Hepatocyte-specific MCT1KO accelerated fibrosis, while hepatic stellate cell-specific MCT1KO decreased it.** Male MCT1<sup>f/f</sup> mice (6 wk, n=10) were intravenously injected with 2X10<sup>11</sup>gc of AAV-TBG-Cre or AAV-Lrat-Cre or both. The same amount of AAV-TBG-null or AAV-Lrat-null was used as a control. A week after the injection, mice were fed a CDHFD for 8 weeks and sacrificed. **(A)** Collagen 1 protein levels were compared between the control and the hepatocyte MCT1KO groups. **(B)** Collagen 1 protein levels were compared between the control and the hepatic stellate cell MCT1KO groups. **(C)** Collagen 1 protein levels were compared between the control group and MCT1KO in both hepatocyte and hepatic stellate cell groups. **(D)** Livers were stained with Trichrome and the representative images of each group were shown. **(E)** Trichrome staining images were quantified. **(F)** Liver stiffness was monitored 4 weeks after CDHFD feeding via SWE. **(G)** Liver stiffness was monitored 8 weeks after CDHFD feeding via SWE. **(H)** ALT levels were measured in every CDHFD-fed group. (t-test, One way ANOVA, \*: p<0.05, \*\*: p<0.01, \*\*\*: p<0.001, \*\*\*\*: p<0.0001)

**Supplement Figure 1. Biodistribution of GN-MCT1-siRNA and Chol-MCT1-siRNA.** Male C57BL/6 wild-type mice (16-18 wk, n=4) were subcutaneously injected with 10mg/kg of siRNAs twice within 15 days. Livers were perfused through inferior vena cava and multiple liver cells were isolated using different gravity centrifugations and gradient solutions. **(A, D)** Purity of isolated hepatocytes, **(B, E)** hepatic stellate cells, **(C, F)** and Kupffer cells was validated with representative marker genes, *Alb*, *Des*, and *Clec4f* expression, respectively. **(G, H)** MCT1 protein expression levels in multiple fat tissues (inguinal white adipose tissue, gonadal white adipose tissue, and brown adipose tissue) were monitored by immunohistochemistry. % MCT1 positive areas were quantified. **(H, J)** MCT1 protein expression levels in multiple tissues (heart, lung, kidney, spleen, and intestine) were monitored by immunohistochemistry. % MCT1 positive areas were quantified. (t-test, One way ANOVA, \*: p<0.05, \*\*: p<0.01, \*\*\*: p<0.001, \*\*\*\*: p<0.0001)

**Supplement Figure 2. GN-MCT1-siRNA decreased food intake and body weight.** Male ob/ob mice (10 wk, n=6) were subcutaneously injected with 10mg/kg of siRNA. Mice were fed a GAN diet for 3 weeks and sacrificed. **(A)** Accumulative food intake and **(B)** body weight were monitored during the study. (t-test, \*: p<0.05)

**Supplement Figure 3. Neither Chol-MCT1-siRNA administration nor hepatocyte-specific MCT1KO improved glucose tolerance.** **(A)** Male ob/ob mice (8 wk, n=4) were

subcutaneously injected with 10mg/kg of either Chol-NTC-siRNA or Chol-MCT1-siRNA once every 10 days. Mice were fed a GAN diet for 3 weeks. Then, a GTT was performed after 16 hours of fasting. **(B)** Male  $MCT1^{fl/fl}$  mice (n=6) were intravenously injected with  $1 \times 10^{11}$ gc of either AAV-TBG-Cre or AAV-Lrat-Cre. Mice were fed a high-fat diet for 12 weeks. Then, a GTT was performed after 16 hours of fasting.

**Supplement Figure 4. Both Chol-MCT1-siRNA and GN-MCT1-siRNA significantly decreased hepatic DNL gene expression.** Representative DNL gene expression levels were measured in **(A, B)** mRNA and **(C, D)** protein upon Chol-siRNA or GN-siRNA administration, respectively. Protein expression levels were quantified. **(E, F)** pAMPK and AMPK protein levels and their expression ratio were quantified. (t-test, \*: p<0.05, \*\*: p<0.01, \*\*\*: p<0.001, \*\*\*\*: p<0.0001).

**Supplement Figure 5. Intravenous injection of AAV9-Lrat-Cre in  $MCT1^{fl/fl}$  mice specifically targets hepatic stellate cells.** **(A)** Male mice (9-10wk, n=4) were intravenously injected with  $1 \times 10^{11}$ gc of either AAV9-Lrat-null or AAV9-Lrat-Cre. Mice were fed a chow diet for 3 weeks and sacrificed. Isolation of either **(B)** primary hepatocytes or **(C)** stellate cells was validated with each cell type's representative marker, albumin and desmin, respectively. *Mct1* mRNA expression levels in **(D)** hepatocyte or **(E)** stellate cell fractions were examined. **(F)** MCT1 protein expression levels in multiple fat tissues (inguinal white adipose tissue, gonadal white adipose tissue, and brown adipose tissue) were monitored by immunohistochemistry. % MCT1 positive areas were quantified. **(G)** MCT1 protein expression levels in multiple tissues (heart, lung, kidney, spleen, and intestine) were monitored by immunohistochemistry. % MCT1 positive areas were quantified. (t-test, \*: p<0.05, \*\*\*\*: p<0.0001)

**Supplement Figure 6. MCT1 silencing enhanced fibrogenic gene expression levels in human hematoma cell lines, HepG2.** Cells were transfected with either NTC-siRNA or MCT1-siRNA for 6 hours. 48 hours after, cells were harvested, and media were collected. LX2 cells were provided with the conditioned media (40% conditioned media + 60% fresh media) and harvested in 48 hours. **(A)** MCT1 mRNA expression levels were measured in HepG2 cells upon MCT1-siRNA treatment. **(B)** Fibrogenic gene expression levels were measured in HepG2 cells upon MCT1-siRNA treatment. **(C)** Fibrogenic gene expression levels were measured in LX2 cells upon conditioned media treatment. (t-test, \*: p<0.05, \*\*: p<0.01)

**Supplement figure 7. Graphical abstract.** Hepatocyte MCT1KO enhances fibrosis, while stellate cell MCT1KO decreases it.

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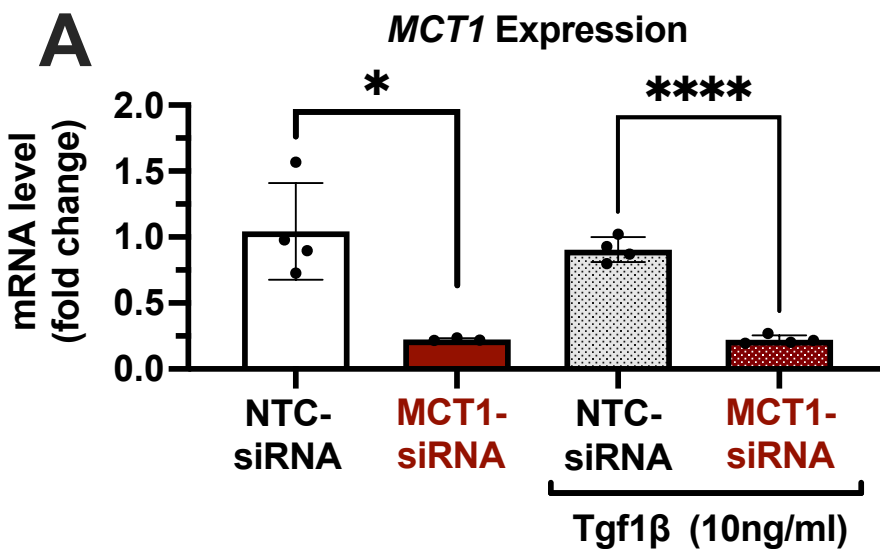
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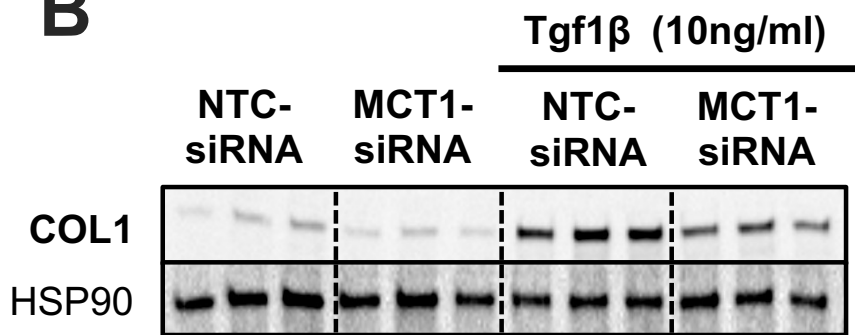
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Figure 1

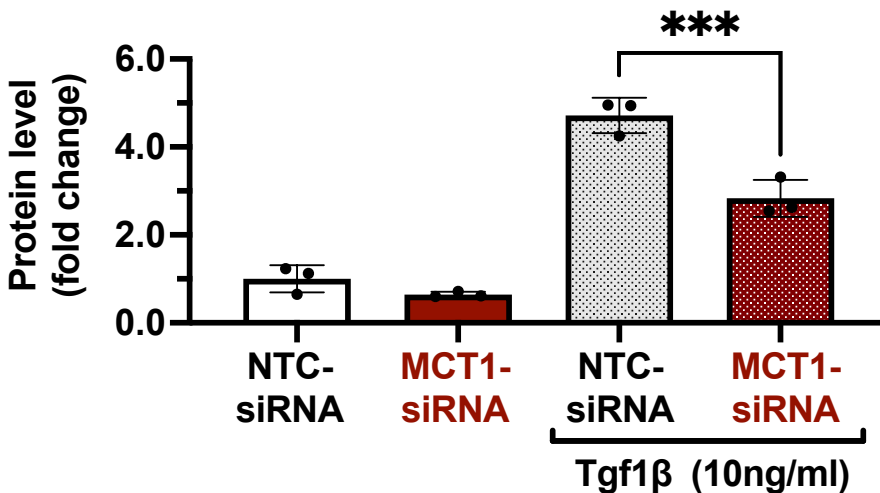
A



B



### Collagen 1 Expression



C

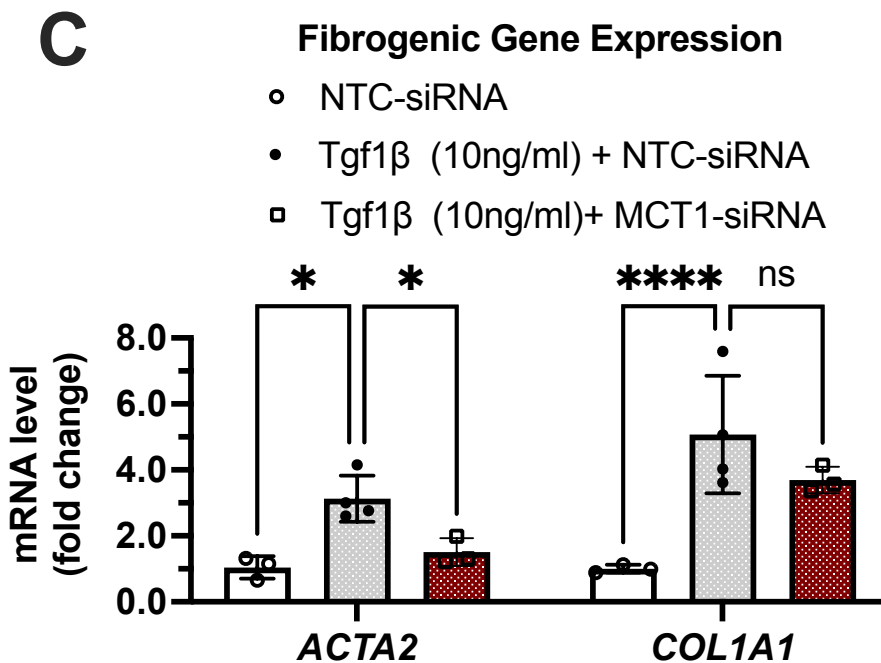
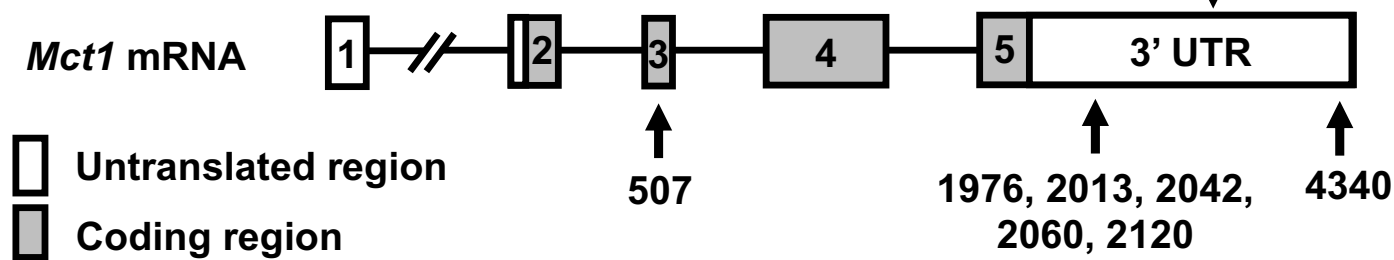
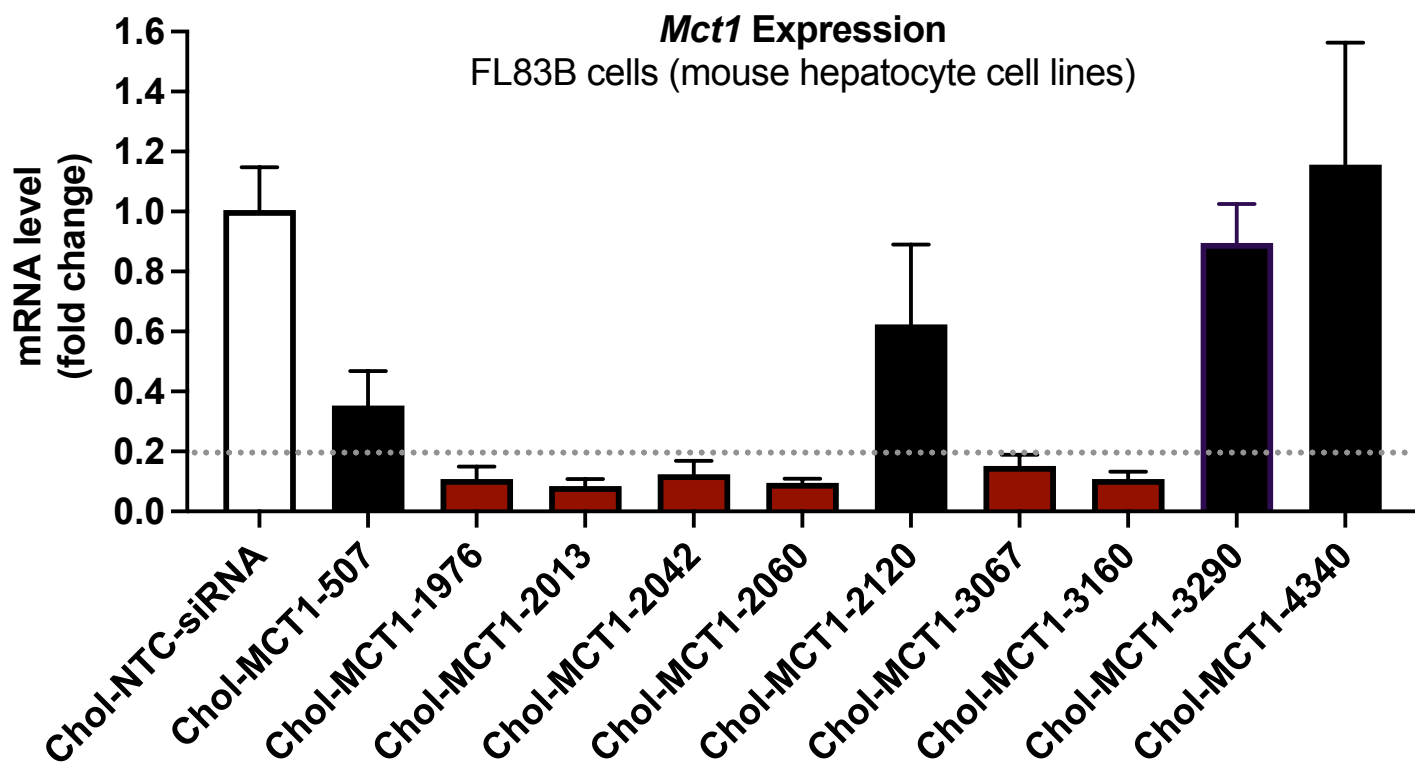
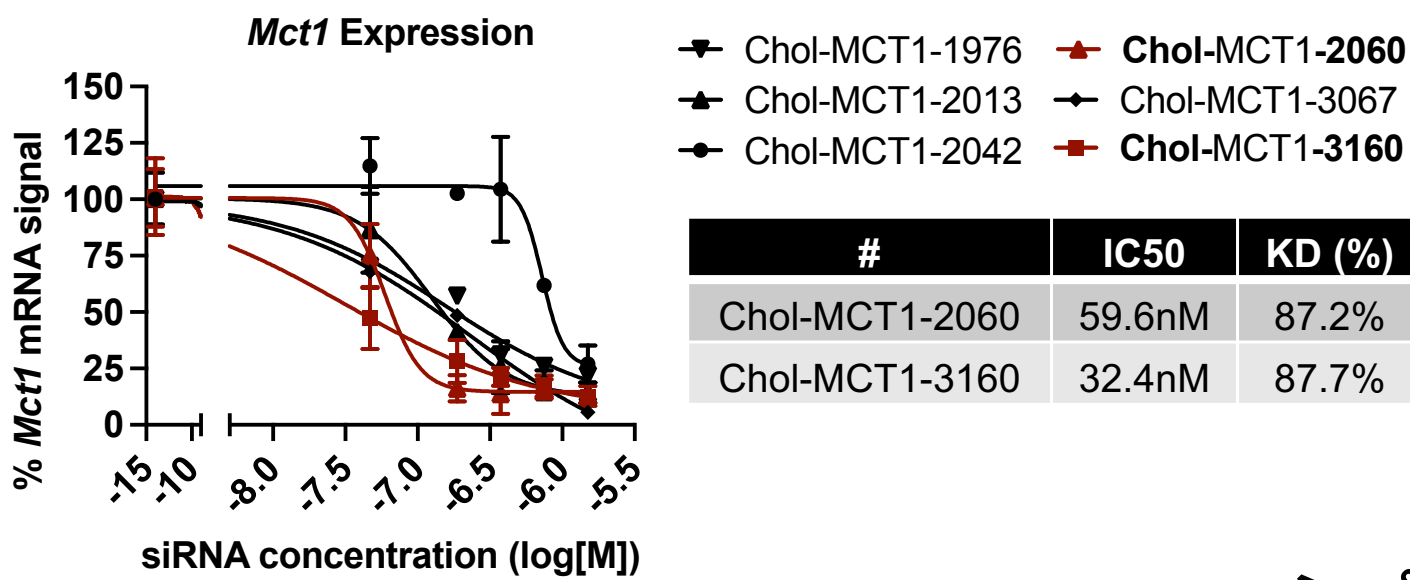
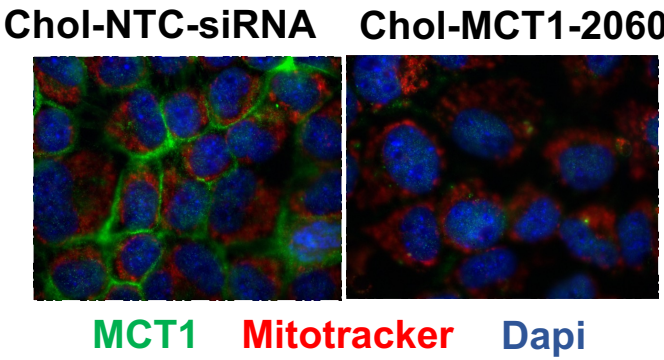
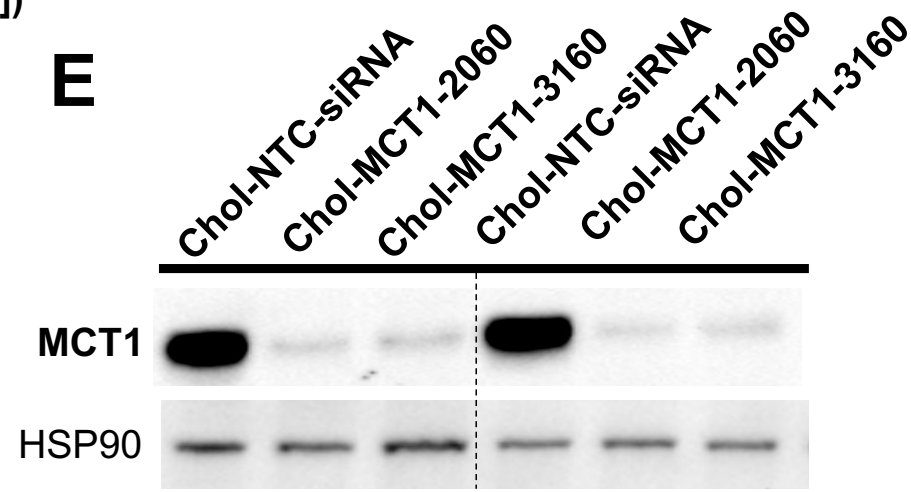


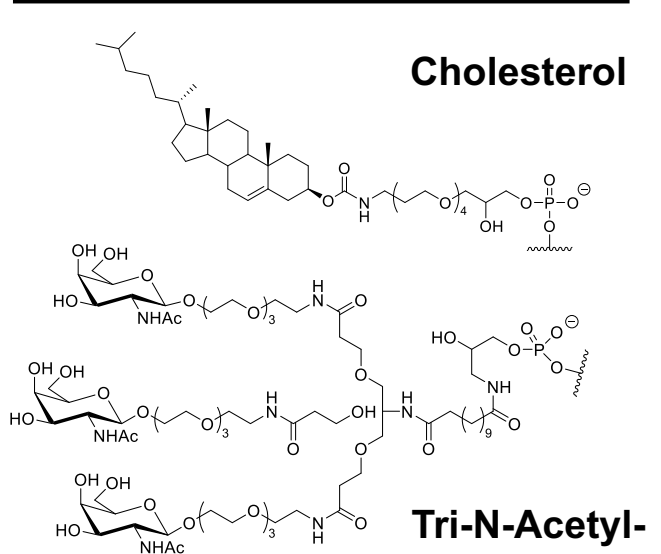
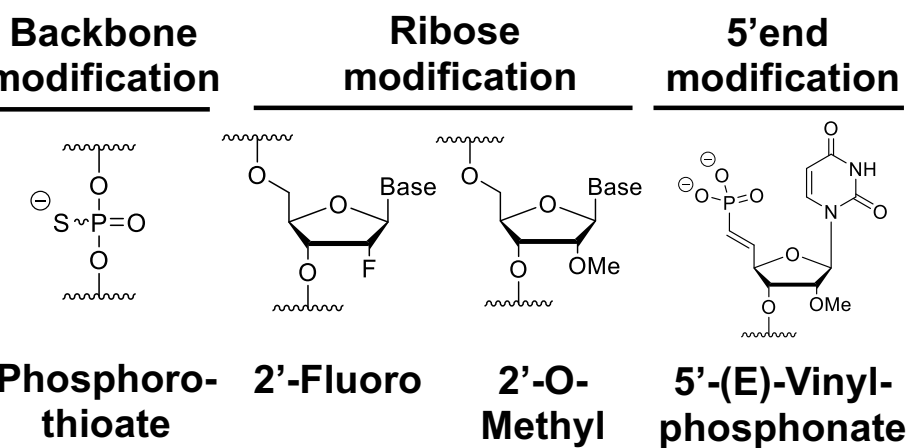
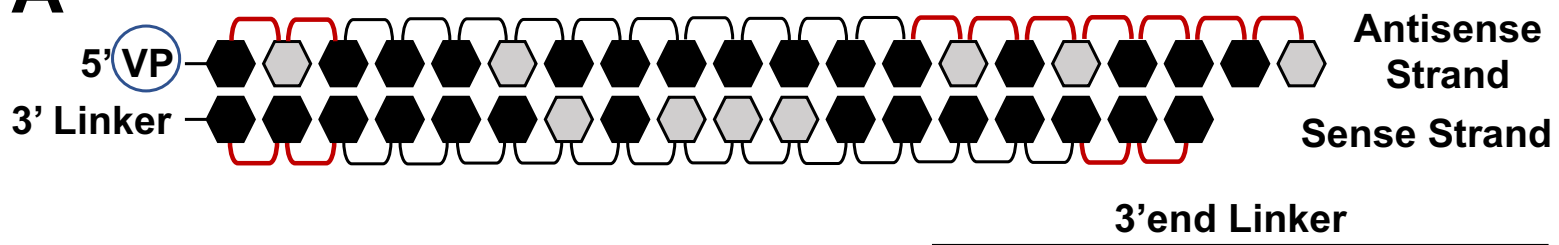
Figure 2

3067, 3160, 3290

**A****B****C****D****E**

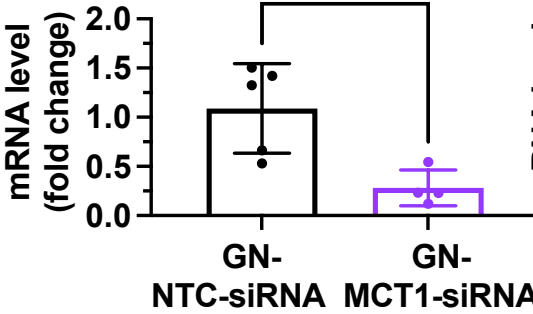
# Figure 3

**A**



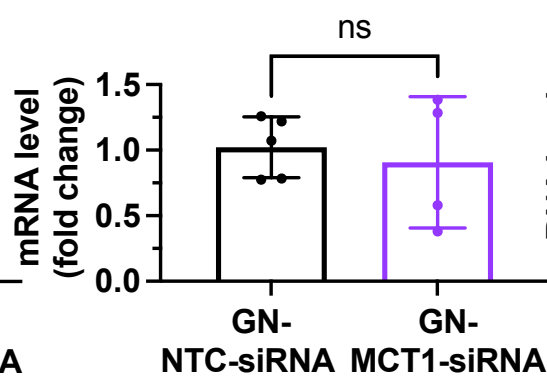
**B**

*Mct1* Expression  
(Hepatocyte fraction)



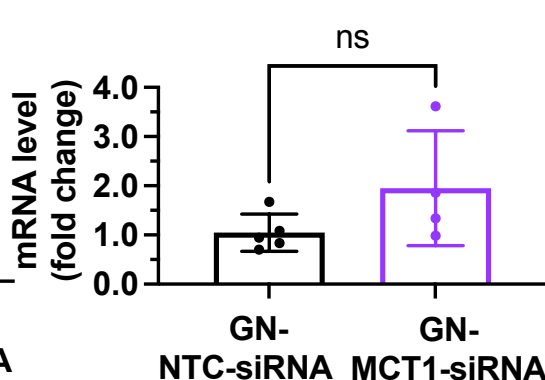
**C**

*Mct1* Expression  
(Stellate cell fraction)



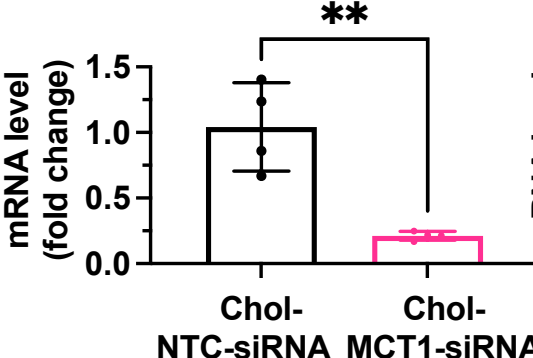
**D**

*Mct1* Expression  
(Kupffer cell fraction)



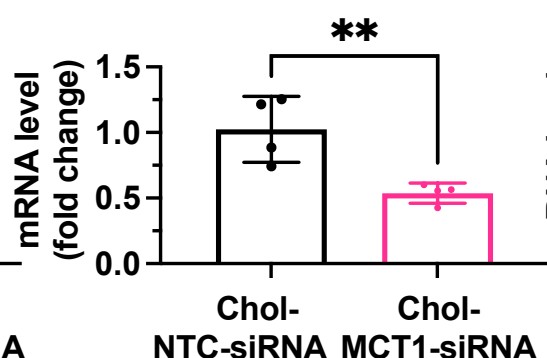
**E**

*Mct1* Expression  
(Hepatocyte fraction)



**F**

*Mct1* Expression  
(Stellate cell fraction)



**G**

*Mct1* Expression  
(Kupffer cell fraction)

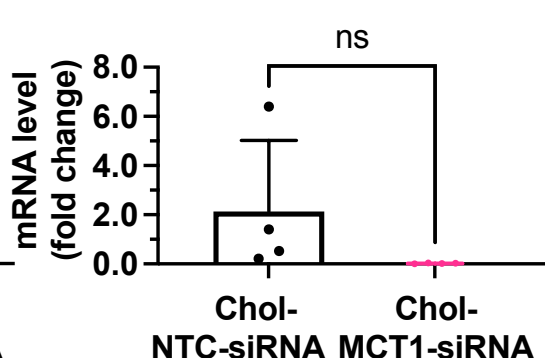


Figure 4

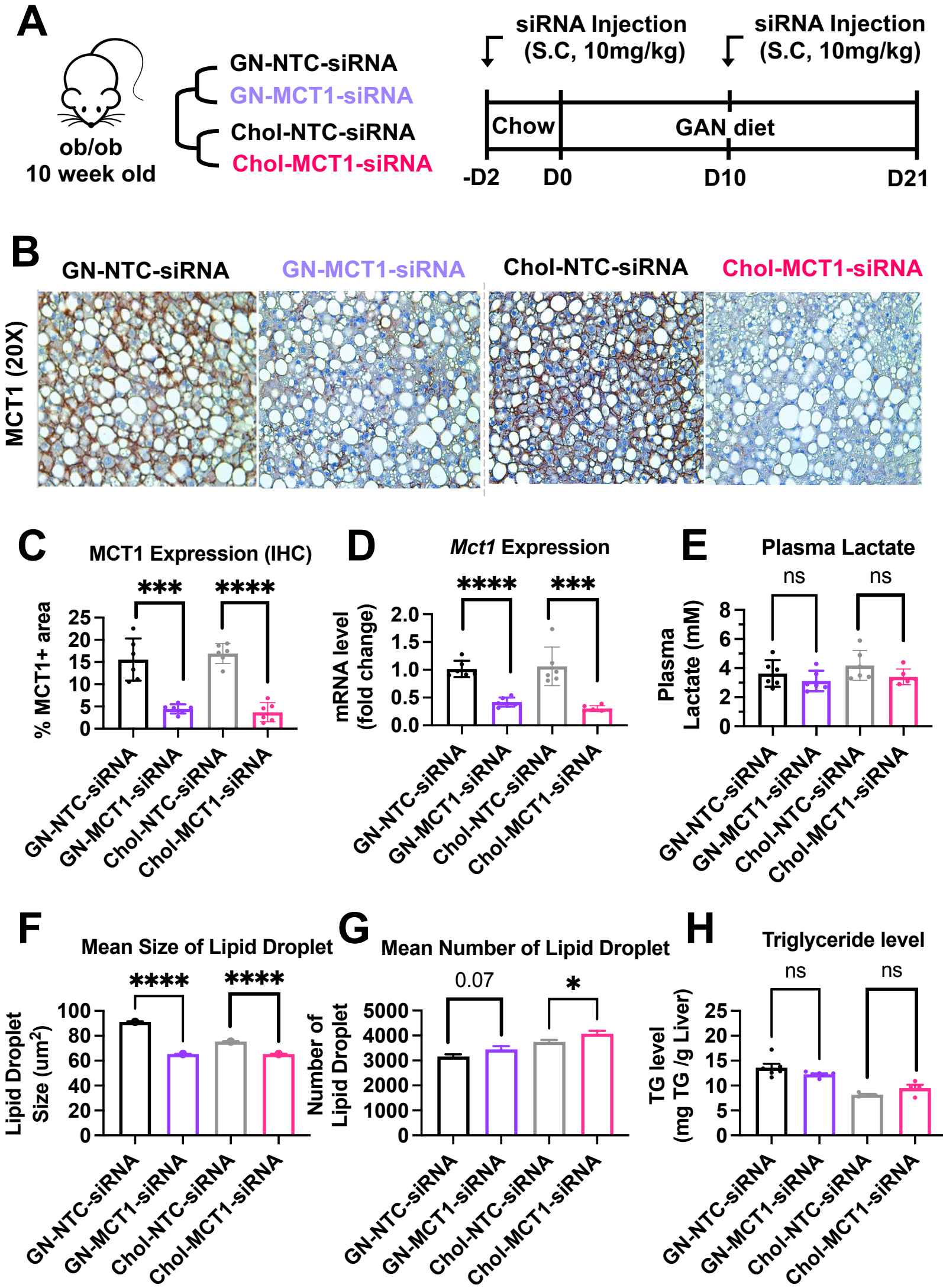
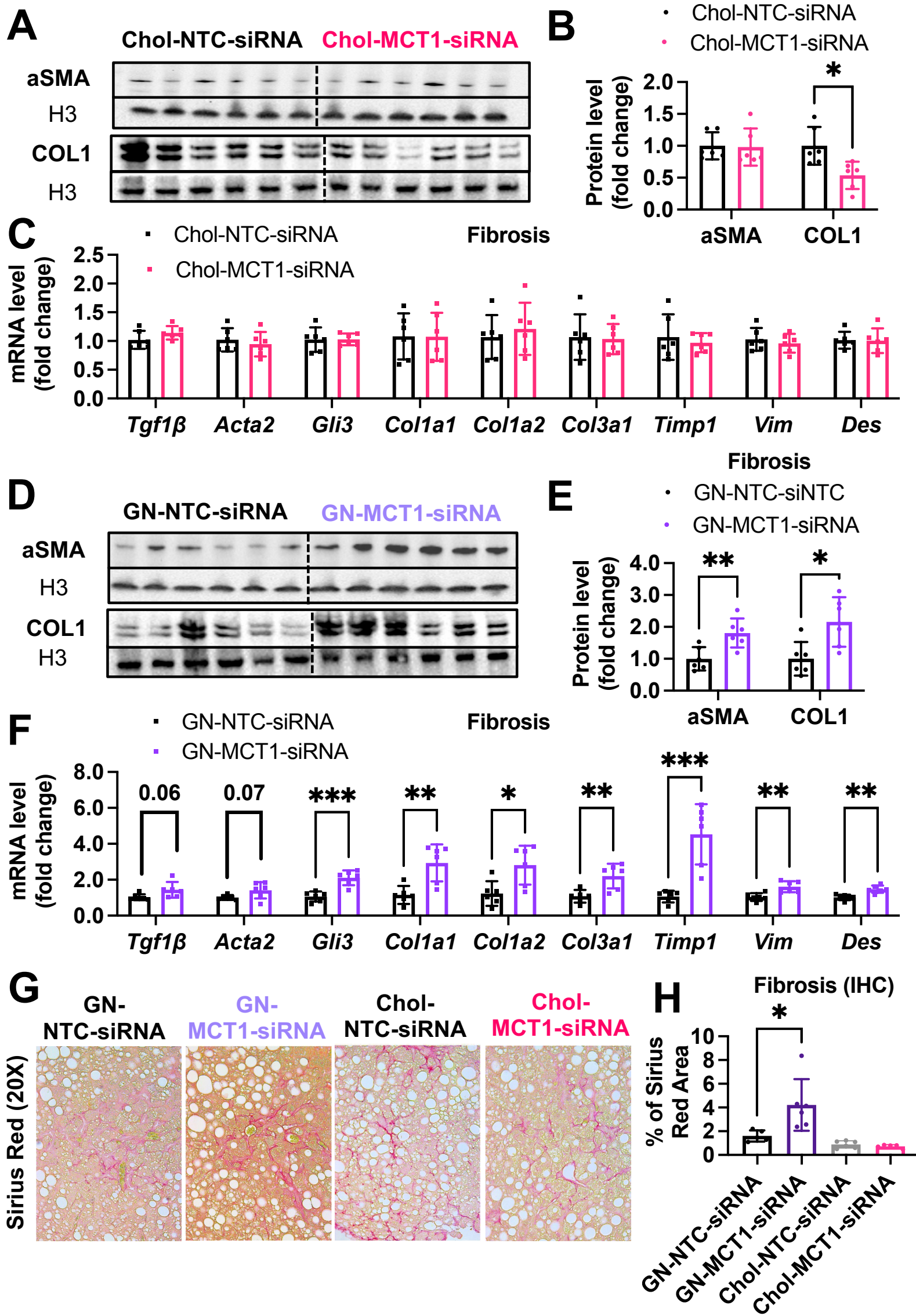


Figure 5



# Figure 6

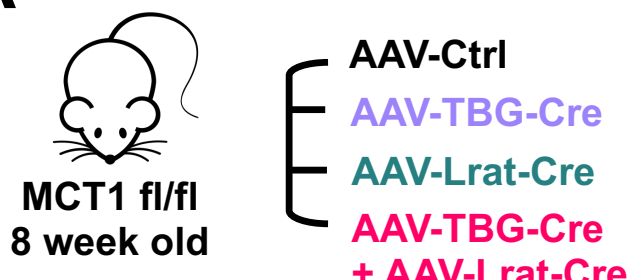
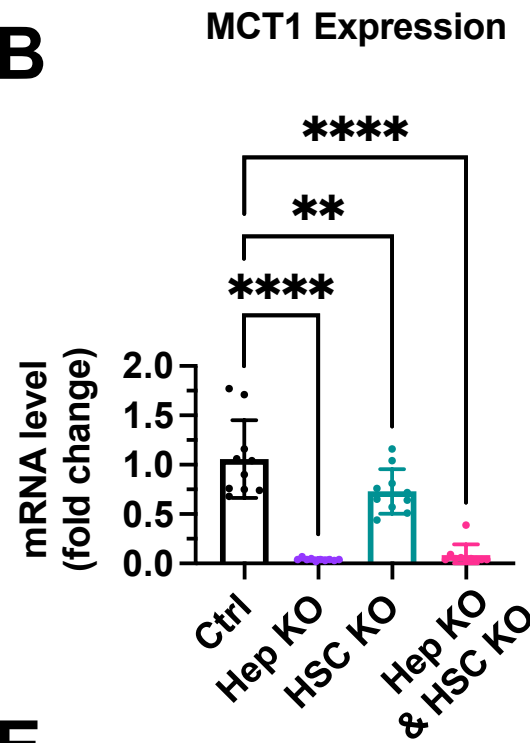
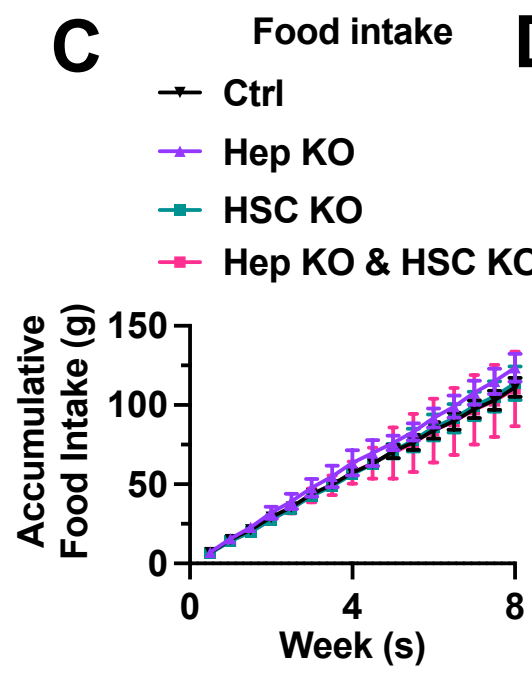
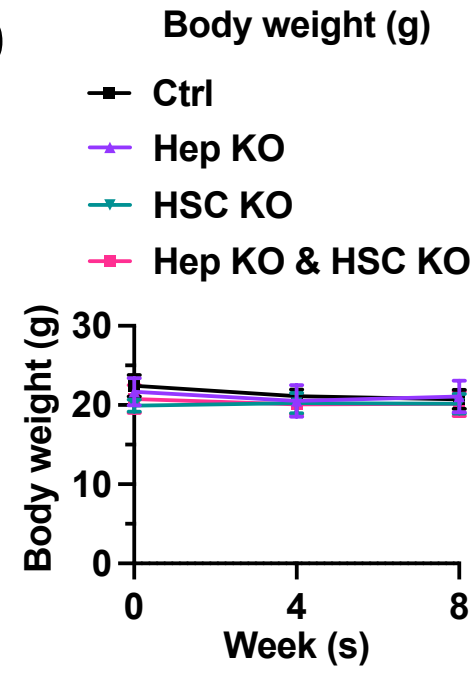
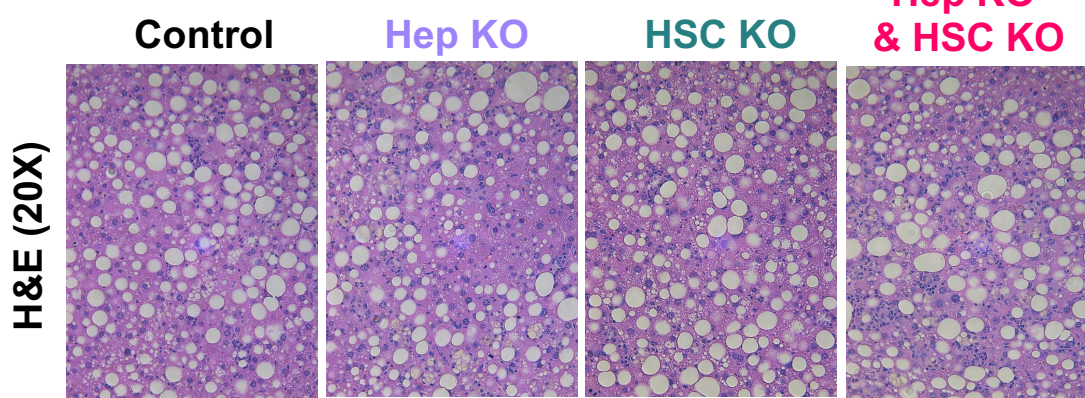
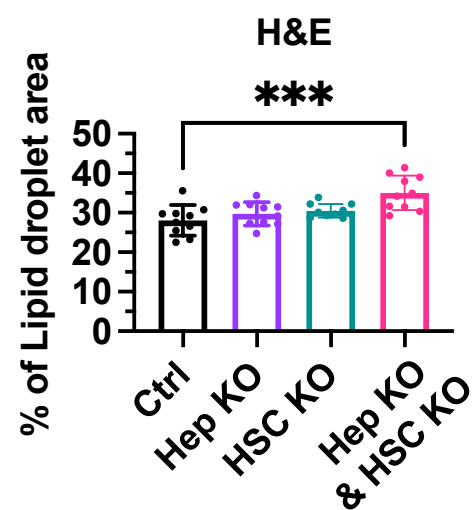
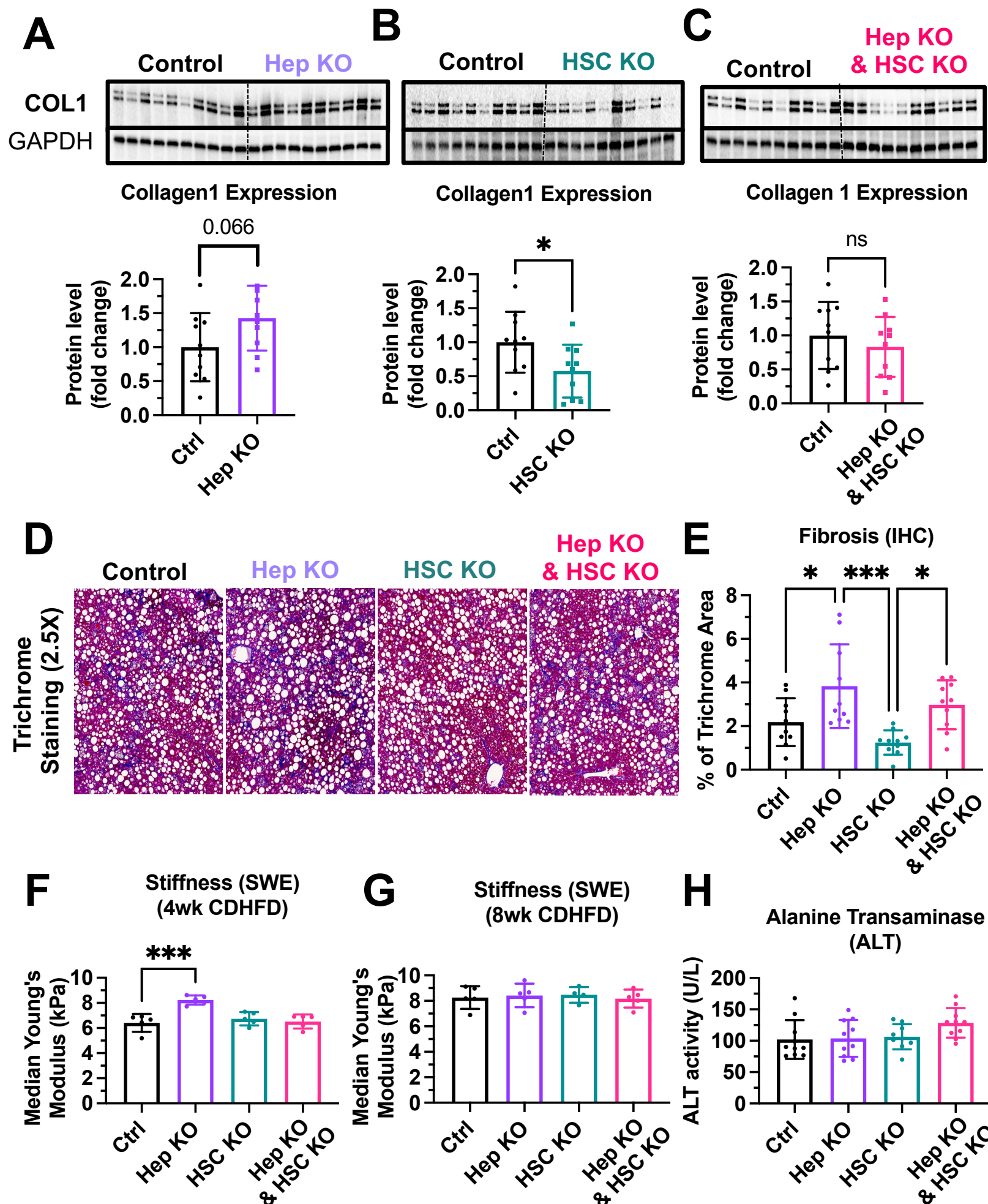
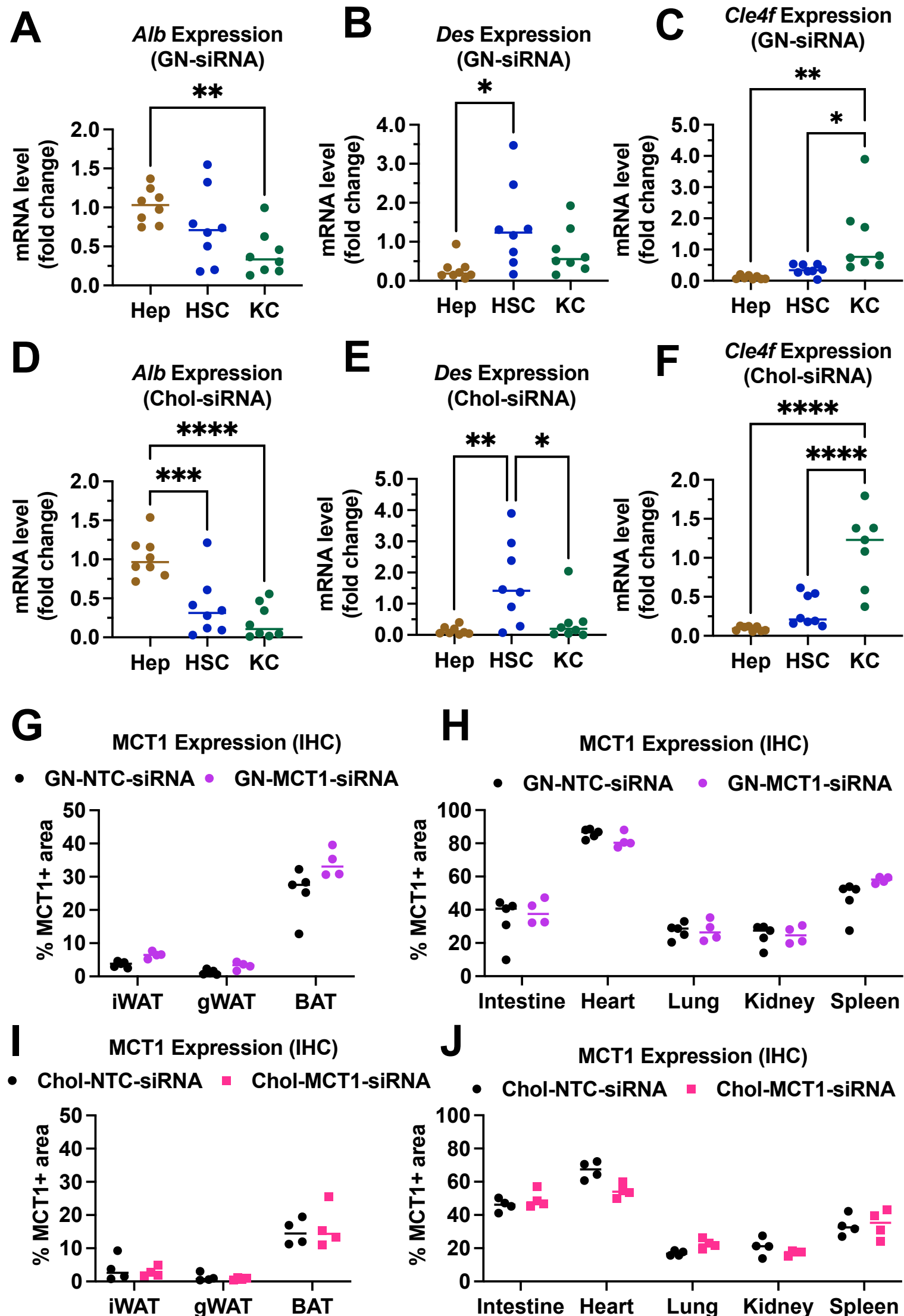
**A****B****C****D****E****F**

Figure 7



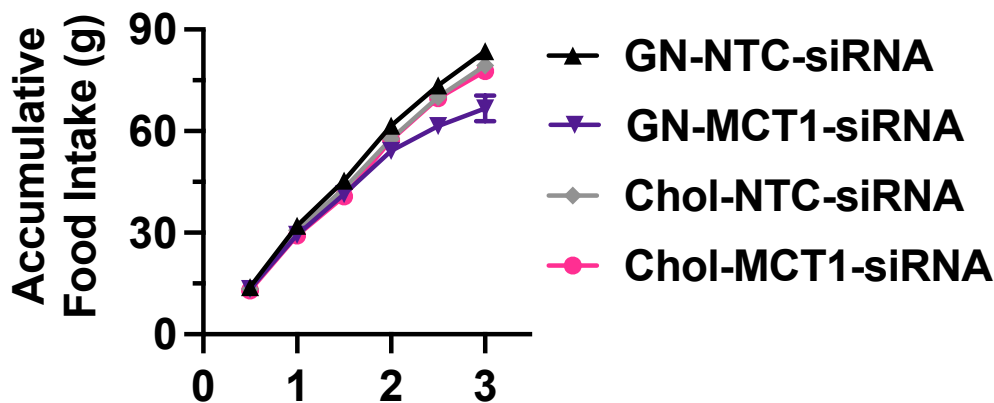
# Supplement Figure 1



# Supplement Figure 2

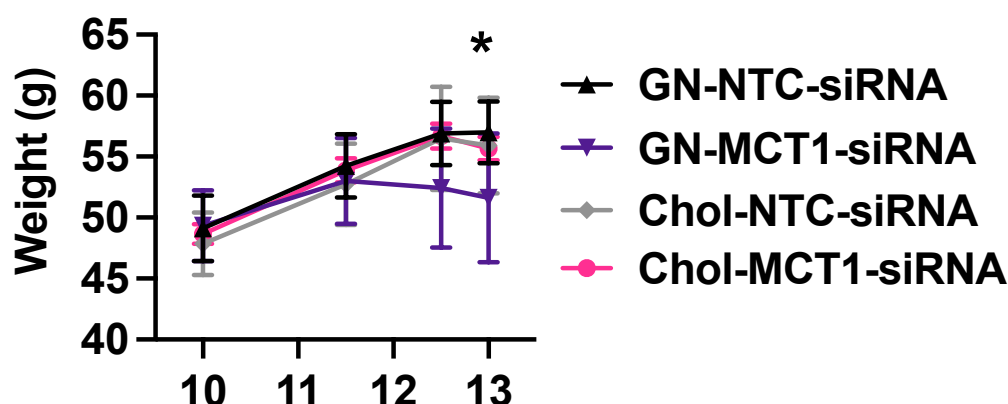
**A**

## Food Intake

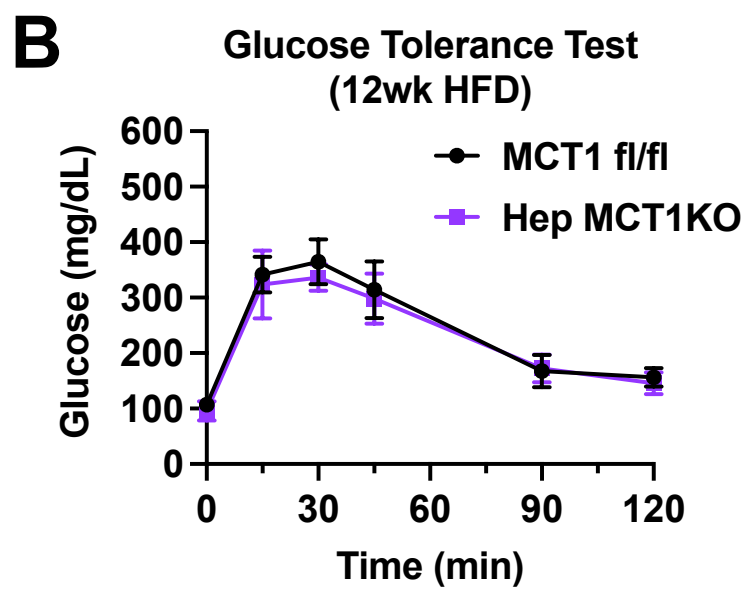
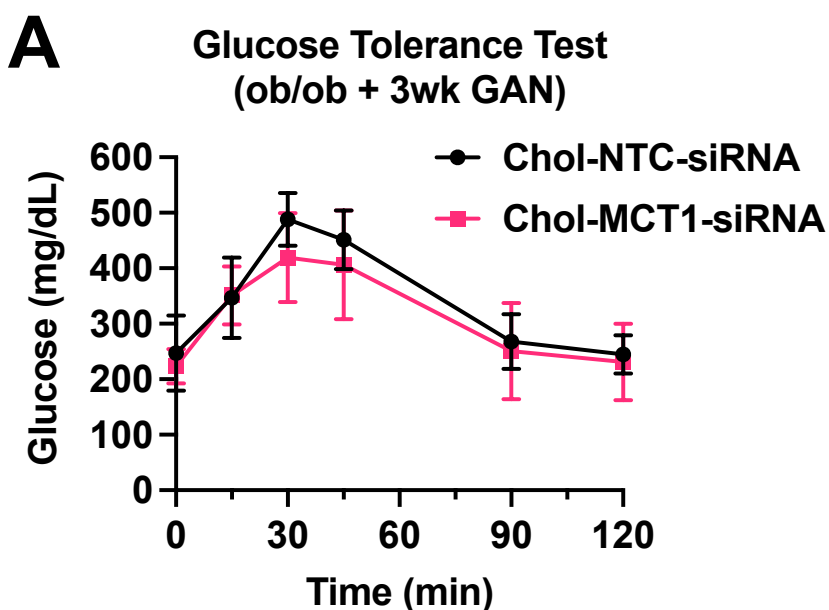


**B**

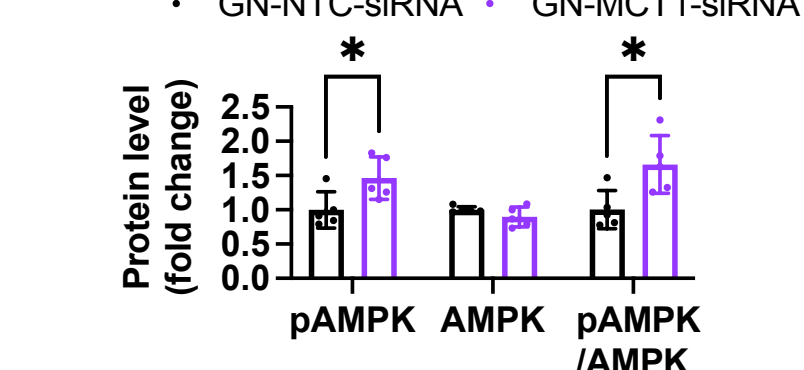
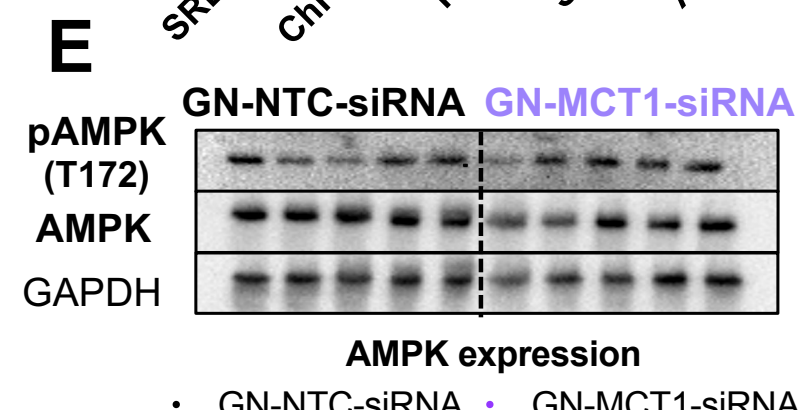
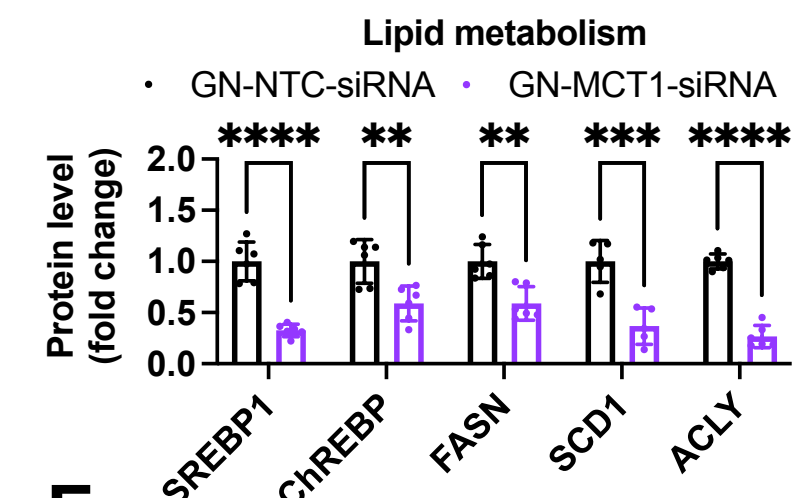
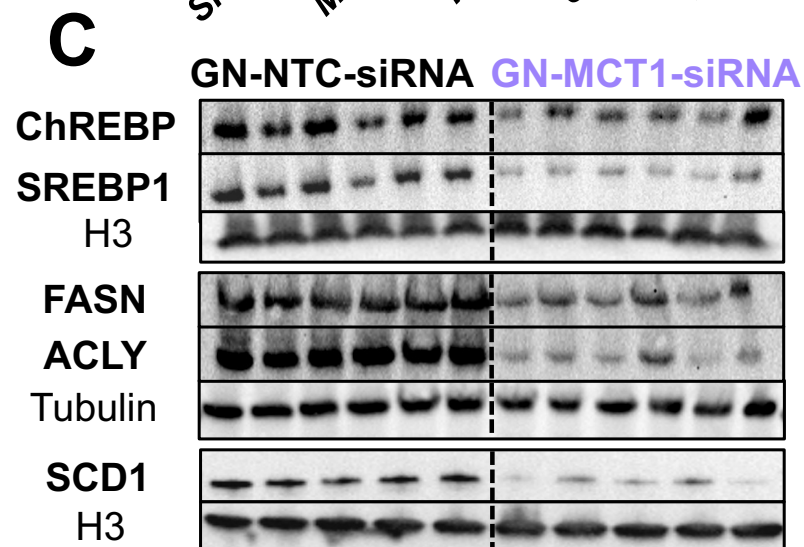
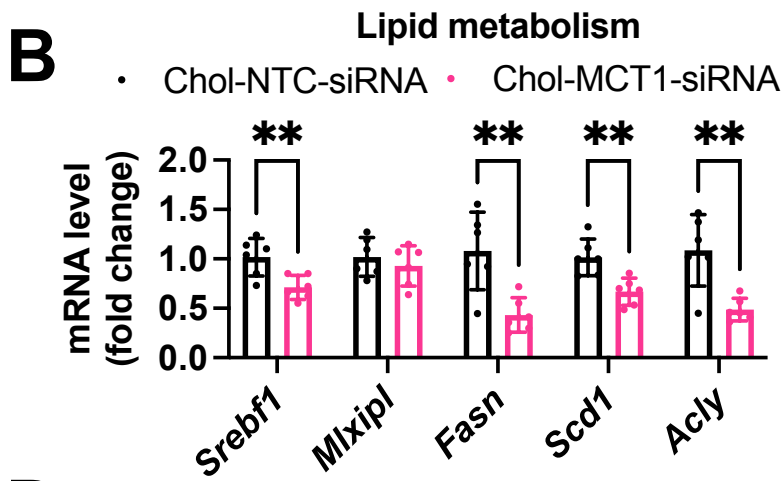
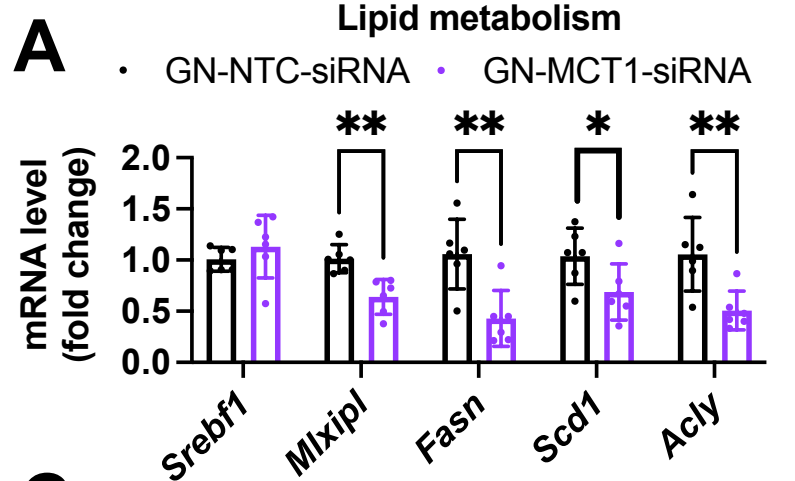
## Body Weight



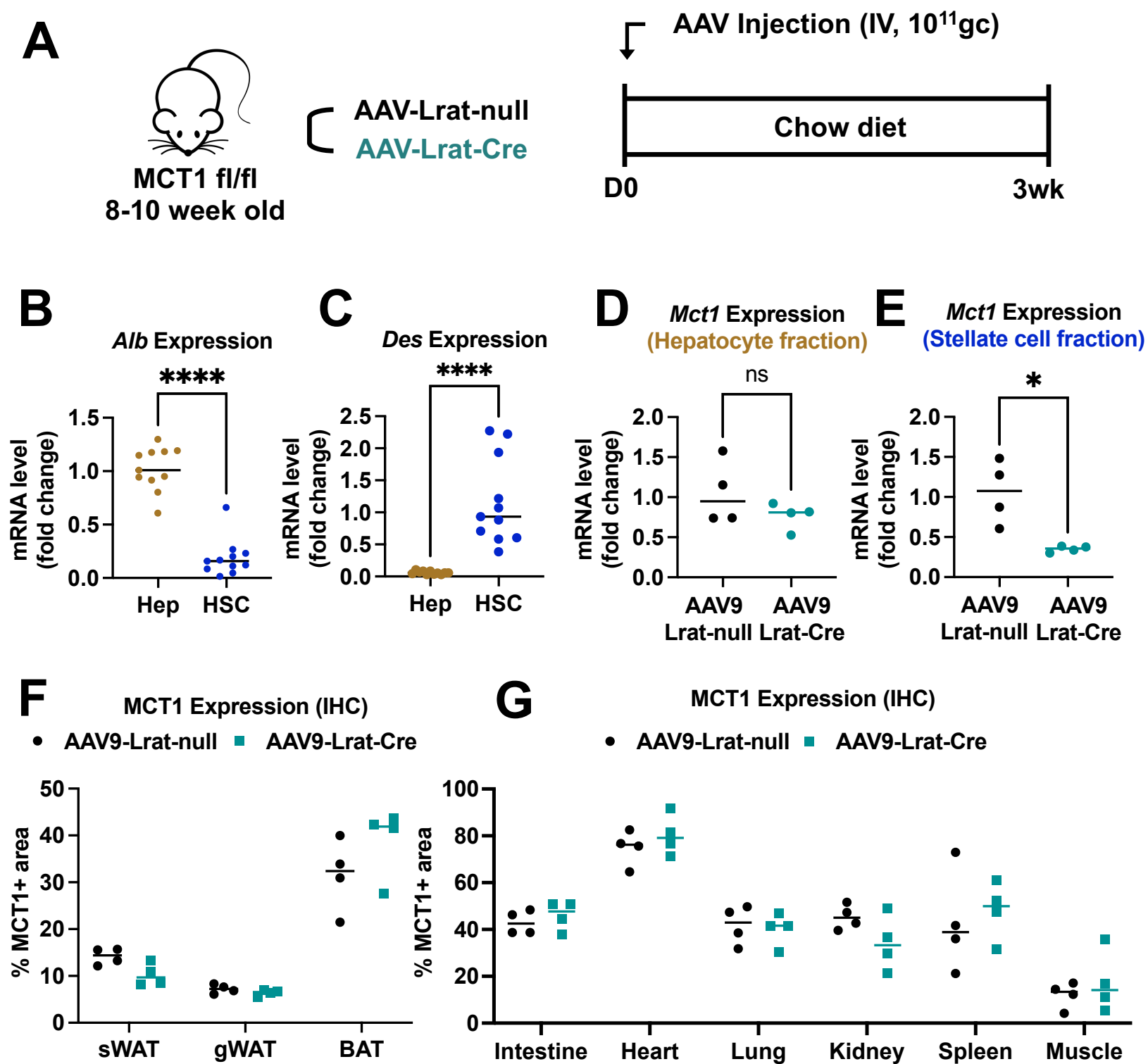
# Supplement Figure 3



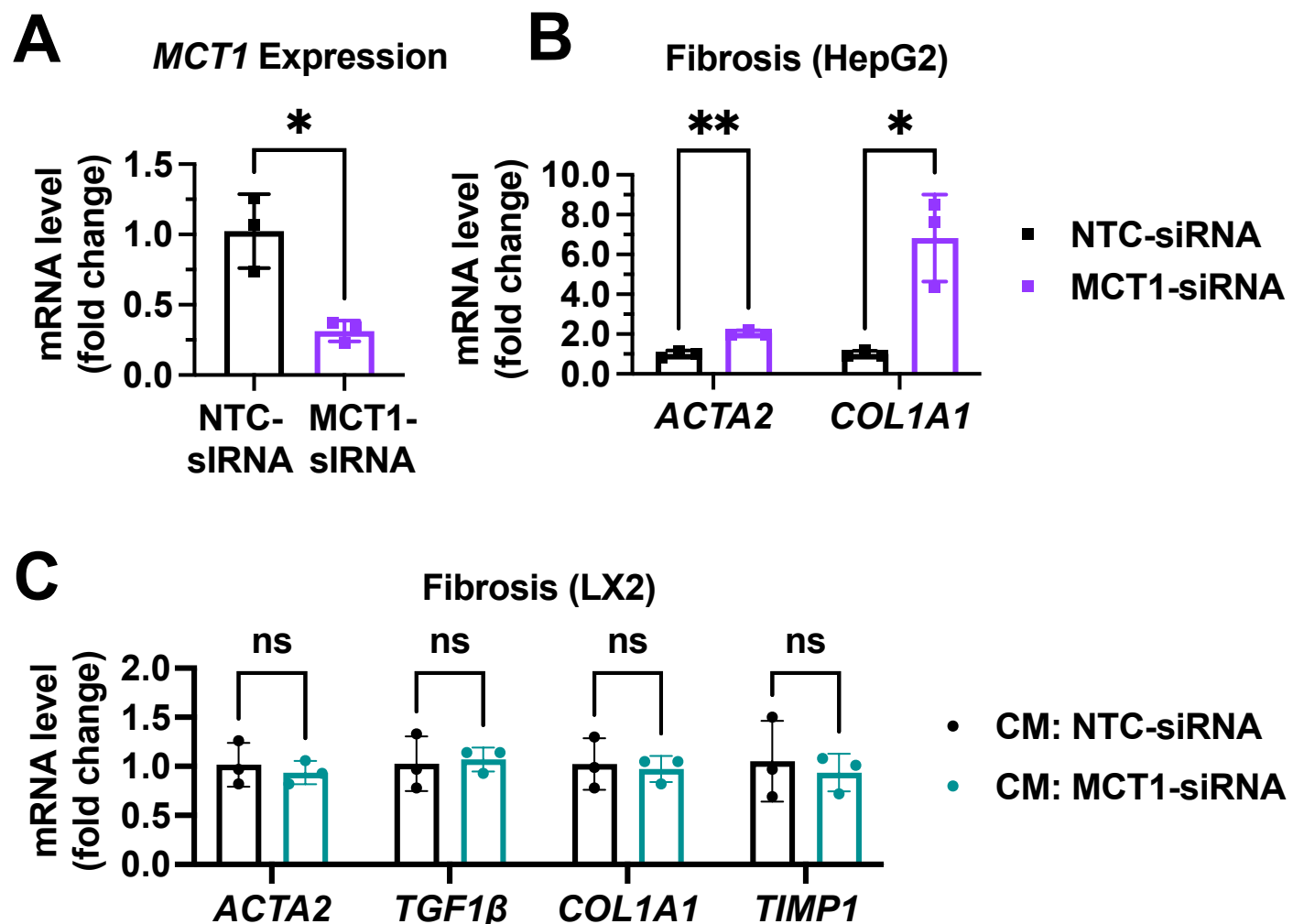
# Supplement Figure 4



## Supplement Figure 5



# Supplement Figure 6



# Supplement Figure 7

