Hepatocyte TAZ/WWTR1 Promotes Inflammation and Fibrosis in Nonalcoholic Steatohepatitis

Highlights
- TAZ/WWTR1 is increased in hepatocytes in human and mouse NASH liver
- Hepatocyte TAZ silencing in steatosis blocks inflammation, cell death, and fibrosis
- Hepatocyte TAZ silencing in NASH reverses inflammation, cell death, and fibrosis
- Hepatocyte TAZ promotes fibrosis by inducing Ihh, a hepatic stellate cell activator

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In Brief
Hepatic fibrosis is a key feature of nonalcoholic steatohepatitis (NASH) affecting morbidity. Wang et al. investigate the underlying mechanisms for progression from benign steatosis to NASH and show that the transcription factor TAZ/WWTR1 is increased in mouse and human NASH. Silencing TAZ can prevent or reverse NASH features, notably fibrosis.
Hepatocyte TAZ/WWTR1 Promotes Inflammation and Fibrosis in Nonalcoholic Steatohepatitis

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http://dx.doi.org/10.1016/j.cmet.2016.09.016

SUMMARY

Nonalcoholic steatohepatitis (NASH) is a leading cause of liver disease worldwide. However, the molecular basis of how benign steatosis progresses to NASH is incompletely understood, which has limited the identification of therapeutic targets. Here we show that the transcription regulator TAZ (WWTR1) is markedly higher in hepatocytes in human and murine NASH liver than in normal or steatotic liver. Most importantly, silencing of hepatocyte TAZ in murine models of NASH prevented or reversed hepatic inflammation, hepatocyte death, and fibrosis, but not steatosis. Moreover, hepatocyte-targeted expression of TAZ in a model of steatosis promoted NASH features, including fibrosis. In vitro and in vivo mechanistic studies revealed that a key mechanism linking hepatocyte TAZ to NASH fibrosis is TAZ/TEA domain (TEAD)-mediated induction of Indian hedgehog (Ihh), a secretory factor that activates fibrogenic genes in hepatic stellate cells. In summary, TAZ represents a previously unrecognized factor that contributes to the critical process of steatosis-to-NASH progression.

INTRODUCTION

The epidemic of obesity has led to the occurrence of fatty liver, or steatosis, in hundreds of millions of people worldwide. While simple steatosis is a relatively benign condition, approximately 20%–30% of these subjects will develop liver inflammation, dysfunctional fibrosis, and hepatocyte death, a serious condition known as nonalcoholic steatohepatitis (NASH) (Rinella, 2015). NASH can progress to cirrhotic liver disease and hepatocellular carcinoma and has become the leading cause of liver failure (Corey and Kaplan, 2014; Rinella, 2015). Despite the high prevalence and clinical importance of NASH, many gaps remain in our understanding of its pathophysiology, leading to a lack of mechanism-based therapeutic targets and treatment options (White et al., 2012).

NASH most likely develops as a result of multiple hits (Day and James, 1998), including obesity/insulin resistance-mediated steatosis and other insults that promote inflammation, fibrosis, and hepatocyte death (Singh et al., 2015). However, the molecular mechanisms corresponding to these pathogenic processes and their integration are poorly understood. In particular, there is a great need to elucidate the mechanisms leading to hepatic fibrosis, which is the leading determinant of long-term mortality in patients with NASH (Angulo et al., 2015a, 2015b; Puche et al., 2013). Extensive data indicate that activation of hepatic stellate cells (HSCs) plays a key role in NASH fibrosis (Mederacke et al., 2013), and although a number of factors have been proposed to activate HSCs in NASH, the work in this area is far from complete and has not yet led to FDA-approved treatment strategies (Angulo et al., 2015b; Puche et al., 2013).

In this context, we became interested in the Hippo pathway transcriptional activator TAZ (also known as WWTR1), which has been shown to have a pathophysiologic role in fibrosis in the lung (Liu et al., 2015). TAZ and the related protein YAP are orthologs of Drosophila Yorkie, a regulator of gene transcription that promotes tumor development in flies (Zanconato et al., 2016; Zhao et al., 2010). Although the roles of TAZ/YAP in normal physiology in mammals appear modest, they have emerged as key factors in carcinogenesis, tumor progression, and metastasis. At a cell biological level, TAZ/YAP mediate cellular responses to both intracellular shape changes and changes in the extracellular matrix environment. In the basal state, TAZ/YAP are relatively inactive as nuclear transcriptional regulators, because kinases in the so-called Hippo pathway maintain...
TAZ/YAP in their phosphorylated, cytoplasmic state. Through dephosphorylation and other processes, however, TAZ/YAP translocate to the nucleus and become active transcriptional regulators. In the nucleus, they interact with TEA domain (TEAD) proteins, leading to transcription of genes possessing consensus TEAD-binding cis-regulatory sequences in promoter and enhancer regions.

While there are some hints in the literature that TAZ or YAP may be involved in various liver processes, their possible role in fatty liver disease or NASH progression has not been reported. We now demonstrate that TAZ expression is elevated in the livers of humans with NASH-related fibrosis and in the livers of several murine NASH models. Silencing of hepatocyte Taz in NASH mice prevents and even reverses the development of liver inflammation and fibrosis without affecting steatosis, and forced expression of TAZ in hepatocytes can promote the progression of NASH from steatosis. In vitro, conditioned medium from Taz-silenced versus control hepatocytes decreases fibrotic responses in hepatic stellate cells (HSCs), which we link both in vitro and in vivo in murine NASH to decreased hepatocyte secretion of the TAZ/TEAD target Indian hedgehog (Ihh). Thus, TAZ promotes NASH progression, including fibrosis, and therefore emerges as a potential therapeutic target to prevent the progression of steatosis to NASH.

RESULTS

TAZ Levels Are Increased in the Livers of Humans and Mice with NASH

We conducted TAZ immunofluorescence microscopy on human liver samples from obese individuals with normal, neutrophic, and NASH histology. While there was similar TAZ staining in normal and steatotic livers, we observed a significant increase in TAZ staining in the NASH samples (Figure 1A). The specificity of the anti-human TAZ antibody for immunofluorescence is demonstrated by a silencing RNA-targeting Taz (siTaz) experiment conducted with human HepG2 liver cells (Figure S1A). TAZ localization in NASH liver was not confined to any one zone (Figure S1B), and most of the TAZ-stained cells were hepatocytes, as identified by HNF4α staining (Figure S1C). We then analyzed TAZ protein by immunoblot in liver extracts from subjects with NASH versus early non-alcoholic fatty liver disease (NAFLD) and normal liver and found that TAZ was expressed at the highest level in NASH liver (Figure 1B).

In preparation for TAZ causation studies, we explored TAZ expression in various mouse models of NASH. We began with the widely used methionine-choline-deficient (MCD) diet model, which induces NASH-like liver pathology despite weight loss and insulin sensitivity (Hebbard and George, 2011), and found that TAZ expression was markedly higher in MCD liver compared with control liver (Figure 1C). We next sought to study TAZ in a NASH model that had weight gain and insulin resistance, as is the case with humans. For this purpose, we modified two previously described diet-induced weight-gain models (Charlton et al., 2011; Kohl et al., 2010) to achieve NASH features within an experimentally acceptable time frame. We devised a diet that was rich in fructose, palmitate, cholesterol (FPC), and trans-fat, with other features as detailed in Tables S1 and S2. Cholesterol (1.2% by food weight) was added to the diet in view of multiple links between liver cholesterol and NASH in humans (Ioannou, 2016). Note that a high dietary content of cholesterol is needed to achieve adequately increased cholesterol absorption in C57BL/6J mice, the strain used here (Jolley et al., 1999).

After 16 weeks, FPC-fed mice had higher body weight and liver:body weight ratio than chow-fed mice (Figures S2A and S2B). Additionally, FPC mice showed significant increases in fasting blood glucose, plasma insulin, and alanine aminotransferase (ALT) and aspartate aminotransferase (AST) levels (Figures S2C–S2E). Analyses of liver sections revealed that the following parameters were greater in the livers of FPC-fed mice versus chow-fed mice: lipid droplet area (H&E and Oil Red O [ORO]), inflammatory cells, fibrosis (aniline blue component of trichrome [Trichr] and Sirius red [Sir red]), and cell death (TUNEL) (Figures S2F–S2K).

As further evidence of inflammation, FPC liver had elevated liver mRNA levels for Tnfa, Mcp1, and F4/80 (Adgre1; macrophages) and a higher percentage of F4/80+ cells (Figures S2L and S2M). With regard to fibrosis-associated parameters, hepaticTgfb1 and Acta2 (α-smooth muscle actin, α-SMA) mRNAs were higher in FPC-fed mice compared with chow-fed mice (Figure S2N), and there was also an increase in α-SMA+ cells (Figures S2O). Moreover, as designed, liver cholesterol was elevated in the livers of FPC mice (Figures S2P and S2Q). Additional features of the model in terms of blood parameters and liver assays are shown in Figure S3, and Table S3 shows hepatic fatty acid changes similar to those reported for human NASH liver (Yamada et al., 2015). Finally, Figures S4A–S4G show a direct comparison with one of the models from which it was derived, referred to as the “fast food” (FF) model (Charlton et al., 2011).

Most importantly, for the purpose of our study, the livers of FPC mice express high levels of TAZ protein and Wwtr1 (Taz) mRNA (Figures 1D and 1E). By immunoblot analysis, TAZ was abundant in extracts of primary hepatocytes from FPC-fed mice, but it could not be detected in extracts of combined non-hepatocytes (Figure S4H). Moreover, in view of the fact that the transcriptionally active form of TAZ is non-phosphorylated and nuclear (Liu et al., 2011), we found that the livers of FPC-fed mice had a lower phospho-TAZ:total TAZ ratio and higher nuclear TAZ compared with the livers of chow-fed mice (Figures S4I and S4J). Collectively, these data show that hepatocyte TAZ is induced in the livers of humans and mice with NASH, which led us to explore the possibility that TAZ may be a contributor to the progression from benign steatosis to NASH.

TAZ Silencing in Hepatocytes Suppresses Liver Inflammation, Fibrosis, and Cell Death in Mouse Models of NASH

To study TAZ function in NASH development, we used an adenovirus type 8 vector in which short hairpin RNA-targeting Taz (shTaz) was driven by the H1 promoter (AAV8-H1-shTaz), which was designed to silence genes in liver (Lisowski et al., 2014). (NB: to conform to the literature, we use the common name Taz, rather than Wwtr1, for sh terminology.) Compared with AAV8-H1 control vector, AAV8-H1-shTaz led to robust silencing of TAZ (Figure 2A) and a decrease in the known TAZ/TEAD transcriptional targets, Birc5 and Rhamm (Figure S5A).
In terms of specificity within the liver, note that TAZ expression in NASH liver was abundant in hepatocytes but not detected by immunoblot in combined non-hepatocytes (above). In extracts of adipose, heart, lung, and skeletal muscle, TAZ was not decreased in mice treated with AAV8-H1-shTaz; note that TAZ was not detected in kidney extract (Figure S5B).

Mouse body weight, liver:body weight ratio, fasting blood glucose, plasma insulin, and plasma cholesterol were similar in the shTaz and control groups (Figures S5C–S5G). Liver sections showed marked reductions in both inflammatory cell infiltration and fibrosis endpoints in the shTaz cohort (Figures 2B–2D), as reflected in fibrosis stage and NAFLD activity score (NAS) (Figures S5H and S5I), while steatosis was not affected (Figure S5J). Plasma ALT was decreased in shTaz-treated mice (Figure 2E), and this was associated with a decrease in TUNEL+ and 4-HNE+ liver cells (Figure 2F; Figure S5K), indicating that Taz silencing reduced both cell death and oxidative stress in liver cells. At the mRNA level, Taz silencing caused a robust reduction in the expression of mRNAs related to hepatic inflammation—Tnfa, Mcp1, and F4/80 (Adgre1)—and fibrosis (Figures 2G and 2H), including the NASH-relevant genes Acta2 (α-SMA), Timp1, Des, Col1a1, Col1a2, Col3a1, and Vim (Friedman, 2008; Younossi et al., 2011). These changes were accompanied by decreases in both F4/80+ macrophages and α-SMA+ cells (Figures 2I and 2J; Figure S5L). Note that silencing TAZ did not change filipin staining of liver, suggesting...
Figure 2. TAZ Silencing Reduces Liver Inflammation, Fibrosis, and Cell Death in FPC-Fed Mice

The following parameters were measured in male C57BL/6J mice treated with AAV8-H1-shTaz or control vector and then fed the FPC diet for 16 weeks (*p < 0.05, **p < 0.01, ***p < 0.0002, mean ± SEM; n = 10 mice/group):

(A) Immunoblot of TAZ in liver, with β-actin as loading control.

(B) Staining of liver sections for H&E (upper panels; bar, 100 μm), Masson’s trichrome (Trichr) (middle panels; bar, 100 μm), and Sirius red (Sir red) (lower panels; bar, 500 μm).

(C) Hepatic inflammatory cells.

(D) Aniline blue area (%)

(E) ALT (IU/L)

(F) TUNEL+ cells (%)

(G) Relative mRNA expression

(H) Relative mRNA expression

(I) F4/80

(J) α-SMA

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that TAZ is downstream of liver cholesterol accumulation (Figure S5M).

We conducted a similar study in which hyperphagic Mc4r−/−mice (Butler and Cone, 2002) were fed the FPC diet for 16 weeks. These mice develop more liver fibrosis compared with FPC-fed wild-type (WT) C57BL/6J mice, and silencing of Taz in the liver of these mice resulted in decreased staining for aniline blue, Sirius red, and α-SMA; lower hydroxyproline content; and decreased liver inflammation (Figure S6). We confirmed these results in the MCD model, where AAV8-H1-shTaz decreased both hepatic inflammation and fibrosis without affecting steatosis and also reduced inflammatory and fibrotic gene expression, α-SMA+ cells, and macrophages in the liver (Figures S7A–S7J). In contrast, in a model of liver fibrosis in which TAZ is not increased in the liver—CCI4-treated mice—silencing Taz did not suppress fibrosis or fibrotic gene expression (Figures S7K–S7M). We also conducted an experiment in MCD-fed Wwtr1fl/fl mice (Xin et al., 2013) treated with AAV8-TBG-LacZ control or AAV8-TBG-Cre, which deletes floxed genes specifically in hepatocytes versus other types of liver cells (Mu et al., 2015) or other organs (Figures S8A and S8B). As above, inflammatory endpoints, α-SMA+ cells, fibrosis-related gene expression, and cell death were decreased by hepatocyte TAZ knockdown, as was fibrosis itself, despite the fact that the level of fibrosis in the control cohort in this particular experiment was less than usual (Figures S8C–S8H).

We next asked whether features of NASH, notably fibrosis, could be reversed by treating mice with AAV8-H1-shTaz after NASH had developed. For this purpose, mice were placed on the FPC diet for 16 weeks, treated with shTaz or control vector, and then continued on the diet for an additional 12 weeks. As expected for 28 weeks of FPC feeding, the control mice had a marked increase in hepatic TAZ compared with mice mutated sequence in control AML12 cells, with a marked decrease in the ChIP signal in the livers of FPC-fed versus chow-fed mice, which was dependent on anti-TAZ and was not seen when a non-consensus sequence was amplified. Most importantly, the ChIP signal in the livers of FPC mice was lowered to the chow level by TAZ silencing. Next, we conducted a dual-luciferase reporter assay using this intron 1 consensus sequence versus a mutated sequence, with 500 bp of upstream and downstream flanking sequence. The luciferase vectors were transfected into siControl- or siTaz-treated AML12 cells, a non-cancerous mouse hepatocyte cell line (Dumenco et al., 1995). The data showed robust luciferase activity with the consensus sequence versus a mutated sequence, with 500 bp of upstream and downstream flanking sequence. The luciferase vectors were transfected into siControl- or siTaz-treated AML12 cells, a non-cancerous mouse hepatocyte cell line (Dumenco et al., 1995).

To investigate the role of TAZ specifically in steatosis-to-NASH progression, mice were first fed the FPC diet for 8 weeks and then injected with AAV8-H1-shTaz or control virus, followed by an additional 8 weeks on the diet. After 8 weeks, there was only a slight increase in hepatic TAZ compared with mice fed the FPC diet for 16 weeks, and the livers showed steatosis but no appreciable inflammation or fibrosis (Figures 4A and 4B). The livers of mice who received shTaz at week 8 showed marked reductions at 16 weeks in fibrosis endpoints, inflammatory cells, inflammatory- and fibrosis-related genes, F4/80+ macrophages, and α-SMA+ cells, but not steatosis, and plasma ALT was also decreased by shTaz (Figures 4B–4H). These data suggest that TAZ is particularly important in the critical processes that promote steatosis-to-NASH progression.

Finally, whereas the above experiments addressed necessity of TAZ in NASH, we addressed the question of sufficiency by treating mice on the aforementioned FF diet, which is a model of steatosis in the time frame studied (above), with AAV8-albumin-TAZ. We achieved a level of hepatic TAZ protein in these mice that was similar to that in FPC-fed mice, and this was associated with an increase in all parameters of inflammation and fibrosis (Figure S9). Thus, TAZ is both necessary and sufficient to promote the development of NASH in mice.

Hepatocyte TAZ Silencing in Hepatocytes Blocks Steatosis-to-NASH Progression, and Genetically Induced TAZ Expression in Hepatic Stellate Cells and Mediates TAZ-Induced NASH

We sought to explore possible mechanisms linking TAZ to fibrosis progression. Previous studies have demonstrated that HSCs, the main source of collagen-producing myofibroblasts in NASH-related fibrosis (Mederacke et al., 2013), can be activated by the hedgehog pathway (Angulo et al., 2015b). In this context, chromatin immunoprecipitation (ChIP) array data suggested that the gene encoding Indian hedgehog, Ihh, is a TAZ/TEAD target. We therefore considered the hypothesis that increased TAZ in hepatocytes during NAFLD progression leads to the secretion of Ihh, which then acts on HSCs to promote the expression of pro-fibrotic genes.

To begin, we conducted TAZ ChIP analysis of livers of chow-fed and FPC-fed mice with or without TAZ silencing, focusing on a TAZ/TEAD consensus sequence in intron 1 of murine Ihh that is conserved among species, including humans (Zanconato et al., 2015)(Figures 5A and 5B). The results show a significant increase in the ChIP signal in the livers of FPC-fed versus chow-fed mice, which was dependent on anti-TAZ and was not seen when a non-consensus sequence was amplified. Most importantly, the ChIP signal in the livers of FPC mice was lowered to the chow level by TAZ silencing. Next, we conducted a dual-luciferase reporter assay using this intron 1 consensus sequence versus a mutated sequence, with 500 bp of upstream and downstream flanking sequence. The luciferase vectors were transfected into siControl- or siTaz-treated AML12 cells, a non-cancerous mouse hepatocyte cell line (Dumenco et al., 1995). The data showed robust luciferase activity with the consensus sequence versus a mutated sequence, with 500 bp of upstream and downstream flanking sequence. The luciferase vectors were transfected into siControl- or siTaz-treated AML12 cells, a non-cancerous mouse hepatocyte cell line (Dumenco et al., 1995).

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with TAZ (above), the expression of Ihh was greater in NASH liver than in normal and steatotic liver (Figure 5D). Similarly, the livers of FPC-fed mice had markedly higher levels of Ihh compared with the livers of chow-fed mice (Figure 5E). We then compared chow and FPC liver extracts for gene expression of Ihh and the Ihh pathway downstream genes Gli2 and Gli3. All three mRNAs were elevated in FPC liver, as was an Ihh target gene, osteopontin (Opn) (Razzaque et al., 2005), which is involved in HSC-induced fibrosis in NASH (Syn et al., 2011) (Figure 5F). To explore causation with regard to TAZ, we repeated these assays in FPC-fed mice with or without TAZ silencing. All four mRNAs and Ihh protein were substantially lower in the TAZ-silenced mice (Figures 5G and 5H), as was OPN as assessed by immunohistochemistry (Figure 5I). Thus, TAZ induces transcriptionally active Ihh during NASH progression in FPC-fed mice, and one of the targets of Ihh, Opn, has been linked to NASH fibrosis (Bohinc and Diehl, 2012; Syn et al., 2011).

Figure 3. TAZ Silencing after the Development of NASH Reduces Liver Inflammation and Fibrosis in FPC-Fed Mice

The following parameters were measured in male C57BL/6J mice fed the FPC diet for 28 weeks, with AAV8-H1-shTaz or control vector administered at the 16 week time point (*p < 0.05; **p < 0.01, ***p < 0.0001, mean ± SEM; n = 5 mice/group):

(A) Staining of liver sections for H&E (upper panels; bar, 100 μm), Masson’s trichrome (Trichr) (middle panels; bar, 500 μm), and Sirius red (Sir red) (lower panels; bar, 500 μm).

(B) Hepatic inflammatory cells.

(C) Aniline blue- and Sirius red-positive area.

(D) mRNA levels of Mcp1, Acta2 (α-SMA), Col1a1, Col3a1, Des, Vim, and Timp1.

(E) F4/80 immunofluorescence (red) and quantification; DAPI counterstain for nuclei is shown in bottom panels; bar, 500 μm.

(F) α-SMA immunofluorescence (red) and quantification; DAPI counterstain for nuclei is shown in bottom panels; bar, 500 μm.
Figure 4. TAZ Silencing after the Development of Steatosis Reduces Liver Inflammation and Fibrosis in FPC-Fed Mice

(A) TAZ immunoblot of liver extracts from C57BL/6J mice fed chow or FPC diet for 8 or 16 weeks.

(B) H&E- and Masson’s trichrome (Trich)-stained liver sections and quantified lipid droplet area, inflammatory cells, and aniline blue-positive area for C57BL/6J mice fed chow or FPC diet for 8 weeks. Bar, 100 μm.

(C–H) The following parameters were measured in male C57BL/6J mice fed the FPC diet for 16 weeks, with AAV8-H1-shTaz or control vector administered at the 8 week time point (*p < 0.05; **p < 0.01, ***p < 0.0001, mean ± SEM; n = 5 mice/group): staining of liver sections for H&E (upper panels; bar, 100 μm), Masson’s trichrome (Trich) (middle panels; bar, 100 μm), and Sirius red (Sir red) (lower panels; bar, 500 μm) (C); Aniline blue- and Sirius red-positive area and hepatic inflammatory cells (D); mRNA levels of Tnfa, Mcp1, Tgfb1, and Acta2 (α-SMA) (E); F4/80 immunofluorescence (red) and quantification; DAPI counterstain for nuclei is shown in bottom panels; bar, 100 μm (F); α-SMA immunofluorescence (red) and quantification; DAPI counterstain for nuclei is shown in bottom panels; bar, 100 μm (G); and plasma ALT (H).
Figure 5. TAZ Induces Ihh, and TAZ Silencing Lowers the Expression of Pro-Fibrotic Hedgehog Pathway Genes in the Livers of FPC-Fed Mice

(A) Conserved TAZ/TEAD consensus sequence in intron 1 of the mouse Ihh gene.

(B) Liver nuclear extracts from mice fed chow diet or FPC diet for 16 weeks with or without TAZ silencing were subjected to TAZ ChIP analysis using anti-TAZ or IgG control. The intronic region containing the TAZ/TEAD binding sequence, or a non-consensus sequence as control, was amplified by qPCR and normalized to the values obtained from input DNA (*p = 0.03; mean ± SEM; n = 3).

(C) A dual-luciferase reporter assay in control and TAZ-silenced AML12 cells was conducted using the Ihh intron 1 region in (B), or a mutated version, together with 500 bp of flanking upstream and downstream sequence. (**p < 0.01, ***p < 0.0001; mean ± SEM; n = 6).

(D) Immunoblot of Ihh in normal human livers or those with steatosis or NASH.

(E) Immunoblot of Ihh in the livers of mice fed chow or FPC diet for 16 weeks.

(F) Relative expression of Ihh, Gli2, Gli3, and Opn mRNAs in the livers of mice fed chow or FPC diet for 16 weeks (*p < 0.04, **p < 0.0001; mean ± SEM; n = 6).

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To explore the possibility that TAZ-induced Ihh is secreted by hepatocytes and activates HSCs, we moved to an in vitro model using both AML12 cells and primary murine HSCs. Consistent with our data in vivo, siTaz treatment of AML12 cells lowered cellular Ihh mRNA in the cells and Ihh protein in both the cells and media (Figures 6A–6C). We then added conditioned medium (CM) from control or Taz-silenced AML12 cells, as well as medium not exposed to cells (non-CM), to primary murine HSCs. Compared with non-CM, hepatocyte CM markedly increased Opn mRNA as well as the mRNAs for two proteins involved in fibrosis, Timp1 and Colla1. Most importantly, CM from TAZ-silenced hepatocytes lowered the levels these mRNAs and protein compared to CM from control hepatocytes (Figure 6D). Conditioned medium from Ihh-silenced hepatocytes also decreased the three mRNAs in HSCs (Figure 6E), although the absolute degree of Timp1 lowering was somewhat greater in this experiment than in the siTaz experiment. Finally, to make a more direct link between TAZ-induced Ihh in hepatocytes and activation of HSCs, we restored Ihh in siTaz-treated hepatocytes by Ihh transfection and then asked whether this lessened the suppressive effect of CM from these cells on the expression of the fibrosis-related genes in HSCs. Transfection of TAZ-silenced hepatocytes with Ihh led to a level of Ihh in the CM that was similar to that in the CM of control hepatocytes (Figure 6F—compare first and fourth bars). As before, the CM of TAZ-silenced hepatocytes suppressed Opn, Timp1, and Colla1 mRNA in HSCs (Figure 6G—compare second and fourth bars), and we found that restoration of Ihh in these TAZ-silenced cells rescued CM-induced HSC gene expression (Figure 6G—compare fourth and fifth bars). Finally, we wondered whether TAZ might also directly regulate fibrogenic gene expression or proliferation in HSCs. However, we found that silencing of Taz in primary HSCs affected neither fibrogenic gene expression nor proliferation (Figure 6H). Together, these studies suggest that hepatocyte-induced Ihh in hepatocytes promotes the expression of pro-fibrotic genes in HSCs.

To provide causation evidence in vivo for the role Ihh in TAZ-induced NASH, we first used AAV8-Ihh to restore Ihh in Taz-silenced FPC-fed mice. As designed, the level of hepatic Ihh mRNA in Ihh-restored shTaz mice was similar to that in control mice (Figure 7A, first set of bars). Consistent with the TAZ-Ihh hypothesis, Ihh restoration prevented the improvement in all NASH features that we see with Taz silencing, including liver inflammation, fibrosis and fibrogenic mRNA expression, and cell death (Figures 7A–7D). Next, we directly silenced Ihh in FPC-fed mice using AAV8-H1-shIhh and found that this intervention, like silencing Taz, improved all NASH-like features (Figures 7E–7L). These combined in vitro and in vivo data support the conclusion that hepatocyte TAZ promotes NASH progression in large part through inducing Ihh.

**DISCUSSION**

NASH, characterized by inflammation, cell death, and fibrosis, can progress to advanced liver disease, cirrhosis, and the need for liver transplant. Steatosis alone is believed to be little to no risk for progressive liver disease. Given the clinical significance of NASH compared to steatosis (Rinella, 2015), a critical objective of research in this area is to identify factors and pathways that promote the conversion of steatosis to NASH and the development of fibrosis. The importance of this objective is underscored by the fact that NASH is becoming the leading cause of liver disease worldwide and yet lacks any definitive, evidence-based drug therapies approved by the US Food and Drug Administration (Rinella, 2015). In this context, the finding that a hepatocyte TAZ-Ihh pathway plays a key role in steatosis-to-NASH conversion and the development of fibrosis provides new insight into NASH and may suggest new targets for therapy.

Hepatic fibrosis is a key feature of NASH that distinguishes it from steatosis and determines long-term mortality in patients with NASH (Angulo et al., 2015a). While both TAZ and YAP have been implicated in organ fibrosis in other settings, particularly in the lung with links to TGFβ-SMAD signaling or induction of plasminogen activator inhibitor-1 (Liu et al., 2015; Mitani et al., 2009; Piersma et al., 2015; Saito and Nagase, 2015), there are only scattered reports about their roles in liver fibrosis and none in the setting of NAFLD. For example, a recent study reported an association between microRNA-130-301, which can regulate TAZ and YAP, and CCl4-induced liver fibrosis (Bertero et al., 2015), but there were no direct causation or mechanistic data related to the role of TAZ in this process, and we now show that TAZ is not involved in CCl4-induced liver fibrosis. Another report showed that knockout of a pair of Hippo factors called Mps One Binder Kinase Activator (MOB)1A/1B in lean, non-NASH mice caused elevated TGFβ-2/3 and liver fibrosis in a manner that was partially dependent on TAZ (Nishio et al., 2016).

Although the mechanism of TAZ in liver fibrosis in NASH is likely to be multifactorial, we show here that the TAZ target Ihh is elevated in human and mouse NASH liver in a TAZ-dependent manner and is important in NASH progression and fibrosis. Previous work has implicated hedgehog signaling in NASH fibrosis, particularly Shh signaling in HSCs (Bohinc and Diehl, 2012), both directly and via the induction of the pro-fibrotic cytokine IL-13 by immune cells (Shimamura et al., 2008; Syn et al., 2010). ChIP array data (Zhao et al., 2008) suggested to us that Ihh might be a TAZ/TEAD target gene by interacting with a consensus site in intron 1 of the Ihh gene, which we have now confirmed using complementary ChIP and luciferase reporter assays. Most importantly, hepatocyte TAZ-induced Ihh activated a fibrosis program in HSCs in cell culture, restoration of Ihh in hepatocyte-TAZ-silenced mice restored NASH progression, and direct silencing Ihh in hepatocytes suppressed NASH.

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(G) Relative expression of Ihh, Gli2, Gli3, and Opn mRNAs in the livers of mice fed the FPC diet for 16 weeks with or without TAZ silencing (*p < 0.05, **p < 0.002; mean ± SEM; n = 10).
(H) Immunoblot of Ihh in the livers of mice fed the FPC diet for 16 weeks with or without TAZ silencing.
(I) OPN immunohistochemistry and quantification in the livers of mice fed the FPC diet for 16 weeks with or without TAZ silencing (*p < 0.0001; mean ± SEM; n = 10); bar, 200 μm.
Figure 6. TAZ-Induced Hepatocyte Ihh Increases the Expression of Fibrosis-Related Genes in Hepatic Stellate Cells

(A) Expression of Wwtr1 and Ihh mRNA in control and TAZ-silenced AML12 cells (*p < 0.0003; mean ± SEM; n = 3).
(B) Immunoblot of TAZ and Ihh in control and TAZ-silenced AML12 cells.
(C) Ihh concentrations, assayed by ELISA, in the media of control and TAZ-silenced AML12 cells (*p < 0.003; mean ± SEM; n = 3).
(D) Primary hepatic stellate cells (HSCs) were incubated for 72 hr with conditioned medium (CM) obtained from control or Taz-silenced AML12 cells or with medium not exposed to cells (non-CM). The HSCs were then assayed for Opn, Timp1, and Col1a1 mRNA (upper panel; *p < 0.05; mean ± SEM; n = 4) and the respective proteins by immunoblot (lower panel).

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Two other critical features of NASH, inflammation and cell death, were also ameliorated by TAZ silencing in our study. Little is known about the pro-inflammatory roles of TAZ, and, in general, YAP and TAZ inhibit, rather than promote, apoptosis during development and in cancer (Yu et al., 2015). However, there is one report showing that siTaz decreased TNFα-induced apoptosis in salivary gland epithelial cells (Hwang et al., 2014). RIP3-mediated necroptosis may also be important in hepatocyte death in NASH (Gautheron et al., 2014), and therefore it is possible that TAZ promotes this pathway. Given the various consequences of cell necrosis, this action of TAZ could contribute to inflammation and fibrosis, as well as cell death, in NASH (Chan et al., 2015; Luedde et al., 2014).

Our study focused on TAZ in hepatocytes, but a recent report showed that YAP was significantly increased in progenitor-like reactive-appearing ductular cells (RDCs) in human and mouse NASH liver (Machado et al., 2015). We also found increased YAP in the livers of FPC mice, and consistent with the data of Machado et al., most of the YAP-positive cells did not co-localize with HNF4α-positive hepatocytes (data not shown). While the role of YAP in NASH remains to be elucidated, the authors showed correlations among YAP-α RDCs, fibrosis, accumulation of myofibroblasts, and expression of Shh and Opn. Of note, silencing TAZ in the livers of FPC mice did not affect YAP expression (data not shown).

Most of our experiments were conducted in a mouse model of insulin resistance and NAFLD that was a modification of previously described models (Charlton et al., 2011; Kohli et al., 2010). The diet was based on human dietary risk factors for NASH, and the key improvement over previous models was the development of inflammation, hepatocyte death, and fibrosis in 16 weeks without the need for genetically engineered mutations and in the background of weight gain and insulin resistance. Whereas the fructose component of the diet likely contributes to steatosis (Abdelmalek et al., 2010; Ishimoto et al., 2013), the palmitate and palmitic acid components may be important in NASH progression and perhaps TAZ induction. The accumulation of unesterified cholesterol in the liver has been implicated in the development of NASH in humans (Ioannou, 2016) and in various mouse models (Subramanian et al., 2011; Van Rooyen et al., 2011; Wouters et al., 2008). For example, studies using mice fed high-cholesterol diets have suggested that cholesterol can directly activate HSCs by inducing TLR4, promote oxidative stress and cell death in hepatocytes via excess mitochondrial cholesterol, and promote inflammation in Kupffer cells through lysosomal cholesterol enrichment (Bieghs et al., 2013; Rawson, 2006; Teranati et al., 2012). If increased liver cholesterol is important in TAZ induction in NASH, further studies will be needed to define the mechanisms. Likewise, palmitic acid has been reported to induce pro-inflammatory cytokine production by hepatocytes and Kupffer cells during NASH (Joshi-Barve et al., 2007; Miura et al., 2013), but other mechanisms may be involved as well.

In summary, we show that the Hippo pathway transcription factor TAZ and its gene target Ihh are elevated in the livers of humans with NASH, which is recapitulated in mouse models. In these models, silencing of TAZ or Ihh suppresses key features of NASH progression, but not steatosis. Moreover, silencing TAZ after NASH has developed can partially reverse NASH features, including fibrosis. Although further mechanistic work is needed to fully understand these new findings, the data provide new insight into the pathophysiology of NASH and raise the possibility of liver-directed TAZ inhibition as a new therapeutic strategy to prevent NASH progression.

**EXPERIMENTAL PROCEDURES**

**Animal Studies**

Male wild-type mice C57BL/6J (#000664, 8–10 weeks old) and MC4R-negative loxTB Mc4r mice (#006414, 6 weeks old), referred to here as Mc4r−/− mice, were obtained from Jackson Laboratory and were allowed to adapt to housing in the Columbia University Medical Center Institute of Comparative Medicine for 1 week prior to random assignment to experimental cohorts. Wvtr+/− mice were generously provided by Dr. Eric Olson, University of Texas Southwestern. The mice were then fed the following diets for the times indicated in the figure legends: (1) chow diet (Picolab rodent diet 20, #5053); (2) “fast-food” (FF) diet (TestDiet 1810066): high-fat diet with drinking water containing 42 g/L glucose and fructose (55%/45%, w/w); or (3) fructose-palmitate-cholesterol (“FPC”) diet (Teklad, TD.140154): similar to FF diet but with a 1.25% added cholesterol and with palmitic acid, anhydrous milk fat, and Primex as the sources of fat and with an ~80% decrease in vitamin E and an ~35% decrease in choline compared with typical mouse diets. The detailed composition of these diets appear in Tables S1 and S2. For several experiments, groups of mice were placed on a methionine-choline-deficient diet (MCD Diet, Teklad, TD. 90262) for 8 weeks, as described (Dixon et al., 2012). AAV8-TBG-Cre or control AAV8-TBG-LacZ (1 × 1011 genome copies/mouse) was administered to Wvtr+/− mice by tail vein injection 1 week prior to initiation of the MCD diet. AAV8-H1 shRNA virus or control AAV8-H1 virus (2 × 1011 genome copies/mouse) was delivered by tail vein injection either 1 week prior to FPC diet initiation or after 8 or 16 weeks of the diet. For CCl4-induced liver fibrosis, 6-week-old C57BL/6J mice were injected intraperitoneally (i.p.) biweekly for 6 weeks with 0.5 µL/g body weight of CCl4, which was dissolved in olive oil at a ratio of 1:1 (Mederacke et al., 2013). For the imaging of organs undergoing AAV8-TBG-Cre-mediated recombination, ZsGreen reporter mice (#007906, Jackson Laboratory) (Mu et al., 2015) were injected intravenously (i.v.) with AAV8-TBG-Cre (1 × 1011 genome copies/mouse). 1 week later, the mice were sacrificed, and selected organs were fixed and then imaged using a Leica MZ 16F fluorescence stereomicroscope. Animals were housed in standard cages at 22 °C in a 12 hr light/12 hr dark cycle. All animal experiments were performed in accordance with institutional guidelines and regulations and approved by the Institutional Animal Care and Use Committee at Columbia University.
Figure 7. Ihh Restoration in Taz-Silenced FPC-Fed Mice Promotes NASH Features, and Ihh Silencing Reduces NASH Features

(A–D) The following parameters were measured in male C57BL/6J mice treated with AAV8-H1-shTaz or control vector and then overexpressed Ihh or GFP by AAV8 vector and fed the FPC diet for 16 weeks (*p < 0.05, **p < 0.01, mean ± SEM; n = 6 mice/group): mRNA levels of Wwtr1, Ihh, Col1a1, Timp1, Opn, and Mcp1 (A); quantification of inflammatory cells, Sirius red positive area, F4/80+ cells, and α-SMA+ area (B); TUNEL+ cells (C); and plasma ALT (D).

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Human Samples
Liver biopsy specimens from individuals undergoing weight loss surgery were selected from the MGH NAFLD Biorepository. Patients gave informed consent at the time of recruitment, and their records were anonymized and de-identified. Studies were approved by the Columbia University Institutional Review Board and by the Partners Human Research Committee and conducted in accordance with National Institutes of Health and institutional guidelines for human subject research. Additional anonymized and de-identified liver biopsy sections were obtained from Dr. Jay Lefkowitch, Columbia University Medical Center. Cases with NAFLD activity score of 1–3 were classified as early NAFLD (no fibrosis), while cases with NAS ≥ 5 and fibrosis stage 1a/b–4 were classified as NASH. Cases with steatosis score ≥ 1 and inflammation and ballooning scores of 0 and no fibrosis were classified as steatosis. Cases with NAS 0 were classified as normal.

Blood and Plasma Analyses
Fasting blood glucose was measured using a glucometer (One Touch Ultra, LifeScan) in mice that were fasted for 4–5 hr with free access to water. Complete blood counts were obtained with the FORCYTE Veterinary Hematology Analyzer (Oxford Science). Total plasma triglyceride and cholesterol were assayed using a commercially available kit from Wako. For insulin, MCP1, AST, ALT, TC, and TG are measured following kit instruction by using plasma.

Statistical Analysis
All results are presented as mean ± SEM. p values were calculated using the Statistic Analysis System statistical software package.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Supplemental Experimental Procedures, nine figures, and four tables and can be found with this article online at http://dx.doi.org/10.1016/j.cmet.2016.09.016.

AUTHOR CONTRIBUTIONS
X.W., R.F.S., and I.T. developed the study concept and experimental design; X.W. performed all animal study procedures and most of the in vitro experiments; Z.Z. conducted the CHp assays; J.M.C. conducted the fluorescence-activated cell sorting (FACS) analysis; T.M.H. conducted the liver lipid profiling experiment; and Dr. Peter Nagy, Columbia University, for assisting with the ChIP experiments. This work was supported by NIH grants HL087123 and HL075662 to I.T.; DK078772 to R.T.C.; DK09422 to K.E.C.; and R03DK101863 to J.M.C. Z.Z. is a Russell Berrie Foundation Scholar in Diabetes Research. I.M.S. was supported by a Pinnacle Clinical Research Network (PCRNet) Program grant. This work was supported by NIH Grant Number UL1 TR000040 from the National Center for Advancing Translational Science.

Received: March 17, 2016
Revised: August 6, 2016
Accepted: September 24, 2016
Published: October 27, 2016

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