Two-signal requirement for growth-promoting function of Yap in hepatocytes

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Abstract The transcriptional coactivator Yes-associated protein (Yap) promotes proliferation and inhibits apoptosis, suggesting that Yap functions as an oncogene. Most oncogenes, however, require a combination of at least two signals to promote proliferation. In this study, we present evidence that Yap activation is insufficient to promote growth in the otherwise normal tissue. Using a mosaic mouse model, we demonstrate that Yap overexpression in a fraction of hepatocytes does not lead to their clonal expansion, as proliferation is counterbalanced by increased apoptosis. To shift the activity of Yap towards growth, a second signal provided by tissue damage or inflammation is required. In response to liver injury, Yap drives clonal expansion, suppresses hepatocyte differentiation, and promotes a progenitor phenotype. These results suggest that Yap activation is insufficient to promote growth in the absence of a second signal thus coordinating tissue homeostasis and repair.

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Introduction

The transcriptional coactivator Yap is an evolutionarily conserved mediator of cell fate decisions such as proliferation, differentiation, and survival (Yu and Guan, 2013). Together with a related protein Taz, Yap is the central target of the Hippo pathway, a growth-suppressive network of kinase complexes that inactivates Yap and Taz (Hong and Guan, 2012). The Hippo pathway has been implicated in organ size control, as overexpression of Yap or inactivation of the Hippo components leads to tissue overgrowth and organomegaly in Drosophila and mouse models (Camargo et al., 2007; Dong et al., 2007; Lin et al., 2013). These effects are largely mediated by Yap-TEAD complexes that activate transcription of genes promoting cell proliferation and inhibiting apoptosis (Wu et al., 2008; Zhang et al., 2008; Zhao et al., 2008). Yap is interconnected with RTK (Reddy and Irvine, 2013), GPCR (Yu et al., 2012), PI3K (Fan et al., 2013), Wnt (Bernascone and Martin-Belmonte, 2013), and TGF-beta (Ferrigno et al., 2002; Attisano and Wrana, 2013; Mullen, 2014) signaling, and Yap co-regulates transcription by interacting with Smads (Ferrigno et al., 2002; Beyer et al., 2013), TCF/LEF (Konsavage and Yochum, 2013), Tbx5 (Beyer et al., 2013), Runx2 (Zaidi et al., 2004), FoxO1 (Shao et al., 2014), and p73 (Strano et al., 2001), among others (Barry and Camargo, 2013). The intestinal epithelium in both Drosophila (Karpowicz et al., 2010; Ren et al., 2010; Shaw et al., 2010; Staley and Irvine, 2010) and mice does not depend on Yap for homeostatic tissue turnover (Zhou et al., 2011), but it does respond very strongly to Yap overexpression (Barry et al., 2013) and requires Yap for tissue repair (Cai et al., 2010). Thus, in some tissues Yap may be dispensable for...
eLife digest As we grow up, the organs in our body tend to stop growing and then remain roughly the same size for the rest of our lives. This is possible because of control systems that determine how often the cells within the organ can divide and when they should die. If these controls fail, the cells may divide rapidly and not die when they should, which can cause tumors to grow. In healthy cells, the proteins that promote cell division and/or prevent cell death are strictly controlled—usually by at least two different ‘on’ signals—so that they are only active at specific times.

Yap is one such protein that promotes cell division and inhibits programmed cell death. Previous studies have reported that artificially increasing the levels of Yap in cells is sufficient to make tumors grow in a seemingly unrestrained fashion. This suggests that only one signal is required to over-activate Yap. However, it is also possible that these experiments were carried out under conditions where a second unknown growth-promoting signal was also present.

Here, Su, Bondar et al. genetically altered mice to produce more Yap in some, but not all, cells in the liver. This revealed that when surrounded by normal cells, the high levels of Yap in individual cells do not lead to excessive liver growth. The cells do divide more, but this is balanced by an increase in the numbers of cells that die.

However, if the liver is injured, the high levels of Yap can keep the cells in a stem cell-like state and cause them to grow and divide excessively. This helps the liver to recover, and once the recovery is complete, the cells that produce more Yap are killed. Su, Bondar et al. suggest that the excessive cell growth observed in previous studies may be due to the researchers unintentionally simulating the conditions of inflammation by increasing the levels of Yap in all the cells.

These findings reveal that Yap needs to be activated by both a signal from within the cell and a second signal from other cells—caused by injury or inflammation—to promote the growth of tumors.

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from *Drosophila* (*Chen et al., 2012*) and mammalian cell culture (*Norman et al., 2012*) suggest that the outcome of Yap activation depends on Yap activity in neighboring cells. The regulation of Yap activity by such mechanism has not been characterized in mammalian tissues in vivo.

Here, we describe a mosaic model of Yap activation in the mouse liver, where a high level of Yap expression is induced in a fraction of hepatocytes surrounded by normal tissue. We present evidence that a high level of Yap in this context is insufficient to drive clonal expansion, as increased proliferation of Yap-overexpressing hepatocytes is balanced by their increased susceptibility to apoptosis. Yap-mediated clonal expansion requires a second signal provided by disruption of tissue homeostasis, such as injury or inflammation. Upon returning to homeostasis, excessive Yap-overexpressing cells are eliminated by apoptosis.

### Results

#### Generation of mosaic Yap mouse model

To study the role of cell environment in the regulation of the mammalian Hippo pathway, we generated a conditional mosaic mouse model of Yap overexpression (*YapKl* mice). We used *Yap1* mutant S112A which has enhanced nuclear localization and has been previously shown to cause a several-fold increase in liver size when overexpressed (*Schlegelmilch et al., 2011; Zhang et al., 2011*) in the entire organ. A targeting construct driving expression of the *Yap1* S112A-IRES-GFP downstream of a floxed transcriptional STOP cassette was knocked into the Gt(ROSA)26Sor (*Rosa26*) locus. Upon expression of Cre recombinase, the transcriptional STOP cassette is removed, allowing expression of Yap together with GFP (*YapKl* mice; *Figure 1—figure supplement 1*).

We used the *Rosa26*-IRE-S-GFP targeting construct which is expressed bimodally (*Bondar and Medzhitov, 2010*) (likely due to epigenetic regulation of the locus), generating GFP\textsuperscript{low} and GFP\textsuperscript{high} populations in all cells of hematopoietic lineages (*Bondar and Medzhitov, 2010*) and solid tissues tested (liver, pancreas, muscle, intestine; *Figure 1—figure supplement 2* and 2 data not shown). Both populations have a recombined STOP cassette, but the gene of interest is expressed above physiological level only in GFP\textsuperscript{high} cells. GFP\textsuperscript{low} cells serve as an internal control population of the identical genetic background for the GFP\textsuperscript{high} cells. In the absence of selective pressure, the ratio of GFP\textsuperscript{low} to GFP\textsuperscript{high} cells remains constant throughout life. Thus, crossing *YapKl* mice to *Albumin-Cre* (*Alb-Cre*) yields mosaic Yap expression in the hepatic lineage, where individual cells overexpressing Yap can be traced by GFP fluorescence (*YapKl*\textsuperscript{Ab-Cre} mice). Similar to the previously reported *Rosa26*-DNp53 mice (*Bondar and Medzhitov, 2010*), the proportion of GFP\textsuperscript{low} to GFP\textsuperscript{high} cells remained stable in the blood of *YapKl*\textsuperscript{CreER} mice upon tamoxifen induction for many months (*Figure 1—figure supplement 2B–D*). In the colons of *YapKl*\textsuperscript{Ab-Cre} mice, crypts were either entirely GFP\textsuperscript{high} or GFP\textsuperscript{low} (GFP\textsuperscript{low} appearing as GFP− due to limited sensitivity of GFP detection by tissue immunostaining). There were no crypts that have mixed populations, as would be expected if cells randomly switched expression level (*Figure 1—figure supplement 2E*). Collectively, these data illustrate stable inheritance of the *Rosa26* allele expression level.

First, we verified the correlation of Yap and GFP expression by immunofluorescence staining of the *YapKl*\textsuperscript{Ab-Cre} liver sections. Indeed, Yap and GFP in hepatocytes displayed mosaic and overlapping expression pattern, with cytosolic as well as nuclear Yap localization (*Figure 1A*). Utilizing the GFP marker, we tested the recombination efficiency by flow cytometry. More than 90% of hepatocytes were GFP-positive, indicating high deletion efficiency of the STOP cassette. Expression of Cre recombinase, the transcriptional STOP cassette is removed, allowing expression of Yap together with GFP (*YapKl* mice; *Figure 1—figure supplement 1*). Recombination was also confirmed by genomic qPCR (*Figure 1—figure supplement 3*). The hepatocytes formed two distinct populations differing in GFP brightness. To determine the level of Yap activity in these cells, we sorted each of these populations by FACS. Since GFP fluorescence varied 10-fold within the GFP\textsuperscript{low} population, we further subdivided it into GFP\textsuperscript{low} and GFP\textsuperscript{med-low} populations (*Figure 1B*). The exogenous transcripts of Yap are highly expressed only in the GFP\textsuperscript{high} population, but not in the GFP\textsuperscript{low} and GFP\textsuperscript{med-low} populations. The small amount of the exogenous Yap mRNA in the GFP\textsuperscript{low} and GFP\textsuperscript{med-low} populations is insignificant due to a potential negative feedback mechanism that reduces the endogenous Yap expression, resulting in a similar level of total Yap mRNA compared to that of wild-type (WT) (*Figure 1—figure supplement 4*). Additionally, CTGF, a known Yap downstream target gene (*Zhao et al., 2008*), was significantly upregulated only in the GFP\textsuperscript{high} population, whereas its expression remained unchanged in GFP\textsuperscript{low} and GFP\textsuperscript{med-low} populations, indicating normal levels of Yap activity in these cells (*Figure 1C*). Thus, GFP and Yap protein levels correlate in *YapKl*\textsuperscript{Ab-Cre} mice, and Yap is overexpressed and has higher activity specifically in the GFP\textsuperscript{high} hepatocytes (to which we will refer as Yap\textsuperscript{high} cells).
Yap activation in hepatocytes does not lead to clonal expansion at steady state

The first question we addressed is whether liver size is affected by mosaic Yap overexpression. Previous studies showed that hyperactivation of Yap in the entire liver leads to rapid and progressive hepatomegaly: liver weight increases fivefold in 1 month in Yap transgenic mice (Camargo et al., 2007; Dong et al., 2007) and fourfold by 3 months of age in Stk3, Stk4 (Mst1/2) double null mice (Lu et al., 2010; Song et al., 2010) (where Yap is activated due to an absence of upstream negative regulators). However, the liver/body weight ratio of the mosaic YapKI Alb-Cre mice was only slightly elevated (Figure 2A). Hepatocyte numbers (Figure 2B) and liver morphology (Figure 2C) also remained normal. Moreover, Yaphigh cells did not have a growth advantage over the WT neighbors, since neither the average size of Yap high cells clusters in the liver sections (Figure 2—figure supplement 1) nor the average frequency of Yap high hepatocytes increased over time (Figure 2D).

These results were unexpected, considering the dramatic increase in hepatocyte proliferation and survival driven by Yap in the previously published whole-liver transgenic models (Camargo et al., 2007; Dong et al., 2007). We therefore tested whether Yap influenced proliferation and apoptosis in YapKIAlb-Cre liver. Indeed, more Yaphigh hepatocytes incorporated BrdU than their GFPlow neighbors or hepatocytes in WT livers (Figure 2E). Meanwhile, TUNEL staining indicated more apoptosis in the YapKI Alb-Cre livers (Figure 2F). It is noteworthy that only a minor fraction of the Yaphigh cells proliferated, and over 95% remained quiescent throughout the 3-day BrdU labeling period. These data suggest that increased Yap levels sensitize hepatocytes to both mitogenic and apoptotic signals but are insufficient to drive cell expansion, resulting in slightly increased cell turnover without net growth advantage.
Figure 2. Yap overexpression in hepatocytes does not induce hepatomegaly or clonal expansion at steady state. (A) Liver/body weight ratios of 1- to 3-month-old YapKIAlb-Cre mice (KI) and littermate controls (WT), n ≥ 11. ****p ≤ 0.0001. (B) Total hepatocyte numbers were determined by quantitative flow cytometry of primary hepatocytes isolated from 1- to 3-month-old mice of the indicated genotypes, n ≥ 14. (C) A representative image of H&E staining performed on liver sections of 6-week-old control (WT) and YapKIAlb-Cre mice (KI). (D) Primary hepatocytes were
This insufficiency was not due to an overall lack of Yap activation, as Yap overexpression induced a significant change in gene expression in our model. Sorted WT, GFP<sup>low</sup> and GFP<sup>high</sup> hepatocytes were compared by RNA-sequencing analysis. Gene expression of GFP<sup>low</sup> cells isolated from YapKI<sup>Ab-Cre</sup> livers was similar to that of the WT hepatocytes, whereas GFP<sup>high</sup> hepatocytes showed robust transcriptional changes: 821 genes were significantly up-regulated (p ≤ 0.005 and fold change ≥2) in Yap<sup>high</sup> hepatocytes compared to WT hepatocytes (Figure 2—figure supplement 2A). 37% of these genes overlapped with genes upregulated in the previously characterized Yap overexpression model (Dong et al., 2007) (Figure 2—figure supplements 2B, 3A–B), further validating our model. Dong et al (Dong et al., 2007) used an ApoE promoter which is not restricted to hepatocytes, and assayed the whole liver that includes stromal, endothelial and immune cells with microarray, whereas we assayed sorted hepatocytes in a model with hepatocyte-specific Yap induction with RNA-sequencing. Thus, a complete overlap in gene profile would not be expected. Yap-induced genes showed an enrichment in cell adhesion, extracellular matrix (ECM), and cell–cell junction functional groups (Figure 2—figure supplement 2C), according to DAVID functional cluster analysis (Huang et al., 2009a, 2009b). In contrast, the cell cycle category was not significantly enriched, likely because only a small fraction of all Yap<sup>high</sup> hepatocytes were proliferating, as evidenced by BrdU labeling.

It is well known that pro-growth effects of Yap overexpression are mediated through TEAD (Wu et al., 2008; Zhang et al., 2008; Zhao et al., 2008), and the dominant-negative TEAD mutant largely suppresses effects of Yap transgene in the liver (Liu-Chittenden et al., 2012). To evaluate how many of the Yap-induced genes are potentially regulated directly by Yap-TEAD complexes, we compared these genes to genes with TEAD binding sites identified previously by ChIP-seq (ENCODE Project Consortium, 2011). Indeed, over 25% of all Yap-activated genes are associated with TEAD compared to only 9% of all genes in the genome. Moreover, a large fraction of genes induced in Yap<sup>high</sup> hepatocytes and related to cell adhesion (33%), ECM (36%), and cell–cell junctions (36%) contain TEAD binding regions (Figure 2—figure supplement 2D). The gene expression data shown here suggest that inability of Yap to drive cell expansion in our model is not due to lack of transcriptionally competent Yap-TEAD complexes.

To test whether mosaic Yap activation can lead to clonal expansion in previously characterized mouse models, we injected Stk4<sup>−/−</sup>;Stk3<sup>flox/flox</sup> mice with a low dose of adenovirus expressing Cre and GFP, and compared patterns of BrdU incorporation and GFP+ hepatocytes 5 and 35 days later. At both time points, we could only observe single GFP+ hepatocytes, BrdU negative (Figure 2—figure supplement 4A). Semi-quantitative PCR of the recombined Stk3<sup>ΔMst2</sup> allele also did not show an increase at 35 days, and liver/body mass ratio remained normal (Figure 2—figure supplement 4B,C).

Taken together, these data suggest that clonal activation of Yap in hepatocytes is not sufficient to overrule growth restricting mechanisms at steady state, but is sufficient to regulate transcription of a large number of genes related to extracellular environment.
Yap promotes hepatocyte proliferation in response to injury

Previous studies demonstrated that Yap is activated during liver injury (Bai et al., 2012; Wang et al., 2012; Anakk et al., 2013) and protects mice from oxidative stress-induced damage (Wu et al., 2013). We therefore tested whether Yap induction provides selective advantage during liver injury. We chose carbon tetrachloride (CCl4) as a damaging agent due to its localized effects around the central vein (CV) (Weber et al., 2003).

We first characterized the kinetics of liver damage after a single injection of CCl4. Blood ALT levels in YapKI Alb-Cre and WT mice rose to similar levels, indicating comparable hepatocyte damage (Figure 3—figure supplement 1). By day 4 the acute damage phase ceased completely (Figure 3—figure supplement 1) and very few hepatocytes incorporated BrdU (Figure 3—figure supplement 2). We therefore chose day 4 to study the outcome of the tissue repair response.

In response to CCl4 damage Yaphigh cells expanded dramatically within 4 days in the pericentral area, comprising over 80% of all hepatocytes in this zone (Figure 3A,B). These pericentral Yaphigh cells incorporated significantly more BrdU during days 1–3 after CCl4 administration compared to Yaplow hepatocytes in the same location (Figure 3C). In contrast, Yaphigh cells proliferated much less in the CV-distal areas than in the pericentral area (Figure 3C). In fact, the BrdU index outside the pericentral zone did not differ between Yaphigh and Yaplow hepatocytes within the same liver. Even though Yaphigh cells expanded locally at the injury sites, the effect was strong enough to significantly increase total percentage of Yaphigh cells in the liver (Figure 3D). Nevertheless, this expansion of Yaphigh cells did not overrule normal liver size control, as the total number of hepatocytes was similar in YapKI Alb-Cre and WT mice (Figure 3—figure supplement 3). Liver/body weight ratios in both groups also returned to steady-state levels within a month after undergoing a similar increase in response to CCl4, (likely due to the infiltration of immune cells during early phase of tissue repair) (Figure 3—figure supplements 3–5). These results indicate that Yaphigh cells undergo expansion in response to local tissue injury signals.

To investigate whether Yap activity in hepatocytes is required for tissue repair, we generated Yap conditional knockout mice and crossed them to the AlbCre strain to obtain hepatocyte-specific Yap knockout Yap1floxflox.AlbCre (YapCKO Alb-Cre) mice (Figure 4—figure supplements 1, 2). These mice were treated with CCl4 using the same protocol as described above. A similar increase in blood ALT level in YapCKO Alb-Cre and WT mice indicated comparable liver damage in the first 2 days (Figure 4—figure supplement 3). By day 4, ALT levels returned to baseline (Figure 4—figure supplement 3), normal pericentral morphology was restored, and expression of the CV-proximal marker glutamine synthase was reestablished in both groups (Figure 4A). However, YapCKO Alb-Cre hepatocytes proliferated significantly less than their WT counterparts throughout the repair phase, as shown by BrdU labeling on days 1–3 (Figure 4B,C). Consistent with the defective regeneration, YapCKO Alb-Cre livers displayed abnormal collagen deposition around the CV (Figure 4D). Interestingly, the steady-state liver/body mass ratio (Figure 4—figure supplement 4) and BrdU incorporation rate (Figure 4—figure supplement 5) were higher in the YapCKO Alb-Cre mice than in the WT mice, suggesting that Yap is not required for homeostatic hepatocyte proliferation.

Thus, while Yap is not required for liver development and hepatocyte proliferation under normal conditions, Yap function is required for hepatocyte proliferation during tissue repair. Additionally, hepatocytes with a higher level of Yap have growth advantage during the early repair phase.

Yap promotes hepatocyte proliferation in response to inflammation

Given differential effects of Yap on hepatocyte proliferation during homeostasis vs injury, we wondered how growth control mechanisms might differ under these conditions. Tissue injury is accompanied by inflammation, which can promote proliferation by mechanisms distinct from homeostasis. To test whether Yap cooperates with inflammatory signals to drive cell cycle progression, we measured hepatocyte BrdU incorporation in YapKI Alb-Cre mice after 6 continuous days of intraperitoneal injections of LPS. This chronic LPS treatment induced a massive expansion of Yaphigh hepatocytes, similar to what was seen after CCl4 treatment (Figure 3D). Consistently, the BrdU incorporation index indicated that Yaphigh hepatocytes proliferated significantly more than hepatocytes in the WT livers (Figure 5A,B). YapKAlb-Cre livers displayed significant hyperplasia, as visualized by increased hepatocyte density (Figure 5C) and measured by total number of hepatocytes per liver (Figure 5D).

We then tested whether IL-6, an inflammatory cytokine induced by LPS and by liver injury, can cooperate with Yap in promoting hepatocyte growth. When IL-6 was produced by LLC cells growing as subcutaneous tumors in YapKAlb-Cre mice, it caused expansion of Yaphigh hepatocytes in the livers of
Figure 3. Injury induces proliferation and clonal expansion of Yap-overexpressing hepatocytes. (A) Liver sections of control (WT) and YapKIAlb-Cre (KI) mice isolated at steady state (SS) or on days 4 and 14 after CCl4 treatment were stained with antibodies to GFP and BrdU. BrdU was injected in all groups for 3 consecutive days before harvesting the livers. (B) The percentage of Yap^∗^ (GFP^+^) cells among all hepatocytes was quantified within the pericentral zone (‘damage site’) and outside the pericentral zone (‘outside’) and in the liver sections of YapKIAlb-Cre mice on day 4 after CCl4 treatment. 3–4 mice were used for each group and 3–4 images were taken for each mouse. Each dot represents cell count from each image. ****p ≤ 0.0001. (C) The percentages of BrdU^+^ hepatocytes among GFP^−^ and GFP^+^ hepatocytes were quantified outside the CV zone (‘outside’) and around CV (‘damage site’) in liver sections of YapKIAlb-Cre mice on day 4 after CCl4 treatment. 3–4 mice were used for each group and 3–4 images were taken for each mouse. Each dot represents a cell count from each image. ****p ≤ 0.0001. (D) Hepatocytes were isolated by collagenase perfusion from 1- to 2-month-old YapKIAlb-Cre mice at steady state (SS), at day 4 (D4) and day 14 (D14) after CCl4 treatment, or after a 6-day LPS treatment (LPS), and percentage of GFP^high^ cells determined by flow cytometry.

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The following figure supplements are available for figure 3:

Figure supplement 1. Kinetics of CCl4-induced liver damage in the YapKIAlb-Cre mice.
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Figure supplement 2. Proliferation rates of WT and YapKIAlb-Cre hepatocytes are similar on day 4 after CCl4 treatment.
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Figure supplement 3. Hepatocyte numbers undergo similar changes in response to CCl4 in the WT and YapKIAlb-Cre mice.
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Figure supplement 4. Liver/body weight ratios undergo similar changes in response to CCl4 in the WT and YapKIAlb-Cre mice.
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Figure supplement 5. Hematopoietic cell numbers undergo similar changes in response to CCl4 in the WT and YapKIAlb-Cre mice.
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mice that had the highest levels of systemic IL-6 (Figure 5—figure supplement 1A,B), suggesting that IL-6 may cooperate with Yap to promote hepatocyte proliferation.

**Yap upregulates progenitor markers and represses hepatocyte differentiation in response to injury**

4 days after CCl₄ treatment in WT mice, normal liver tissue morphology was restored around the CV. In contrast, we observed high cell density, strong cytoplasmic eosinophilia, round to ovoid nuclei, and increased nuclear/cytoplasmic ratio in the pericentral zones of livers from CCl₄-treated YapKI Alb-Cre mice (Figure 6A). These morphological changes are often seen in hepatocellular carcinomas (HCC) and are associated with dedifferentiation (Chang et al., 2010; Balani et al., 2013). Immunofluorescent staining indicated that these cells were GFP⁺ and therefore originated from Yap⁺⁺ hepatocytes (Figure 3A). To determine whether Yap induction affects differentiation status of hepatocytes, we
compared the gene expression of hepatocytes sorted from YapKI Alb-Cre mice at steady state and 4 days after CCl4 damage. The majority of liver-specific transcripts (Pan et al., 2013) (which roughly represent hepatocyte differentiation state) were strongly repressed in Yap high hepatocytes from CCl4-treated mice, whereas WT and Yap low hepatocytes showed minimal alterations and no bias towards decreased expression (Figure 6B). Immunofluorescent staining and Western blot for hepatocyte-specific gene glutamine synthase also showed a strong downregulation of expression in YapKI Alb-Cre livers after CCl4 damage (Figure 6C,D). Downregulation of other hepatocyte-specific genes (acute phase protein Orm1 and urea cycle component Asl) was also confirmed by qPCR (Figure 6—figure supplement 1). To
Figure 6. Yap activation cooperates with tissue injury to repress hepatocyte differentiation and promote progenitor phenotype. (A) H&E staining of liver sections of control (WT) and YapK<sup>Alb-Cre</sup> (KI) mice isolated on day 4 after CCL<sub>4</sub> treatment. (B) Heatmap of RNA-sequencing data comparing expression of liver specific genes (see 'Materials and methods'). Hepatocytes were sorted from wild-type livers (WT) or from YapK<sup>Alb-Cre</sup> livers based on GFP levels (Lo
and Hi), at steady state (SS) or on day 4 after CCl4 treatment (CCl4). The fold change is calculated between the indicated samples and WT in steady state. Data represent the mean of the duplicates. (C) Liver sections of control (WT) and YapKIAlb-Cre (KI) mice isolated on days 4 CCl4 treatment were stained with antibodies to glutamine synthase. (D) Glutamine synthase (GS), Yap and beta-actin protein levels in whole-liver protein lysates prepared on day 4 after CCl4 treatment were determined by Western blotting. Higher migrating band in the middle panel corresponds to the exogenous Yap (due to the in-frame triple flag tag). (E and F) Heatmaps showing the RNA-sequencing data for genes enriched in embryonic tissues (E) or progenitor markers (F). Hepatocytes were sorted from wild type livers (WT) or from YapKIAlb-Cre livers based on GFP levels (Lo and Hi), at steady state (SS) or on day 4 after CCl4 treatment (CCl4). The fold change is calculated between the indicated samples and wild type in steady state. (G and H) Flow cytometric analysis of primary hepatocytes isolated by collagenase perfusion from CCl4-treated control (WT) or YapKIAlb-Cre (KI) mice. (G) Representative flow cytometry plots gated of CD45+CD31 hi population. The numbers on the plots indicate the percentages of the gated population (EpCam+ progenitors). (H) The results from (G) were combined with the total hepatocyte numbers to calculate the number of hepatic EpCam+ progenitors per liver.

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The following figure supplement is available for figure 6:

Figure supplement 1. Verification of the RNA-sequencing results.
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Further verify RNA-sequencing data and evaluate levels of Yap activation in our model, we performed qPCR analysis of the genes previously shown to be induced in Yap-Tg liver by Pan’s group (Dong et al., 2007) and found comparable effects (Figure 6—figure supplement 1).

The transcriptional repression was not global: a number of genes expressed predominately during embryogenesis (Pan et al., 2013) were induced in Yaphigh hepatocytes, and this effect was strongly enhanced by injury (Figure 6E). In addition, several hepatocyte progenitor markers were upregulated in Yaphigh cells after injury, and some of them were already elevated in the steady state (Figure 6F). FACS staining of liver cells isolated on day 4 post-CCl4 confirmed confirmed upregulated expression of CD45+CD31hi EpCam+ hepatic progenitors in YapKI;AlbCre livers (Figure 6G,H). These data are in line with the recent report by Yimlamai et al. (2014).

Altogether, these data suggest that Yap cooperates with injury signals to repress hepatocyte differentiation and promote a progenitor phenotype.

Excessive Yaphigh cells are eliminated during repair resolution phase

The expansion of Yaphigh cells induced by CCl4 did not persist, and by day 14 after the treatment, the proportion of Yaphigh cells per liver returned to the average value of the untreated group (Figure 3D). Yaphigh cells formed a ring of only 1–2 cell layers around CVs, in contrast to multiple layers observed on day 4 (Figure 3A), indicating that the Yaphigh cell pool underwent contraction. TUNEL staining suggested that the contraction was mediated by apoptosis: the number of TUNEL positive cells was markedly elevated in the YapKIAlb-Cre livers specifically around the CV on day 4 after CCl4 treatment (Figure 7A). Accordingly, RNA sequencing revealed a number of pro-apoptotic genes (e.g., Bok, Bax, NGFRAP1, and Tnrsf10b (Dr5)) (Figure 7B) upregulated as a consequence of Yap overexpression and further induced after damage. We verified by qPCR that Dr5 gene expression was strongly upregulated in the Yaphigh cells after CCl4 (Figure 7C,D). Indeed, Yaphigh cells sorted from CCl4-treated livers were more sensitive to killing mediated by TRAIL in vitro, compared to WT or GFPlow counterparts (Figure 7E), suggesting that Yap activation sensitizes hepatocytes to Dr5-mediated cell death. This higher sensitivity to cell death was induced by a combination of injury and Yap overexpression, as it was not observed neither in Yaphigh hepatocytes isolated from untreated mice nor in WT hepatocytes from CCl4 livers.

In summary, the expanded population of Yaphigh cells contracts to baseline at later stages of the tissue injury response. This process is likely mediated, in part, by sensitization of Yaphigh cells to TRAIL-induced apoptosis via upregulation of Dr5.

Discussion

Yap has emerged as a central mediator of signals promoting proliferation, survival, and stemness (Barry and Camargo, 2013). Not surprisingly, an expanding number of pathways restricting the pro-growth potential of Yap have been identified, including the Hippo pathway, the actin cytoskeleton and cell junction complexes (Boggiano and Fehon, 2012), C/EBPa, and Trib2 (Wang et al., 2013). Despite these multifactorial control mechanisms, mere overexpression of Yap in the whole liver in mouse models leads to hyperplasia, organomegaly, dedifferentiation, and cancer
Figure 7. Yap sensitizes hepatocytes to TRAIL-mediated apoptosis. (A) Cell death as reflected by TUNEL-positive cell numbers quantified outside the CV zone ('outside') and around CV ('damage site') in liver sections of YapKIAlb-Cre mice on day 4 after CCl4 treatment. 3–4 mice were used for each group and 3–4 images were taken from each mouse. Each dot represents cell count from each image. (B) RPKM data from RNA-sequencing illustrating expression of apoptosis-related genes in WT or Yaphigh (Hi) hepatocytes sorted from untreated (SS) livers or on day 4 after CCl4 treatment (CCl4). (C and D) Dr5 mRNA levels in WT and YapKIAlb-Cre livers (KI) at indicated time points after CCl4 treatment were determined by qPCR in whole liver extracts (C), or in primary hepatocytes sorted by flow cytometry based on GFP levels from steady-state livers (SS) or on day 4 after CCl4 treatment (CCL4) (D). 2–3 mice were used for each group. *p ≤ 0.05. (E) Primary hepatocytes sorted as in (D) were cultured on collagen-coated plates with or without TRAIL, and cell viability measured by CellTiter-Blue the next day. 3–4 mice were used for each group and 3–4 wells were seeded with hepatocytes from each mouse. Each dot represents reading from each well. DOI: 10.7554/eLife.02948.030
(Camargo et al., 2007; Dong et al., 2007; Schlegelmilch et al., 2011), raising a question as to whether some layer of homeostatic growth control may be lacking in such models. Organ-wide Yap activation in the liver does occur during embryonic development (Septer et al., 2012) and adaptive hepatomegaly (Kowalk et al., 2011). Interestingly, these are also examples of physiologically relevant settings of liver size increase. Transgenic Yap overexpression in the entire liver may thus be viewed as a model of these physiological scenarios. In contrast, during response to a localized injury, only the cells within the damage site need to proliferate, and Yap activation in this case should conform to the physiological organ size limits. Our study suggests that under these circumstances, activation of a pro-growth gene program by Yap requires an additional local signal (e.g., growth factor, cytokine, or release of contact inhibition) provided by the injury. Our preliminary data implicate IL-6 as at least one of such signals (Figure 5—figure supplement 1). Proto-oncogenes typically require two signals in order to induce proliferation. In case of Yap, which acts as a sensor of cell density, its overexpression mimics a condition in which a cell has lost its contacts with the neighbors. This in itself should not induce proliferation unless cell-extrinsic signals of tissue damage such as inflammatory cytokines are also present. The mechanism of cooperation between activation of Yap and injury/inflammatory signals may occur at the level of gene expression, as many proliferation genes have TEAD as well as NFkB binding sites (Figure 5—figure supplement 2). While it is also possible that cooperation is mediated by increased Yap stability or recruitment of TEAD, this possibility is less likely as we observed induction of high number of TEAD-dependent genes in Yap-overexpressing hepatocytes whereas proliferation genes were not induced.

Our data suggest that the tissue environment imposes a selective pressure on cells that activate Yap. The mosaic model of Yap induction described here reveals selective pressures that determine the fate of Yap^high^ hepatocytes: no growth advantage at steady state, positive selection during early stages of tissue injury, and negative selection at later stages when injury induced signals subside. The mechanisms opposing clonal expansion of Yap^high^ hepatocytes under homeostatic conditions may have many components, but one implicated by our data is apoptosis. Pro-apoptotic effects of Yap have been extensively documented in vitro (Strano et al., 2001; Lapi et al., 2008; Tomlinson et al., 2010), and increased apoptosis has been reported in Mst1/2-deficient mouse livers (Lu et al., 2010), but to our knowledge there has been no evidence of Yap promoting cell death in vivo. We show that Yap activation in the mouse liver leads to increased apoptosis in vivo and sensitization to TRAIL measured ex vivo. The induction of proliferation or apoptosis by Yap is modulated by tissue damage and inflammatory signals. This is analogous to the well-characterized functions of c-myc, which is also known induce proliferation or apoptosis, depending on growth factor availability (Evan et al., 1992; Harrington et al., 1994). These examples may represent a general principle of mitogenic pathway design, aimed at elimination of aberrant cells.

YapKI^Ab-Cre^ mice display many features described in other in vivo models of Yap induction, including increased proliferation, dedifferentiation, and activation of TEAD target genes. However, the unexpected phenotype unique to YapKI^Ab-Cre^ mice is that the pro-growth effects of Yap are only manifested in a state of altered homeostasis, such as tissue injury or inflammation. These differences are unlikely to be due to an insufficient level of Yap expression, as a large number of genes are highly induced by Yap under steady-state conditions in YapKI^Ab-Cre^ mice. One relevant difference may be mosaic vs tissue-wide Yap activation. We show that mosaic deletion of Mst2 in Mst1−/− background does not lead to clonal expansion or liver size increase. Furthermore, in another mouse model recently reported by Camargo et al. (Yilmamai et al., 2014) while this manuscript was under consideration, mosaic Yap activation in hepatocytes did not induce proliferation of clonal expansion in the first weeks. Instead, it promoted hepatocyte dedifferentiation into progenitors, which then expanded. Taken together, these results suggest that activation of Yap in hepatic progenitor cells is sufficient for clonal expansion but in mature hepatocytes it requires the second signal to drive proliferation. Another possible cause of different phenotypes may lie in different levels of inflammation between our mouse model and the whole liver transgenics. Of note, livers with a hepatocyte-specific ablation of Mst1/2 or Sav (both of which result in Yap activation) have elevated expression of immune response genes, including TNFa and IL-6 (Lu et al., 2010). Many variables may contribute to differences in systemic inflammation. One likely contributor is the intestinal microbiome composition, which varies greatly across mouse facilities (Bleich and Hansen, 2012) and can induce inflammatory responses in the liver (Henao-Mejia et al., 2012). Another source of inflammation in case of whole-liver Yap overexpression may be the disruption of systemic metabolism due to Yap-mediated repression of...
liver-specific genes (Yimlamai et al., 2014) and our RNA-sequencing analysis. As liver-specific genes are still expressed at normal levels in a large fraction of hepatocytes (Yaplow cells), systemic metabolism is not compromised in YapKI Alb-Cre mice. It is therefore possible that the level of systemic inflammation is low in our mice, and this is why they require an exogenous inflammatory signal to enable Yap pro-growth effects. Inflammation is an essential component of carcinogenesis (Ben-Neriah and Karin, 2011) and anti-inflammatory treatments can inhibit aberrant proliferation and delay tumor development (Pribluda et al., 2013). Inflammation can promote tumor development through multiple mechanisms (Ben-Neriah and Karin, 2011), including provision of a permissive signal for proliferation in the settings of disturbed homeostasis (Bondar and Medzhitov, 2013; Pribluda et al., 2013).

Our RNA-sequencing analysis has revealed two important functions of Yap in hepatocytes: first, the largest category of Yap-regulated genes is related to ECM and cell adhesion. These transcriptional targets of Yap appear to be the most common across many cell types and contexts of Yap activation; for example, the gene widely used as a hallmark of Yap activation is a matricellular growth factor CTGF. Since loss of cell contacts is known to activate Yap, it makes sense that Yap induces a restorative transcriptional program. Second, Yap strongly suppresses liver-specific genes in response to injury. Hepatocytes are known to downregulate liver-specific transcripts as they enter the cell cycle during liver regeneration (Ito et al., 1991). Proliferation and tissue-specific activities are often segregated between progenitor and differentiated cells, especially in tissues with high cell turnover. This suggests that extensive proliferation may be generally incompatible with specialized tissue functions.

In conclusion, tissue remodeling, cell cycle progression, and repression of tissue-specific functions all need to be orchestrated in response to injury, and our findings suggest Yap as a key coordinator of these activities during tissue repair. Furthermore, our data highlight the role of tissue microenvironment in the outcome of Yap activation and argue that during homeostasis, growth-promoting function of Yap in differentiated cells requires an additional signal, which may be provided by inflammation or injury.

Materials and methods

Generation of YapKI and YapCKO mice
To generate the YapKI and YapCKO lines, targeting vectors were designed and constructed as shown in Figure 1—figure supplement 1A and Figure 4—figure supplement 1. NM-001171147 mouse Yap1 isoform with introduced S112A mutation (corresponding to S127A of the human protein) was used to generate YapKI allele. After the successful construction of the targeting vectors, the purified plasmid DNA was linearized and transfected into the C57BL/6-derived Bruce4 ES cells. Correct recombination and replacement of the DNA were verified by Southern blots. Selected clones were expanded and submitted to the Yale Transgene Facility for injections into BALB/c blastocysts to generate chimeric mice. In case of the YapCKO line, mice with the germline-transmitted Yap floxed allele were then crossed to Flp-deleter transgenic mice (ENCODENew Consortium, 2011) to remove the neomycin cassette.

Mouse treatments
Animals were maintained at the Yale Animal Resources Center. All animal experiments were performed with approval by the Institutional Animal Care and Use Committee of Yale University. All mice were on C57BL/6 background. AlbCre mice were from Jackson Laboratory. For BrdU staining, daily intraperitoneal (i.p.) injections of BrdU (Sigma, St. Louis, MO; 100 μg/mouse) were performed for 3 days before liver dissection. Liver damage was induced by i.p. injection of CCl4 (Sigma, 1 μl/g, diluted 1:10 with peanut oil). For the LLC-IL-6 experiment, 1 million LLC cells stably expressing mouse IL-6 or empty vector were injected into each flank of 1- to 1.5-month-old YapKI Alb-Cre mice. 2–3 weeks later when tumors reached approximately 1 cm in diameter, the mice were eye-bled for IL-6 ELISA and sacrificed for hepatocyte cell preparation. For the in vivo deletion of Mst2flox/flox allele, AdCMVCre-RSV-GFP adenovirus from KeraFast was amplified in 293A cells, purified by ViraPur kit, dialized against PBS and injected into the tail vein of the 1- to 2-month-old Mst1−/− Mst2flox/flox mice.

Immunofluorescence microscopy
Liver lobes were fixed in 4% PFA overnight and embedded in paraffin. Sections were deparaffinized and heated for 20 min at 95°C for antigen retrieval in Tris-EDTA buffer pH 9.
TUNEL staining was performed using In Situ Cell Death Detection Kit (Roche, Germany). For GFP, Yap, and GS immunofluorescence staining, samples were then blocked in TBS-T with 3% BSA for 2 hr (staining buffer), incubated with the primary antibody overnight and with the secondary antibody for 1 hr. For BrdU staining, after the described steps, the samples were fixed in 4% PFA in PBS for 20 min at RT. DNA was then denatured in 2 N HCl for 30 min at 37°C. After neutralization with 0.1 M sodium tetraborate, the sections were stained with mouse anti-BrdU antibodies for 2 hr and visualized with fluorophore-conjugated anti-mouse secondary antibodies. The slides were mounted in Vectashield anti-fade media containing Hoescht dye to counterstain the nuclei. 20–30 20× tiled images were obtained using Zeiss Axioplan microscope and AxioVision software. Cells were counted manually in AxioVision software. Only hepatocytes (as determined by morphology and GFP expression where applicable) were scored for BrdU quantification.

Hepatocyte FACS
Primary hepatocytes were prepared by collagenase perfusion at Yale Liver Center. Hepatocytes were washed twice with FACS buffer (2 mM EDTA, 2% FBS in PBS), stained with 7-Aminoactinomycin D (7AAD) or propidium iodide, and analyzed on BD Accuri or LSRII, or sorted on MoFlo cytometers. Samples were gated on live hepatocytes based on forward and side scatter and live dye exclusion.

RNA sequencing analysis
Duplicate hepatocyte samples were prepared in several independent experiments. Each of the duplicates contained material pooled from multiple mice. Hepatocytes were isolated by flow cytometry, and RNA was extracted using RNeasy kit (Qiagen, Germany). 1–10 μg of each sample was submitted to the Yale Center of Genome Analysis.

Sequencing libraries were constructed and were sequenced by Illumina Hiseq 2500 with 76 bp single-end reads, which generated 20 million raw reads per sample on average. After removing the low-quality reads (around 0.18% of all reads) and low-quality portions (Q value <30) of each of the raw reads, the RNA-sequencing data for each sample were mapped to the GRCm38/mm10 mouse reference first. The GRCm38 reference was downloaded from the Mouse Genome Informatics (MGI). The mapping was performed using TopHat v2.0.8, allowing two mismatches. The percentages of mapped reads were 71.16% on average (from 70.00% to 73.79%). After mapping, Cufflinks v2.0.2 was applied to assemble and quantify the transcripts and discover the differentially expressed genes, with the annotated gene information from the MGI. The gene expression values were calculated for each sample based on the number of fragments per kilobase of exon per million reads mapped (FPKM). The significance of differential expression of genes was detected using Cuffdiff for all comparisons of every two samples.

To select differentially expressed genes, the average of two biological replicates was compared between the conditions examined. To select genes with significant activation or repression in any comparison, the p-value of 0.005 was used together with a fold-change threshold of twofold. A pseudocount of 0.01 was added to RPKM value of each gene as a sequencing background, to avoid inflated fold change caused by lowly expressed genes.

Gene functional analysis
Gene functional analysis was performed with the DAVID functional annotation tools (Huang et al., 2009a, 2009b) (http://david.abcc.ncifcrf.gov). The p-value was adjusted with Benjamini methods for multiple hypothesis testing.

Analysis of tissue-specific genes
Tissue-specific genes were obtained from Pattern Gene Database (Pan et al., 2013) (http://bioinf.xmu.edu.cn/ PaGenBase). Genes expressed specifically (only in one tissue) and selectively (only in a group of samples) were pooled together as the tissue-specific genes in our analysis. Liver-specific genes are identified from mouse liver sample, and mouse embryo specific genes are identified from mouse day 9.5 embryo sample.

Transcription factor binding analysis
Genome-wide binding of Tead4 (identified by ChIP-sequencing) was retrieved from the ENCODE Project Consortium (2011). The uniform peaks determined in all available samples (HepG2 cell line, Su et al. eLife 2015;4:e02948. DOI: 10.7554/eLife.02948 16 of 21
K562 cell line, and hESC) were combined, and genes whose TSS is within 10 kb distance from any Tead4 uniform peak were defined as Tead4-associated genes. Hg19 genome annotation was downloaded from the UCSC database (Karolchik et al., 2014).

**TRAIL sensitivity assay**

Primary hepatocytes were plated in collagen-coated wells in Williams’ E medium (WEM) supplemented with 5% FCS, 10 mM HEPES, 2 mM L-glutamine, 10 mM penicillin/streptomycin, 8 μg/ml gentamicin, 100 μg/ml chloramphenicol, 100 nM dexamethasone, and 1 nM Insulin. 4 hr after plating, the medium was changed to the supplemented WEM without chloramphenicol and dexamethasone. On the second day cells were treated with recombinant TRAIL (R&D Systems, Minneapolis, MN) overnight. Viable cell number was measured by the CellTiter-Blue assay (Sigma) according to the manufacturer protocol.

**Western blotting**

Whole liver tissue was snap frozen in liquid nitrogen, homogenized, and resuspended in TNT buffer (0.1 M Tris-HCL pH 7.5, 150 mM NaCl, 0.1% Tween20) supplemented with the Complete protease inhibitors (Roche) and cleared by centrifugation at 100×g. Protein concentration in the supernatants was normalized using Bradford reagent. 15 μg of the extracts were separated on a gradient SDS-PAGE gel and transferred to PVDF membrane. After blocking with 3% BSA in TBST buffer, the membranes were probed with antibodies to Yap (Cell Signaling, Beverly, MA), glutamine synthase (Genscript, Piscataway, NJ), and beta-actin (Sigma).

**ALT test**

ALT tests were performed using ALT activity assay (Sigma) according to the manufacturer’s protocol.

**Real-time PCR**

RNA was extracted with RNA-bee, and cDNA synthesis was performed using SMART MMLV reverse transcriptase according to the manufacturers’ protocols. Real time PCR was performed in triplicates using Quanta SYBR reagent on C1000 Thermal Cycler (BioRad, Hercules, CA). Primer sequences are listed in the Supplementary file1.

**Liver genomic PCR**

Three 1 mm³ pieces of liver tissue per mouse were digested with Pronase overnight, and genomic DNA isolated by NaCl/ethanol precipitation. The three DNA samples for each mouse were pooled and used at fourfold dilutions to amplify the Rag2 or recombined Stk3floxflox alleles using Taqman polymerase.

**Enzyme linked immunosorbent assay (ELISA)**

Paired antibodies against IL-6 were purchased from BD Biosciences to perform ELISA.

**Statistical analysis**

All statistical analysis was performed by two tailed unpaired Student’s t test, unless specified otherwise.

**Accession numbers**

The RNA sequencing data have been deposited to GEO database under the accession number GSE65207.

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**Author contributions**

TS, TB, Conception and design, Acquisition of data, Analysis and interpretation of data, Drafting or revising the article; XZ, Performed bioinformatic analysis of RNA-seq data to identify enrichment in transcription factor binding sites in the promoters and enhancers of Yap-regulated genes under normal conditions and in response to liver damage; CZ, Performed experiments with MST1/MST2 deficient mice, CCL4 treated mice, as well as experiments with mice injected with IL-6 transfected LLCs; HH, Processed raw RNA-seq data, Performed bioinformatic and statistical analysis; RM, Conception and design, Analysis and interpretation of data, Drafting or revising the article

**Ethics**

Animal experimentation: All animal experiments were performed with approval by the Institutional Animal Care and Use Committee of Yale University (protocol # 2014-08006).

**Additional files**

**Supplementary file**
- Supplementary file 1. List of primer sequences.
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**Major dataset**

The following dataset was generated:

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|-----------|------|---------------|-----------------------|-----------------------------------------------|
| Su T, Bondar T, Zhou X, Zhang C, He H, Medzhitov R | 2015 | RNA sequencing data | http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE65207 | Publicly available at NCBI Gene Expression Omnibus (GSE65207). |

**References**

Anakk S, Bhosale M, Schmidt VA, Johnson RL, Finegold MJ, Moore DD. 2013. Bile acids activate YAP to promote liver carcinogenesis. Cell Reports 5:1060–1069. doi: 10.1016/j.celrep.2013.10.030.

Attisano L, Wrana JL. 2013. Signal integration in TGF-beta, WNT, and Hippo pathways. F1000primers 5:17. doi: 10.12703/P5-17.

Bai H, Zhang N, Xu Y, Chen Q, Khan M, Potter JJ, Nayar SK, Cornish T, Alpini G, Bronk S, Pan D, Anders RA. 2012. Yes-associated protein regulates the hepatic response after bile duct ligation. Hepatology 56:1097–1107. doi: 10.1002/hep.25769.

Balani S, Malik R, Malik R, Kapoor N. 2013. Cytomorphological variables of hepatic malignancies in fine needle aspiration smears with special reference to grading of hepatocellular carcinoma. Journal of Cytology/Indian Academy of Cytologists 30:116–120. doi: 10.4103/0970-9371.112655.

Barry ER, Camargo FD. 2013. The Hippo superhighway: signaling crossroads converging on the Hippo/Yap pathway in stem cells and development. Current Opinion in Cell Biology 25:247–253. doi: 10.1016/j.celbi.2012.12.006.

Barry ER, Morikawa T, Butler BL, Shrestha K, de la Rosa R, Yan KS, Fuchs CS, Magness ST, Smits R, Ogino S, Kuo CJ, Camargo FD. 2013. Restriction of intestinal stem cell expansion and the regenerative response by YAP. Nature 493:106–110. doi: 10.1038/nature11693.
Induction of apoptosis in fibroblasts by c-myc protein. 

| Author(s) | Year | Journal | Volume | Pages | DOI |
|-----------|------|---------|--------|-------|-----|
| Ben-Neriah Y, Karin M. | 2011 | Nature Immunology | 12 | 715–723 | 10.1038/ni.2060 |
| Bernasconi I, Martin-Belmonte F. | 2013 | Trends in Cell Biology | 33 | 380–389 | 10.1016/j.tcb.2013.03.007 |
| Beyer TA, Weiss A, Khomchuk Y, Huang K, Ogunjimi AA, Varelas X, Wrana JL. | 2013 | Cell Reports | 5 | 1611–1612 | 10.1016/j.celrep.2013.11.021 |
| Bleich A, Hansen AK. | 2012 | Comparative Immunology, Microbiology and Infectious Diseases | 35 | 81–92 | 10.1016/j.cimid.2011.12.006 |
| Boggiano JC, Fehon RG. | 2012 | Developmental Cell | 22 | 695–702 | 10.1016/j.devcel.2012.03.013 |
| Bondar T, Medzhitov R. | 2010 | Cell Stem Cell | 6 | 309–322 | 10.1016/j.stem.2010.03.002 |
| Bondar T, Medzhitov R. | 2013 | Cancer Cell | 24 | 143–144 | 10.1016/j.ccr.2013.07.016 |
| Cai J, Zhang N, Zheng Y, de Wilde RF, Maity A, Pan D. | 2010 | Genes & Development | 21 | 585–593 | 10.1016/j.gd.2007.07.019 |
| ENCODE Project Consortium. | 2011 | PLOS Biology | 9 | e1001046 | 10.1371/journal.pbio.1001046 |
| Evan GI, Wylie AH, Gilbert CS, Littlewood TD, Land H, Brooks M, Waters CM, Penn LZ, Hancock DC. | 1992 | Cell | 69 | 119–128 | 10.1016/0092-8674(92)90123-T |
| Fan R, Kim NG, Gumbiner BM. | 2013 | Current Biology | 21 | 2054–2060 | 10.1016/j.cub.2013.08.039 |
| Chang O, Yano Y, Masuzawa A, Fukushima N, Teramura K, Hayashi Y. | 2010 | Oncology Reports | 23 | 1229–1232 | 10.1016/j.tar.2010.07.051 |
| Chen C, Schroeder MC, Kango-Singh M, Tao C, Halder G. | 2012 | Proceedings of the National Academy of Sciences of USA | 109 | 484–489 | 10.1073/pnas.1113882109 |
| Dong J, Feldmann G, Huang J, Wu S, Zhang N, Comerford SA, Gayyed MF, Anders RA, Maity A, Pan D. | 2007 | Cell | 130 | 1120–1133 | 10.1016/j.cell.2007.07.019 |
| ENCODE Project Consortium. | 2011 | Nature Protocols | 6 | 1056–1063 | 10.1038/nprot.2012.202 |
| Harrington EA, Bennett MR, Fanidi A, Evan GI. | 1994 | The EMBO Journal | 13 | 3286–3295 | 10.1016/0261-4189(94)90123-T |
| Hena-O-Meja J, Elinav E, Jin C, Hao L, Mehal WZ, Strowig T, Thaiss CA, Kau AL, Eisenbarth SC, Jurcza MJ, Camporez JP, Shulman GI, Gordon JI, Hoffman HM, Flavell RA. | 2012 | Nature | 482 | 179–185 | 10.1038/nature10809 |
| Hong W, Guan KL. | 2012 | Seminars in Cell & Developmental Biology | 23 | 785–793 | 10.1016/j.semcdb.2012.05.004 |
| Huang DW, Sherman BT, Lempicki RA. | 2009a | Bioinformatics enrichment tools: paths toward the comprehensive functional analysis of large gene lists. | 17 | 1–13 | 10.1093/nar/gkn293 |
| Huang DW, Sherman BT, Lempicki RA. | 2009b | Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. | 4 | 44–57 | 10.1093/nprot.2008.211 |
| Ito Y, Hayashi H, Taira M, Tatibana M, Tabata Y, Isono K. | 1991 | Cell Biology | 51 | 143–147 | 10.1016/0022-4701(91)90085-8 |
| Karolchik D, Barber GP, Casper J, Clawson H, Cline MS, Diekhans M, Dresser TR, Fujita PA, Guruvadoo L, Haeussler M, Harte RA, Heintner S, Hinrichs AS, Learned K, Lee BT, Li CH, Raney BJ, Rhead B, Rosenbloom KR, Sloan CA, Speir ML, Zweig AS, Haussler D, Kuhn RM, Kent WJ. | 2014 | UCSC Genome Browser database: 2014 update. | 24 | D764–D770 | 10.1093/nar/gkt1168 |
| Karpowicz P, Perez J, Perrimon N. | 2010 | The Hippo tumor suppressor pathway regulates intestinal stem cell regeneration. | Development | 137 | 4135–4145 | 10.1242/dev.060483 |
| Konsavage WM Jr, Yochum GS. | 2013 | Hepatology | 53 | 2086–2096 | 10.1002/hep.24289 |
| Su et al. eLife 2015;4:e02948. DOI: 10.7554/eLife.02948 | 19 of 21 | | | | |
Lapi E, Di Agostino S, Donzelli S, Gal H, Domany E, Rechavi G, Pandolfi PP, Givol D, Strano S, Lu X, Blandino G. 2008. PML, YAP, and p73 are components of a proapoptotic autoregulatory feedback loop. Molecular Cell 32: 803–814. doi: 10.1016/j.molcel.2008.11.019.

Lee KP, Lee JH, Kim TS, Kim TH, Park HD, Byun JS, Kim MC, Jeong WI, Calvisi DF, Kim JM, Lim DS. 2010. The Hippo-Salvador pathway restrains hepatic oval cell proliferation, liver size, and liver tumorigenesis. Proceedings of the National Academy of Sciences of USA 107: 8248–8253. doi: 10.1073/pnas.0912203107.

Lin JT, Poon CL, Harvey KF. 2013. The Hippo size control pathway—ever expanding. Science Signaling 6: pe4. doi: 10.1126/scisignal.2003813.

Liu-Chittenden Y, Huang B, Shim JS, Chen Q, Lee SJ, Anders RA, Liu JO, Pan D. 2012. Genetic and pharmacological disruption of the TEAD-YAP complex suppresses the oncogenic activity of YAP. Genes & Development 26: 1300–1305. doi: 10.1101/gad.192856.112.

Lu L, Li Y, Kim SM, Bossuyt W, Liu P, Qiu Q, Wang Y, Halder G, Finegold MJ, Lee JS, Johnson RL. 2010. Hippo signaling is a potent in vivo growth and tumor suppressor pathway in the mammalian liver. Proceedings of the National Academy of Sciences of USA 107: 1437–1442. doi: 10.1073/pnas.0911427107.

Mullen AC. 2014. Hippo tips the TGF-beta scale in favor of pluripotency. Cell Stem Cell 14: 6–8. doi: 10.1016/j.stem.2013.12.009.

Norman M, Wisniewska KA, Lawrenson K, Garcia-Miranda P, Tada M, Kajita M, Mano H, Ishikawa S, Ikegawa M, Shimada T, Fujita Y. 2012. Loss of Scribble causes cell competition in mammalian cells. Journal of Cell Science 125: 59–66. doi: 10.1242/jcs.085803.

O’Reilly LA, Huang DC, Strasser A. 1996. The cell death inhibitor Bcl-2 and its homologues influence control of cell cycle entry. The EMBO Journal 15: 6979–6990.

Pan JB, Hu SC, Shi D, Cai MC, Li YB, Zou Q, Ji ZL. 2013. PaGenBase: a pattern gene database for the global and dynamic understanding of gene function. PLOS ONE 8:e80747. doi: 10.1371/journal.pone.0080747.

Pribluda A, Elyada E, Wiener Z, Hamza H, Goldstein RE, Biton M, Burstein I, Morgenstern Y, Brachya Y, Billauer H, Biton S, Snir-Alkalay I, Vucic D, Schlereth K, Mernberger M, Steewe T, Oren M, Alitalo K, Pikarsky E, Ben-Neriah Y. 2013. A senescence-inflammatory switch from cancer-inhibitory to cancer-promoting mechanism. Cancer Cell 24: 242–256. doi: 10.1016/j.ccr.2013.06.005.

Reddy BV, Irvine KD. 2013. Regulation of Hippo signaling by EGFR-MAPK signaling through Ajuba family proteins. Developmental Cell 24: 459–471. doi: 10.1016/j.devcel.2013.01.020.

Ren F, Wang B, Yue T, Yun EY, Ip YT, Jiang J. 2010. Hippo signaling regulates Drosophila intestine stem cell proliferation through multiple pathways. Proceedings of the National Academy of Sciences of USA 107: 21064–21069. doi: 10.1073/pnas.1012759107.

Schlegelmilch K, Mohseni M, Kirak O, Pruszk J, Rodriguez JR, Zhou D, Kreger BT, Vasioukhin V, Avruch J, Brummelkamp TR, Camargo FD. 2011. Yap1 acts downstream of alpha-catenin to control epidermal proliferation. Cell 144: 782–795. doi: 10.1016/j.cell.2011.02.031.

Septer S, Edwards G, Guenawardena S, Wolfe A, Li H, Daniel J, Apte U. 2012. Yes-associated protein is involved in proliferation and differentiation during postnatal liver development. American Journal of Physiology Gastrointestinal and Liver Physiology 302: G493–G503. doi: 10.1152/ajpgi.00056.2011.

Serrano M, Lin AW, McCurrach ME, Beach D, Lowe SW. 1997. Oncogenic ras provokes premature senescence associated with accumulation of p53 and p16INK4a. Cell 88: 593–602. doi: 10.1016/S0092-8674(00)81902-9.

Shao D, Zhai P, Del Re DP, Sciarretta S, Yabuta N, Nojima H, Lim DS, Pan D, Sadoshima J. 2014. A functional interaction between Hippo-YAP signalling and FoxO1 mediates the oxidative stress response. Nature Communications 5: 3315. doi: 10.1038/ncomms4315.

Shawl RL, Kohlmaier A, Polese N, Veelken C, Edgar BA, Tapon N. 2010. The Hippo pathway regulates intestinal stem cell proliferation during Drosophila adult midgut regeneration. Development 137: 4147–4158. doi: 10.1242/dev.052506.

Song H, Mak KK, Topol L, Yun K, Hu J, Garrett L, Chen Y, Park O, Chang J, Simpson RM, Wang CY, Gao B, Jiang J, Yang Y. 2010. Mammalian Mst1 and Mst2 kinases play essential roles in organ size control and tumor suppression. Proceedings of the National Academy of Sciences of USA 107: 1431–1436. doi: 10.1073/pnas.0914409107.

Staley BK, Irvine KD. 2010. Warts and Yorkie mediate intestinal regeneration by influencing stem cell proliferation. Current Biology 20: 1580–1585. doi: 10.1016/j.cub.2010.07.041.

Strano S, Munarriz E, Rossi M, Castagnoli L, Shaul Y, Sacchi A, Oren M, Sodol M, Cesareni G, Blando S. 2001. Physical interaction with Yes-associated protein enhances p73 transcriptional activity. The Journal of Biological Chemistry 276: 15164–15173. doi: 10.1074/jbc.M010484200.

Tomlinson V, Gudmundsdottir K, Luong P, Leung KY, Knebel A, Basu S. 2010. JNK phosphorylates Yes-associated protein (YAP) to regulate apoptosis. Cell Death & Disease 1: e29. doi: 10.1038/cddis.2010.7.

Wang C, Zhang L, He Q, Feng X, Zhu J, Xu Z, Wang X, Chen F, Li X, Dong J. 2012. Differences in Yes-associated protein and mRNA levels in regenerating liver and hepatocellular carcinoma. Molecular Medicine Reports 5: 410–414. doi: 10.3892/mmr.2011.640.

Wang J, Park JS, Wei Y, Rajurkar M, Cotton JL, Fan Q, Lewis BC, Ji H, Mao J. 2013. TRIB2 acts downstream of Wnt/TCF in liver cancer cells to regulate YAP and C/EBP alpha function. Molecular Cell 51: 211–225. doi: 10.1016/j.molcel.2013.05.013.

Wang K, Degerny C, Xu M, Yang XJ. 2009. YAP, TAZ, and Yorkie: a conserved family of signal-responsive transcriptional coregulators in animal development and human disease. Biochemistry and Cell Biology = Biochimie Et Biologie Cellulaire 87: 77–91. doi: 10.1139/O08-114.

Weber LW, Boll M, Stampfl A. 2003. Hepatotoxicity and mechanism of action of halogenated carbon tetrachloride as a toxicological model. Critical Reviews in Toxicology 33: 105–136. doi: 10.1080/713611034.
Wu H, Xiao Y, Zhang S, Ji S, Wei L, Fan F, Geng J, Tian J, Sun X, Qin F, Jin C, Lin J, Yin ZY, Zhang T, Luo L, Li Y, Song S, Lin SC, Deng X, Camargo F, Avruch J, Chen L, Zhou D. 2013. The Ets transcription factor GABP is a component of the hippo pathway essential for growth and antioxidant defense. *Cell Reports* 3:1663–1677. doi: 10.1016/j.celrep.2013.04.020.

Wu S, Liu Y, Zheng Y, Dong J, Pan D. 2008. The TEAD/TEF family protein Scalloped mediates transcriptional output of the Hippo growth-regulatory pathway. *Developmental Cell* 14:388–398. doi: 10.1016/j.devcel.2008.01.007.

Yimlamai D, Christodoulou C, Galli GG, Yanger K, Pepe-Mooney B, Gurung B, Shrestha K, Cahan P, Stanger BZ, Camargo FD. 2014. Hippo pathway activity influences liver cell fate. *Cell* 157:1324–1338. doi: 10.1016/j.cell.2014.03.060.

Yu FX, Guan KL. 2013. The Hippo pathway: regulators and regulations. *Genes & Development* 27:355–371. doi: 10.1101/gad.210773.112.

Yu FX, Meng Z, Plouffe SW, Guan KL. 2015. Hippo pathway regulation of Gastrointestinal tissues. *Annual Review of Physiology* 77:201–227. doi: 10.1146/annurev-physiol-021014-071733.

Yu FX, Zhao B, Panupinthu N, Jewell JL, Lian I, Wang LH, Zhao J, Yuan H, Tumaneng K, Li H, Fu XD, Mills GB, Guan KL. 2012. Regulation of the Hippo-YAP pathway by G-protein-coupled receptor signaling. *Cell* 150:780–791. doi: 10.1016/j.cell.2012.06.037.

Zaidi SK, Sullivan AJ, Medina R, Ito Y, van Wijnen AJ, Stein JL, Stein GS. 2004. Tyrosine phosphorylation controls Runx2-mediated subnuclear targeting of YAP to repress transcription. *The EMBO Journal* 23:790–799. doi: 10.1038/sj.emboj.7600073.

Zhang L, Ren F, Zhang Q, Chen Y, Wang B, Jiang J. 2008. The TEAD/TEF family of transcription factor Scalloped mediates Hippo signaling in organ size control. *Developmental Cell* 14:377–387. doi: 10.1016/j.devcel.2008.01.006.

Zhang N, Bai H, David KK, Dong J, Zheng Y, Cai J, Giovannini M, Liu P, Anders RA, Pan D. 2010. The Merlin/NF2 tumor suppressor functions through the YAP oncoprotein to regulate tissue homeostasis in mammals. *Developmental Cell* 19:27–38. doi: 10.1016/j.devcel.2010.06.015.

Zhang H, Pasolli HA, Fuchs E. 2011. Yes-associated protein (YAP) transcriptional coactivator functions in balancing growth and differentiation in skin. *Proceedings of the National Academy of Sciences of the USA* 108:2270–2275. doi: 10.1073/pnas.1019603108.

Zhao B, Ye X, Yu J, Li L, Li W, Li S, Yu J, Lin JD, Wang CY, Chinnaiyan AM, Lai ZC, Guan KL. 2008. TEAD mediates YAP-dependent gene induction and growth control. *Genes & Development* 22:1962–1971. doi: 10.1101/gad.1664408.

Zheng T, Wang J, Jiang H, Liu L. 2011. Hippo signaling in oval cells and hepatocarcinogenesis. *Cancer Letters* 302:91–99. doi: 10.1016/j.canlet.2010.12.008.

Zhou D, Zhang Y, Wu H, Barry E, Yin Y, Lawrence E, Dawson D, Willis JE, Markowitz SD, Camargo FD, Avruch J. 2011. Mst1 and Mst2 protein kinases restrain intestinal stem cell proliferation and colonic tumorigenesis by inhibition of Yes-associated protein (Yap) overabundance. *Proceedings of the National Academy of Sciences of USA* 108:E1312–E1320. doi: 10.1073/pnas.1110428108.