The Hippo pathway effector TAZ induces TEAD-dependent liver inflammation and tumors

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The Hippo signaling pathway regulates organ size and plays critical roles in maintaining tissue growth, homeostasis, and regeneration. Dysregulated in a wide spectrum of cancers, in mammals, this pathway is regulated by two key effectors, YAP and TAZ, that may functionally overlap. We found that TAZ promoted liver inflammation and tumor development. The expression of TAZ, but not YAP, in human liver tumors positively correlated with the expression of proinflammatory cytokines. Hyperactivated TAZ induced substantial myeloid cell infiltration into the liver and the secretion of proinflammatory cytokines through a TEAD-dependent mechanism. Furthermore, tumors with hyperactivated YAP and TAZ had distinct transcriptional signatures, which included the increased expression of inflammatory cytokines in TAZ-driven tumors. Our study elucidated a previously uncharacterized link between TAZ activity and inflammatory responses that influence tumor development in the liver.

INTRODUCTION

Liver cancer, which includes hepatocellular carcinoma (HCC) and cholangiocarcinoma (CCA), is a major cause of cancer-related deaths and is a substantial unmet medical need (1). Although the number of cases has substantially increased in recent years, the treatment options have hardly changed. Progress in detection and treatment of localized disease has been made, but the 5-year relative survival rate for distant stage liver cancer patients is 3.1%, whereas that of regional stage liver cancer is 10.7% (www.cancer.org). Hence, there is a clear need for new and better treatment options for liver cancer patients.

The Hippo signaling pathway has emerged in recent years as a central regulator of organ size (2, 3). The linear kinase cascade that activates this pathway is highly conserved between Drosophila and mammals. Multiple factors act on the kinases mammalian STE20-like protein kinase 1/2 (MST1/2) and the adaptor salvador 1 (SAV1/WW45). MST kinases phosphorylate Mob1 homolog (MOB1) and large tumor suppressor 1/2 (LATS1/2) kinases. Upon activation, LATS kinases then phosphorylate Yes-associated protein (YAP) and transcriptional coactivator with PDZ-binding motif (TAZ, also known as WWTR1), the key effectors of the Hippo pathway. In Drosophila, where the Hippo pathway also cross-talks with other developmental signaling pathways (4–11) and several novel members of the pathway, and extracellular stimuli provide physiological modulation of the pathway.

Most studies have focused on YAP as a key effector of the Hippo pathway. Whereas Yorkie (YKI) is the only Hippo pathway effector encoded in flies, mammals have two paralogous effector proteins, YAP and TAZ. On the basis of genetic mouse studies, YAP and TAZ appear to have distinct, nonredundant functions during development (12–14). Whether this is due to differences in expression patterns during development or functional distinctions is not well understood. Although both YAP and TAZ are dysregulated in a variety of cancers, whether there are differences in the tumorigenic properties of YAP and TAZ remains unknown. Therefore, it is important to assess biological differences between YAP and TAZ and to understand their differential regulation in development and disease. Given that the liver is exquisitely sensitive to modulation of the Hippo pathway and several studies have reported a role of YAP in liver cancer (15–17), in this study, we investigated whether TAZ has distinct functions in the liver.

Here, we characterized the effects of expressing hyperactivated TAZ in the liver. We found that cooperation of hyperactivated TAZ (TAZ-4SA) and oncogenic NRasV12 in the liver increased cytokine
production, tissue myeloid cell recruitment, and mortality in mice. We further found that these effects required TAZ-TEAD interaction. Our studies highlight that TAZ has a key role in promoting inflammation and development of liver tumors. Furthermore, tumors generated by hyperactivated YAP or TAZ had distinct transcriptional signatures, which suggests that YAP and TAZ may have distinct and combinatorial roles in liver tumor development.

RESULTS
TAZ expression correlates with CCL2 and CXCL1 mRNA amount in tumors
HCC is associated with increased inflammation at all stages of the disease (18). Given the established role of the Hippo pathway in HCC (19), we assessed the gene expression data across a panel of 57 patient-derived xenograft (PDX) HCC models by RNA sequencing (RNA-seq). We found that Taz expression, but not Yap, was among the most positively correlated with C-C motif chemokine ligand 2 (Ccl2, also Mcp1), C-X-C motif chemokine ligand 1 (Cxcl1, also Kc), and interleukin-6 (Il-6) expression (Fig. 1, A and B). Similar results were obtained when examining The Cancer Genome Atlas (TCGA) RNA expression data for the correlation of cytokines (Mcp1 and Kc) with Taz, but not Yap, across multiple cancer types (Fig. 1C). Furthermore, these cytokines correlated better with Taz expression, but not with Yap expression, in a panel of 20 human liver cancer cell lines as well (Fig. 1D). These data suggested that inflammation may be associated specifically with TAZ expression in multiple cancers, including HCC and liver cancer.

Expression of active YAP stimulates tumor formation in mice
Given the differences between the correlation of Yap and Taz expression with inflammatory cytokines, we investigated the effect of YAP and TAZ overexpression on liver tumorigenesis driven by hydrodynamic tail vein (HTV) injection of DNA. HTV injection of DNA constructs with sleeping beauty (SB) transposase allows for the rapid uptake and expression of the construct within the liver (20). In this system, NRasV12 expression alone is not sufficient to drive tumor formation but facilitates hepatocellular transformation in mice (21). Whereas TAZ plays a critical role in fibrosis and liver inflammation (22), YAP is important for macrophage recruitment by tumor-initiating cells (23). However, the differential effects of YAP and TAZ on liver inflammation or tumor growth remain unclear.

Abolishing all the LATS inhibitory phosphorylation sites in vitro results in hyperactivated YAP (24). When we expressed Yap-5SA, which has all five LATS sites mutated with NRasV12 and SB in the mouse liver by HTV injection of plasmids, we found that activated YAP reduced the survival of mice (Fig. 2A). Similarly, coexpression of NRasV12 and constitutively active Yap-S127A also reduced mouse survival (Fig. 2B). Inhibiting the interaction of YAP with the TEAD transcription factors (YAP/S94A mutant) significantly rescued survival of mice that received activated Yap (Fig. 2, A and B). Histological analysis suggested that expression of activated YAP stimulated multifocal expansion of oval cell populations in 75% of the mice (9 of 12; Fig. 2C) and a 33% (4 of 12) incidence of intrahepatic tumor formation (HCC or anaplastic spindle cell tumor/sarcoma; Fig. 2D). In contrast, in mice injected with NRasV12, a hepatocellular adenoma was only identified in 1 of 10 mice around 20 weeks after injection. Activated YAP–associated tumors tended to be anaplastic, although some tumors had features of HCC and CCA. Flow cytometry analysis of immune cells from the liver of control (NRasV12), Yap-5SA, or Yap-5SA/S94A (TEAD binding–deficient) mice revealed no significant differences between myeloid cell infiltrations in this model (Fig. 2, E and F). These data suggest that overexpression of activated YAP stimulates TEAD-dependent tumor development and impaired survival in mice.

Hyperactivated TAZ stimulates proinflammatory myeloid cell infiltration of tissues
Because abolishing all the LATS inhibitory phosphorylation sites in vitro results in hyperactivated TAZ (24), we also mutated all four LATS phosphosites from serine to alanine (TAZ-4SA) to generate constitutively active Taz constructs. When we injected mice with Taz-4SA and NRasV12 plasmids, we found that they all died within 1 week (Fig. 2G). This rapid mortality was observed on two different genetic backgrounds (FVB and C57BL/6; Fig. 2G). In contrast, we found no effect on mortality when we injected Taz-4SA alone (Fig. 3A).

Death was associated with enlarged livers with morphological features of inflammation in the TAZ-4SA mice (Figs. 2H and 3B and fig. S1). We noted that TAZ abundance was increased in the livers as early as 2 days after injection (Fig. 2I and fig. S2) and that some liver morphology changes occurred when TAZ-4SA was expressed alone (Fig. 3B). Morphologically altered cells were pancytokeratin AE1/AE3–positive and von Willebrand factor (VWF)–negative (Fig. 3C and fig. S3A), suggesting that these cells had an epithelial origin. We also found strong F4/80 (Fig. 3C) and CD68 (fig. S3B) immunolabeling in liver sections of mice that received coinjection of Taz-4SA and NRasV12. Flow cytometry analysis of liver immune infiltrates indicated that overexpression of activated TAZ increased the frequency and number of CD11b+/F4/80+Gr1– macrophages in this tissue (Fig. 3, D and E). Increased inflammatory cells were also found in multiple organs, most notably in the spleen and lung (fig. S4).

To evaluate whether the massive myeloid infiltration in mice co-injected with Taz-4SA and NRasV12 caused the increased mortality that we observed, we depleted myeloid cells by either a single dose of 10-gray (Gy) irradiation or injection with clodronate liposomes. Neither method of myeloid cell depletion rescued the lethality seen after coinjection with Taz-4SA and NRasV12 in mice (fig. S5A). After myeloid depletion, we found that there was no change in the liver dysplasia associated with overexpression of activated TAZ and NRasV12, although inflammatory infiltrates were reduced (fig. S5B). These results indicated that hepatocyte morphology changes in these mice were not dependent on myeloid infiltration.

When we evaluated the serum cytokine profile of mice after HTV injection, we found that circulating proinflammatory cytokines were altered in mice injected with either Taz-4SA alone or both Taz-4SA and NRasV12 (Fig. 3F). Gene expression analyses on FACs (fluorescence-activated cell sorting)–sorted macrophages revealed that expression of genes involved in inflammation and endoplasmic reticulum stress was increased in mice co-injected with Taz-4SA and NRasV12 (Fig. 3G). Transcript expression of Il-6, Stat3 (signal transducer and activator of transcription 3), and Atf4/Eif2A (activating transcription factor 4/eukaryotic translational initiation factor 2A) all increased in macrophages from these mice. Together, these data demonstrated that hyperactivated TAZ increased proinflammatory cytokine production and myeloid cell recruitment. This is strikingly similar to what we observed in the PDX HCC models and across multiple TCGA indications.
Fig. 1. TAZ expression correlates with CCL2 and CXCL1 expression in tumors. (A and B) Pearson correlation between Taz (A) or Yap (B) expression and chemokine and cytokine mRNAs in RNA-seq data from PDX HCC models. Data for the colored genes are means ± SD from 57 tumor models. (C) Pearson correlation between Yap or Taz expression and Ccl2, Il-6, and Cxcl1 mRNAs in TCGA RNA-seq data sets. Boxplot data are median and quartile correlation values between Taz and Yap with the indicated genes within 22 TCGA indications. (D) Correlation between Yap and Taz expression with inflammatory cytokines. Boxplot data are median and quartile values [left, P value: not significant (ns)] and scatterplot data are mean values (right) from 20 liver cancer cell lines. *P < 0.05 by unpaired t test (B and C), **P < 0.01, ****P < 0.0001 (C), Wilcoxon rank sum test (D, left), or Spearman rank correlation (D, right).
3.24 at their interaction site (TAZ-4SA/S51A), we found that this abolished the TAZ and TEAD interaction by generating a second mutant (Fig. 4). When we inhibited the TEAD family of transcription factors, and the TAZ S51A mutation reduced mortality was accompanied by reduced inflammatory infiltration of the liver and grossly normal liver morphology (Fig. 4B). Similarly, the frequency and number of macrophages in the liver determined by flow cytometry analysis depended on the TAZ-TEAD interaction (Fig. 4, C and D). Mutation of TAZ S51A also reduced proinflammatory serum cytokines when compared to coinjection of Taz-4SA and NRasV12 (Fig. 4E). Together, these data demonstrated that TAZ promoted inflammation in a TEAD-dependent manner (fig. S6).

Reducing TAZ expression or activity stimulates TEAD-dependent tumorigenesis
To evaluate the dose-dependent effect of TAZ expression on liver tumorigenesis, we coinjected mice with a lower dose of Taz-4SA and NRasV12 DNA. When mice were injected with 25% of the Taz-4SA and NRasV12 DNA, we found that they survived significantly longer than mice that received a normal dose of Taz-4SA and NRasV12 (Fig. 5A). After injection with this dose of DNA, histological analysis indicated that about 25% of mice developed tumors (Fig. 5, B and E). Tumors in these mice included one HCC and two anaplastic spindle cell tumors (fig. S8). Histological analysis found that these mice still exhibited multifocal inflammation and liver dysplasia, which was consistent with the increased amount of CCL2 found in the serum when compared to control mice (Fig. 5C). These data indicated that, at lower doses that did not stimulate rapid mortality, activated TAZ promoted liver transformation.

We tested whether reduced TAZ activation would similarly stimulate tumor formation by evaluating a second TAZ construct with only a single phosphosite mutation. Abolishing the LATS inhibitory phosphorylation site at serine 89 on TAZ in vitro activates TAZ (24, 25). Coexpression of NRasV12 with a constitutively active mutation at a single site in TAZ (TAZ-S89A, equivalent to YAP-S127A) reduced survival relative to wild-type mice or mice injected with NRasV12 alone and drove liver tumorigenesis with nearly 37.5% penetrance (Fig. 5, D and E). Similar to YAP, inhibiting the interaction of TAZ with the TEAD transcription factors (TAZ S51A mutant) significantly improved survival and reduced tumor incidence with only a single mouse developing a small, discrete hepatocellular adenoma (Fig. 5, D and E). When we profiled the cytokines secreted by a tumor

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TAZ regulates liver inflammation through TEAD-dependent transcription
YAP and TAZ function as coactivators through interaction with the TEAD family of transcription factors, and the TAZ S51A mutation abolishes the interaction of TAZ with TEAD (24). When we inhibited the TAZ and TEAD interaction by generating a second mutation at their interaction site (TAZ-4SA/S51A), we found that this mutation completely rescued the lethality induced by coinjection of Taz-4SA and NRasV12 (Fig. 4A). When mice were injected with 25% of the Taz-4SA and NRasV12 DNA, we found that they survived significantly longer than mice that received a normal dose of Taz-4SA and NRasV12 (Fig. 5A). After injection with this dose of DNA, histological analysis indicated that about 25% of mice developed tumors (Fig. 5, B and E). Tumors in these mice included one HCC and two anaplastic spindle cell tumors (fig. S8). Histological analysis found that these mice still exhibited multifocal inflammation and liver dysplasia, which was consistent with the increased amount of CCL2 found in the serum when compared to control mice (Fig. 5C). These data indicated that, at lower doses that did not stimulate rapid mortality, activated TAZ promoted liver transformation.

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Fig. 3. Hyperactivated TAZ-4SA mutant cooperates with NRasV12 to induce lethality in mice. (A) Survival analysis of mice injected with the indicated TAZ and NRasV12 constructs. Data are from at least four mice per group analyzed in three independent experiments. (B and C) Histology analysis (B) and immunohistochemistry analysis of cytokeratin AE1/AE3 and F4/80 (C) in liver sections from mice injected with indicated constructs. Scale bars, 100 μm. Histology images are representative of at least three mice per group harvested on day 6 after injection. (D and E) Flow cytometry analysis of myeloid cells in the liver of mice injected with the indicated constructs. Dot plots (D) are representative of at least three mice per group from three independent experiments. Quantified total white blood cells (WBCs) and myeloid cells in the liver (E) are means ± SEM pooled from all experiments. (F) Luminex analysis of serum cytokines in mice injected with the indicated constructs. Data are z-scores of means from three to five mice per group, representative of three independent experiments. IFN-γ, interferon-γ; TNF-α, tumor necrosis factor-α; GM-CSF, granulocyte macrophage colony stimulating factor; G-CSF, granulocyte colony stimulating factor; MIP-1α, macrophage inflammatory protein–1α; RANTES, regulated on activation normal T cell expressed and secreted. (G) Fluidigm reverse transcription polymerase chain reaction (RT-PCR) analysis of sorted macrophages from the livers of mice injected with indicated constructs. Data are z-scores of means from three to five mice per group. Rows have been hierarchically clustered using Pearson correlation as a distance metric. *P < 0.05 by log rank test (A) or two-tailed unpaired Student’s t test (E).
cell line derived from a TAZ-S89A tumor by Luminex analysis, we found that these cells secreted more CXCL1 and CCL2 than the remaining cytokines evaluated (Fig. 5F). Together, these data demonstrated that limited TAZ activity is sufficient to stimulate liver transformation and tumors that secrete inflammatory cytokines.

YAP and TAZ drive differential gene expression in liver tumor cells
We performed whole transcriptome RNA-seq on reestablished cell lines from murine liver tumors formed after coinjection of Yap-S127A, Yap-5SA, or Taz-S89A with NRasV12 to further dissect the independent effects of TAZ and YAP. Because of the massive
myeloid cell infiltration in the livers of mice coinjected with TAZ-4SA and NRasV12, but not Yap-5SA and NRasV12, we could not directly compare gene expression in liver samples ex vivo. Instead, we compared transcript expression in cell lines derived from YAP- or TAZ-driven tumors (fig. S7) and found substantial differences. Unbiased clustering of differentially expressed genes revealed four distinct gene clusters that differed between YAP-S127A–, YAP-5SA–, and TAZ-S89A–transformed cells (Fig. 6A and fig. S9). We noted that, in cluster 2 (Fig. 6A), genes had the weakest expression in YAP-S127A–driven tumor cells, followed by YAP-5SA, and the strongest expression in TAZ-S89A cells. Several immune-related genes were among those with the largest fold change in mRNA expression in TAZ-S89A–derived tumor cell lines when compared to YAP-S127A or YAP-5SA cell lines (Fig. 6B). These immune-related genes included the proinflammatory cytokines Cxcl5, Cxcl1, and Ccl2 that were strongly expressed specifically in TAZ-S89A cells (Fig. 6, B and C). This mutant-specific gene expression pattern was validated by quantitative RT-PCR (Fig. 6, D and E). Furthermore, chromatin immunoprecipitation–qPCR analyses identified that, in cell lines derived from both YAP-S127A– and TAZ-S89A–driven tumors, histone H3 was acetylated on K27 at the Ccl2 promoter, indicative of active transcription (Fig. 6F). These data suggest that both YAP and TAZ bind to the Ccl2 promoter. Together, these data suggest that YAP and TAZ differentially regulate expression of many genes in liver tumors, including immune-related genes. Specifically, differential expression of several proinflammatory cytokines was mutation-specific, suggesting overlapping but distinct roles for YAP and TAZ.

**DISCUSSION**

Our study supports a role for TAZ in liver inflammation and cancer and highlights the idea that YAP and TAZ have different roles in these processes. Because HTV injection offers a reliable, quick, and cost-effective tool to generate preclinical murine models of liver cancer [reviewed in (20)], we used this model to compare the roles of YAP and TAZ in the liver. Whereas overexpression of hyperactivated TAZ drove lethal inflammatory disease and TAZ had an important role in inflammatory cytokine production, YAP stimulated tumor formation. Both effects required interaction with TEAD transcription factors, suggesting that the effects of YAP and TAZ are transcriptionally regulated. Consistent with evaluations of PDX HCC models and the TCGA data analyses, TAZ abundance and activity correlated with the proinflammatory cytokines CCL2 and CXCL1 in both liver inflammation and liver tumorigenesis.

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... underlying mechanism remains unknown. What role TAZ may have in any of these contexts and the differential roles of YAP and TAZ in inflammation and tumorigenesis are largely unexplored. Here, we found that TAZ promotes inflammatory cytokine production and tissue macrophage infiltration. Given the links between inflammation and cancer development (29), it is tempting to speculate that TAZ induces inflammation that may promote the oncogenic function of YAP in the liver. This may provide therapeutic opportunities for intervention by targeting both key Hippo signaling and inflammatory pathways in liver cancer. This study highlights the idea that YAP and TAZ have distinct roles in liver tumor development.

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questions of interest in the field (30, 31). The presence of Gram-positive bacteria is a unique signal that activates the Hippo pathway in flies through the Toll-Myd88-Pelle cascade (6). However, whether this function is conserved in mammals remains unknown. Because flies only encode the gene Yki, we speculate that the functions of the mammalian orthologs may have separable functions. Whereas both YAP and TAZ promoted liver tumors, TAZ played a more prominent role in secretion of proinflammatory cytokines and driving inflammation-induced tumors. TAZ is also required for the differentiation of T helper 17 cells (32). In prostate cancer, YAP stimulates expression of the chemokine CXCL5, which recruits myeloid-derived suppressor cells to promote tumor growth (33). YAP activation also underlies macrophage recruitment by tumor-initiating cells (23). Although these studies do not compare the role of TAZ in their specific model systems, collectively, all our data suggest that YAP and TAZ are not completely redundant. It is possible that there are overlapping yet distinct roles of YAP and TAZ in a context- and tissue-dependent manner. Although YAP could play a more important role in tumor initiation, YAP and TAZ could promote inflammation and stimulate transcription of distinct target genes depending on their expression in specific cell types. Future studies will address whether this is broadly applicable in other cancers and provide additional targets for therapeutic intervention.

MATERIALS AND METHODS
Gene expression profiling—PDX and TCGA
Liver cancer PDX RNA-seq data were obtained from OncoWuXi (https://onco.wuxiapptec.com). RNA-seq alignment was performed using the RNA STAR software (v2.4.0). For Homo sapiens samples, hg19 genome build with gencode v19 gene annotation was used. Raw read numbers of each gene count were used counting featureCounts (v1.4.6). Fragments per Kilobase Million values are calculated with R package edgeR (v3.8.5). For TCGA data, raw RNA-seq data were downloaded from TCGA (n = 7304) and aligned to the human reference genome (GRCh37/hg19) using GSNAP version 2013-10-10, allowing a maximum of two mismatches (parameters: “-M 2 -n 10 -B 2 -i 1-N 1 -w 200000 -E 1 --pairmax-rna=200000”). Gene expression values were obtained from a previously published study (21). For Fig. 1D, all correlations calculated for RNA-seq data were performed using the DESeq2 package for the R programming language. All correlations calculated for RNA-seq data are Pearson’s correlation. Statistical significance between correlation values was determined by nonpaired two-sided t test. For Fig. 1D, all correlations calculated for RNA-seq data were performed in R using the cor function (with the method being either Pearson or Spearman as annotated in axes). Cell line cytokine, Yap, and Taz RNA expression values were obtained from a previously published study (34).

Animals
FVB-N (Charles River) or C57BL/6 mice were subjected to HTV infection with SB transposase and relevant TAZ mutant encoding plasmids, as previously described (21). For myeloid depletion studies, mice were injected with clodronate or control liposomes as per the manufacturer’s instructions (Encapsula NanoSciences). For irradiation studies, mice were given one round of 10-Gy irradiation. The Genentech Institutional Animal Care and Use Committee approved all protocols.

Cytokine profiling
Serum was separated from blood that was harvested by orbital bleed or terminal heart bleed using BD microtainer tubes with serum separator additive (Becton Dickinson). Samples were diluted 1:4 in sample diluent (Bio-Rad) and were used for Bio-Plex Pro Mouse Assay (Bio-Rad) with sample dilutions of 1:4 and 1:20 according to the manufacturer’s instructions. Heatmap visualizations were row-normalized by z-scoring.

Liver processing
All the lobes of the liver were collected in ice-cold sterile phosphate-buffered saline (PBS) after perfusion. Briefly, the lobes were minced with a blade and were incubated in digestion medium [RPMI 1640 medium containing deoxyribonuclease I (DNase I; 0.1 mg/ml; Invitrogen), collagenase P (0.4 mg/ml; Roche), and dispase (0.8 mg/ml; Roche) in a 37°C water bath], and cell fractions were collected every 10 to 15 min over a period of 45 min. The single-cell suspensions were filtered and ACK (ammonium-chloride-potassium)–lysed for a period of 5 min at room temperature, followed by neutralization with PBS. The cells were further stained for antibodies for flow cytometry/sorting.

Fluidigm gene expression analysis
RNA (100 ng) was subjected to complementary DNA (cDNA) synthesis reaction using the Applied Biosystems High Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific), as per the manufacturer’s protocol. Preamplification reaction was then performed with Taqman PreAmp Master Mix (Thermo Fisher Scientific). After amplification, samples were diluted 1:4 with tris-EDTA, and qPCR was conducted on Fluidigm 96.96 Dynamic Arrays using the BioMark HD system according to the manufacturer’s instruction. Fluidigm data were analyzed with RealTime StatMiner in SpotFire program. Heatmap visualizations were row-normalized by z-scoring.

Flow cytometry and antibodies
After mechanical homogenization or enzymatic digestion of liver, cells were resuspended in flow cytometry buffer (PBS containing 2% fetal bovine serum and 2 mM EDTA) and incubated for 30 min with the following dyes and antibodies: CD11b (M1/70; BD Pharmingen), GR1 (RB6-8C5; BD Pharmingen), Ly6C (HK1.4; BioLegend), CD45 (30-F11; BioLegend), Ly6g (IAA8; Bioscience), F4/80 (BM8; BioLegend), CD11C (N418; ebioscience), propidium iodide nucleic acid stain (Molecular Probes), and Viability Dye (53-67; BioLegend). The stained cells were acquired on a FACSAria (BD Biosciences) and analyzed using FlowJo software (Tree Star Inc.). The F4/80+ cells were sorted using FACSaria (BD Biosciences), and the cells were collected in RNA protect buffer (Qiagen) for downstream RNA purification and Fluidigm analysis.

Western blotting
Whole cell lysates were prepared in NP-40 buffer [1% NP-40, 120 mM NaCl, 50 mM tris (pH 7.4), 1 mM EDTA (pH 7.4), 20 mM N-ethylmaleimide, and protease and phosphatase inhibitors (Roche)]. All antibodies were used from Cell Signaling Technology.

Real-time RT-PCR
Cells were lysed, and RNA was isolated using the RNeasy Plus Mini Kit (Qiagen), according to the manufacturer’s protocol. Upon cDNA production using the iScript cDNA Synthesis Kit (Bio-Rad), samples were analyzed by real-time PCR using a QuantStudio 7 Flex machine (Applied Biosystems). All probes used were either purchased from Applied Biosystems [including Ccl2 (Mm00441242_m1), Cxcl1 (Mm04207460_m1), Ccl5 (Mm01302427_m1), and Cxcl10 (Mm00445235_m1)] or
custom-designed [including Cxcl1, AAA AGG TGT CCC CAA GTA (forward) and AAG CAG TGA ACT ACC ATC G (reverse); and β-actin, GGA ACT GTG CGT GAC ATC AAA G (forward) and TGT AGT TTC ATG GAT GCC ACA G (reverse)].

### Histology

Histologic evaluations were performed on about 4-μm-thick formalin-fixed, paraffin-embedded tissue sections routinely stained with hematoxylin and eosin. Immunohistochemistry was also performed on 4-μm-thick formalin-fixed, paraffin-embedded tissue sections using primary antibodies against CD68 (Abcam; rabbit polyclonal, 0.25 μg/ml), cytokeratin AE1/AE3 (Dako; mouse monoclonal, 5 μg/ml), F4/80 (AbD Serotec; rat monoclonal, clone Cl:A3-1, 10 μg/ml), and VWF (Dako; rabbit polyclonal, 3.875 μg/ml). CD68 immuno-histochemistry was performed on the Ventana Discovery XT platform using CC1 standard antigen retrieval and the Rabbit OmniMap custom-prefilter was applied: Only genes with at least a median RPKM of 0.25 were transformed to log2 counts per million, quantile-normalized, and precision-weighted with the “voom” function of the limma package (38). Genes statistically significantly up-regulated in TAZ-S89A were determined by genes that were statistically significantly up-regulated in TAZ-S89A when compared against YAP-S127A as well as when combining YAP-S127A and YAP-5SA samples. Four clusters were determined by manual inspection. Fuzzy c-means clustering was performed using R package e1071 v.1.6.8.

### RNA library preparation and sequencing

RNA-seq was performed to identify genes that are differentially expressed between the YAP and TAZ mutants and to understand their contribution toward liver cancer. Total RNA was extracted using the Qiagen RNeasy kit as per the manufacturer’s protocol including the on-column DNase digestion. Quality control of samples was done to determine RNA quantity and quality before their processing by RNA-seq. The concentration of total RNA samples was determined using NanoDrop 8000 (Thermo Fisher Scientific). The integrity of RNA samples was determined using 2100 Bioanalyzer (Agilent Technologies). About 500 ng of total RNA was used as an input for library preparation using the TruSeq RNA Sample Preparation Kit v2 (Illumina). Size of the libraries was confirmed using 2200 TapeStation and High Sensitivity D1K screen tape (Agilent Technologies), and their concentration was determined by a qPCR-based method using the Library Quantification Kit (KAPA). The libraries were multiplexed and then sequenced on Illumina HiSeq2500 (Illumina) to generate 30M of single-end 50–base pair reads.

### RNA-seq alignment and feature counting

RNA-seq reads were first aligned to ribosomal RNA sequences to remove ribosomal reads. The remaining reads were aligned to the mouse reference genome (GRCm38) using GSNAP (35, 36) version “2013-10-10,” allowing a maximum of two mismatches per 75 base sequence (parameters: -eM 2 -n 10 -B 2 -i 1 -N 1 -w 200000 -E 1 --pairmax-rna=200000 --clip-overlap). Transcript annotation was based on the Ensembl genes database (release 77). To quantify gene expression levels, the number of reads mapped to the exons of each RefSeq gene was calculated.

### RNA-seq differential gene expression

Differential expression was performed with DESeq2 (37). A prefilter was applied: Only genes with at least a median RPKM (reads per kilobase per million mapped reads) value of 10 in one condition were analyzed. P values for other genes were simply set to 1 and log fold changes were set to 0 for visualization purposes, but such genes were not included in the multiple testing correction. Q values were obtained by correcting P values for multiple hypotheses using the Benjamini-Hochberg procedure. Genes were considered if they had a Q value of less than 0.05 and were protein-coding. Counts were transformed to log2 counts per million, quantile-normalized, and precision-weighted with the “voom” function of the limma package (38). Genes statistically significantly up-regulated in TAZ-S89A were determined by genes that were statistically significantly up-regulated in TAZ-S89A when compared against YAP-S127A as well as when combining YAP-S127A and YAP-5SA samples. Four clusters were determined by manual inspection. Fuzzy c-means clustering was performed using R package e1071 v.1.6.8.

### REFERENCES AND NOTES


carcinogenesis.


The Hippo pathway effector TAZ induces TEAD-dependent liver inflammation and tumors

The Hippo pathway effectors YAP and TAZ are overexpressed in various cancers. Loss of upstream kinases that inhibit the activity of these transcriptional coactivators promotes inflammation. In patient-derived xenografts and TCGA data sets, Hagenbeek et al. found that only TAZ expression correlated strongly with inflammatory cytokine transcript abundance. Expression of hyperactivated TAZ, but not YAP, in the livers of mice augmented transcription factor TEAD-mediated systemic inflammation and tissue infiltration by myeloid cells. RNA-seq analysis identified distinct gene signatures in tumor cells driven by activated YAP or TAZ, suggesting that these Hippo pathway effectors have nonredundant functions.

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