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**Title:** Yap1 as a New Therapeutic Target in Neurofibromatosis Type 2

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**Abstract:**
We hypothesized that Yap1 activity is critical for the growth-suppressing activity of Merlin, and that down-modulation of Yap1 function might constitute a novel therapeutic approach for the treatment of neurofibromatosis and other NF2-null tumors. In Aim 1, we will test whether ablation or reduction of Yap1 activity suppresses Nf2-null phenotypes in vivo. In aim 2, we will attempt to identify novel proteins and small molecules that negatively regulate Yap1 activity. We have demonstrated in fact that Nf2-ablation induces YAP activity in multiple tissues in mice. We have also established a reporter system for YAP activity and have finished a genetic screen identifying kinases that might activate YAP1.
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INTRODUCTION
Recently, our laboratory has provided evidence that the function of the Hippo pathway is evolutionary conserved in mammals and demonstrated that inactivation of Hippo signaling in mice leads to dramatic organ size overgrowth and cancer. Our preliminary findings indicate that downregulation of Merlin in human cells induces a potent activation of Yap1- (the mammalian homologue of Yorkie) reporter genes. Additionally, others have shown that Yap1 is preferentially nuclei-localized, and therefore active, in mesothelioma cells carrying NF2 mutations. Thus, based on previous studies and our preliminary data, we hypothesize that Yap1 activity is critical for the growth-suppressing activity of Merlin, and that down-modulation of Yap1 function might constitute a novel therapeutic approach for the treatment of neurofibromatosis and other NF2-null tumors.

Two specific aims are proposed. In Aim 1, we will test whether ablation or reduction of Yap1 activity suppresses Nf2-null phenotypes in vivo. Our laboratory has generated mice carrying hypomorphic and conditionally null Yap1 alleles and we will test whether these mutations result in the suppression of overgrowth phenotypes caused by the tissue specific disruption of NF2 in mice. In aim 2, we will attempt to identify novel proteins and small molecules that negatively regulate Yap1 activity. Our laboratory has developed a set of cell lines carrying transcriptional based-reporters of Yap1 activity. These reporter cell lines will be used to perform genome-wide RNAi and small molecule screens with the objective of finding novel modulators of Yap1 function. We believe that results obtained from these studies have the potential to unravel an entirely novel set of therapeutic targets for the treatment of NF2-deficient tumors.

BODY
Over the initial year months of our project we generated novel cell lines carrying the TEAD-based Hippo signaling reporters with an increased dynamic range of reporter activity (See appendix for attached manuscript). The screens performed utilized small interfering (si)-RNAs technology for gene knockdown. After extensive testing of different siRNA and shRNA reagents, we concluded that the siRNA-based Silencer Select oligos from Ambion provided the most efficient knockdown at the lowest concentrations with the minimal off-target effects. Additionally, preliminary tests indicated that siRNA-mediated knockdown of known molecules involved in Hippo signaling resulted in a much higher signal-to-noise ratio than that obtained with shRNAs against the same genes. With these new cell lines and siRNA reagents, we performed two targeted genetic screens. We have initially chosen to screen siRNA libraries against the human kinome and phosphatomes. These collections encompass 710 kinases and 298 phosphatases, with 3 siRNA oligos per gene. The screens have been performed in triplicate in 96-well plates with a cell line that carries a TEAD-reporter driving an mCherry reporter gene. Analysis of reporter output was done 4 days after transfection using high-throughput FACS analysis. Following the first round of screening and secondary validation, we have identified approximately 40 kinases and phosphatases that robustly modulate Hippo and YAP signaling. 16 of these hits have been validated with independent siRNAs. Some of these molecules activate, and others negatively regulate YAP activity. We have characterized the mechanism of action of these hits by analyzing the effects of siRNA knockdown on Yap1 localization and phosphorylation (see Appendix for attached manuscript). For a subset of these hits (Lkb1 and JNK-related molecules) we have performing biochemical and genetic epistatic experiments to determine putative interactions with known components of Hippo signaling. Multiple other pathways have also been discovered to be involved in YAP activation. We have found that a known tumor suppressor gene, known as LKB1 is a crucial component of the Hippo pathway, which functions downstream of Merlin signaling. LKB1, is a common tumour suppressor whose mechanism of action is only partially understood. We demonstrated that LKB1 acts through its substrates of the microtubule affinity-regulating kinase family to regulate the localization of the polarity determinant Scribble and the activity of the core Hippo kinases. Our data also indicate that YAP is functionally important for the tumour suppressive effects of LKB1. Our results identified a signalling axis that links YAP activation with LKB1 mutations, and have implications for the treatment of LKB1-mutant human malignancies. In addition, our findings provide insight into upstream signals of the Hippo-YAP signalling cascade. This work has been recently published (Mohseni et al, Nat Cell Bio 2014)
In parallel over the first 2 years of our work, we have generated and characterized TEAD-reporter cell lines that carry a firefly luciferase and optimized our screen conditions to 384-well plates and liquid-robotic and have began genome-wide screens. Over months 6-12 of this award we had finalized the first pass of a genome-wide siRNA screen aimed at identifying new Hippo pathway regulators. Two screens were performed: the first one in which 293T cells, which at baseline demonstrate low basal pathway activity, looking for genes whose knockdown activated YAP1 activity, and a screen in Nf2-deficient 293T cells, where baseline reporter is high and relevant knockdowns will induce reporter downregulation. This screen was performed in triplicate using 25,000 different oligos. The screens had to be repeated in a different cell line because of an initially poor signal-to-noise-ratio. After the finalization of the optimized screen approximately 1000 ‘hits’ have been selected as positive, and these are currently being tested in secondary and tertiary screens in which three individual oligos targeting each gene are used. The full set of data containing the primary screen is available if requested.

In regards to our animal work proposed in aim 1 of our application, we currently have generated mouse lines required to assess whether YAP deficiency (either in a heterozygous or homozygous state) can rescue tumor growth in NF2 mutant mice in the context of mesothelioma. We have already established the Adenoviral injections into the thorax of animals and using reporter strains, we have demonstrated that we can infect mesothelial tissue. Currently we are testing tumor progression in mice with the genotypes NF2f/f p53f/f that also carry wild type, f/+ or f/f alleles for YAP. The results of these experiments will be reported in our following progress report. We have been unable to obtain P0-Cre mice that carry homozygous YAP conditional alleles. The reason behind this is unclear. We believe that the P0-Cre transgene is likely located in the same chromosome as YAP.

**KEY RESEARCH ACCOMPLISHMENTS**

- Finished a kinome screen for regulators of YAP1 activity
- Published experimental results in Nature Cell Biology.
- Have identified both the LKB1 and JNK-pathways as important signaling cascades involved in YAP activity.
- Optimized high-throughput platform to screen for YAP regulators by RNA interference
- Finalized first pass of two genome-wide screens for regulators of YAP1 activity.
- Generated mouse models that will unequivocally test the requirement for YAP in NF2-driven mesothelioma.

**REPORTABLE OUTCOMES**

- We have generated several cell lines in which genetic and chemical high-throughput screens can be carried out for regulators of the NF2/Yap1 pathway. This resource will be widely beneficial to the NF2 community.
- Developed NF2 P53 YAP triple mutant animal models.
- Published experimental results in Nature Cell Biology.
- Applied and obtained R01 DK099559-01 based on results from this grant.
- PI was promoted to Associate Professor.

**CONCLUSION**

We have been successful in establishing an animal model for the evaluation of the requirement for YAP in NF2-deficient tumors. We have established multiple reporter cell lines and have developed high-throughput screening protocols that will be highly useful to identify novel regulators of the NF2/YAP1 pathway. We have finalized a targeted kinome screen, and have characterized multiple genes and pathways that are involved in YAP activation. This has been recently published in Nature Cell Biology.
APPENDIX

Mohseni et al. 2014 Nat Cell Bio (attached)
A genetic screen identifies an LKB1–MARK signalling axis controlling the Hippo–YAP pathway

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The Hippo–YAP pathway is an emerging signalling cascade involved in the regulation of stem cell activity and organ size. To identify components of this pathway, we performed an RNAi-based kinome screen in human cells. Our screen identified several kinases not previously associated with Hippo signalling that control multiple cellular processes. One of the hits, LKB1, is a common tumour suppressor whose mechanism of action is only partially understood. We demonstrate that LKB1 acts through its substrates of the microtubule affinity-regulating kinase family to regulate the localization of the polarity determinant Scribble and the activity of the core Hippo kinases. Our data also indicate that YAP is functionally important for the tumour suppressive effects of LKB1. Our results identify a signalling axis that links YAP activation with LKB1 mutations, and have implications for the treatment of LKB1-mutant human malignancies. In addition, our findings provide insight into upstream signals of the Hippo–YAP signalling cascade.

Our understanding of human disease has benefited greatly from the study of developmental pathways in model organisms. Characterization of signalling cascades such as Wnt, Hedgehog and Notch has particularly contributed to the understanding and treatment of cancer1. A more recently discovered signalling cascade is the Hippo pathway, originally described in Drosophila, and proposed to be a means by which organ size can be regulated. This pathway is highly conserved in mammals, where the mammalian hpo orthologues, MST1/2, phosphorylate the large tumour suppressor (LATS1/2) kinases, which in turn phosphorylate the transcriptional co-activator YAP, restricting its activity and stability2–4. In the absence of phosphorylation, YAP translocates to the nucleus where it binds to the TEA-domain transcription factors5,6 (TEAD1–4).

Activation of YAP, or loss of upstream negative regulators leads to striking overgrowth and tumour phenotypes in epithelial tissues, in many cases driven by the expansion of tissue-resident stem cells3,4. In addition, studies of human samples have demonstrated widespread Hippo pathway inactivation and nuclear YAP localization in multiple epithelial malignancies7–9. However, genomic analyses of common epithelial cancers have not revealed a significant rate of mutations in the known components of the pathway10. Recent data also suggest the presence of alternative kinases that might be responsible for YAP regulation9,11. Thus, common alterations of Hippo signalling in human cancer might be caused by mutations in genes not associated with the pathway at present.

Here, we have performed a genetic screen to identify kinases that impinge on the Hippo pathway. Our work uncovers kinases associated with multiple aspects of cellular function that are robust regulators of YAP localization and activity. These data provide important insight about the nature of inputs that speak to Hippo kinases. In addition, we identify the tumour suppressor LKB1 and its substrates of the microtubule affinity-regulating kinase (MARK) family as crucial regulators of the Hippo pathway. We present functional evidence suggesting that YAP is a critical component of the LKB1 tumour suppressive pathway. Our data have significant implications for the treatment of Lkb1-mutant cancers.
RESULTS

A genetic screen identifies multiple Hippo-regulating kinases

To identify potential kinases that can repress YAP/TEAD activity, we developed an improved transcriptional reporter containing 14 copies of the known TEAD DNA-binding sequence (SuperTBS reporter; Fig. 1a). Functional assays revealed that this reporter faithfully recapitulated YAP/TEAD transcriptional activity, and was highly responsive to perturbations of endogenous upstream Hippo components such as LATS2 and the cytoskeleton-associated protein NF2 (refs 12,13 and Fig. 1b). Armed with a robust reporter for Hippo–YAP activity, we interrogated the effects of a human kinome short interfering RNA (siRNA) library containing 2,130 unique short interfering RNA (siRNA) library containing 2,130 unique

Figure 1 A kinome RNAi screen identifies regulators of Hippo–YAP signalling. (a) Graphical representation of YAP-mediated STBS reporter activation in cells. (b) Validation of STBS reporter sensitivity using siRNA knockdown of known components of Hippo signalling. CTR, scrambled siRNA. n = 5 independent experiments. (c) Schematic of RNAi screening strategy. The RNAi screen was performed in 96-well plates using a stably expressing HEK293T STBS–mCherry reporter cell line. Activation of the STBS–mCherry reporter was visualized 4 days following siRNA transfection. Fluorescence intensity was captured by flow cytometry. Statistical analysis was performed to identify genes for secondary screening and final selection of hits. (d) Mean Z-score and mCherry reporter fold change (versus scrambled controls) values for each triplicate siRNA oligonucleotide were plotted to identify hits with statistical thresholds of Z-score > 2 and fold change greater than 4. Highlighted rectangle represents hits satisfying these thresholds. Green filled circles represent siRNA knockdown of LATS2 as a positive control. (e) A secondary siRNA screen identifies kinases that reproducibly raise STBS–mCherry reporter activity, performed using an alternative siRNA oligonucleotide source using two reporter systems. The secondary screen was repeated three times using pooled siRNAs. (f) YAP immunolocalization in HaCaT cells following siRNA knockdown of kinases that regulate STBS reporter activity. Representative images are shown; experiment repeated independently three times. Scale bars, 200 µm. (g) Immunoblot for Ser 127 YAP phosphorylation following siRNA knockdown of kinases from secondary screen. CTR represents scrambled siRNA and NF2 siRNA is used as a positive control. Representative blots shown; experiment repeated three times. Also see uncropped figure scan in Supplementary figures. Error bars represent ± s.d. from n = 3 biological replicates.
Figure 2 LKB1 regulates YAP activity through the Hippo kinases. (a) LKB1 knockdown induces YAP-dependent expression of target genes Amot12 and Cyr61. (b) STBS–luciferase reporter (error bars represent mean ± s.d. from n = 3 biological replicates). (c) Immunofluorescence for F-actin (red), YAP (green) and nuclei (blue) in LS174T (W4) cells. Dox-inducible LKB1 YAP nuclear to cytoplasmic translocation. Representative images shown; experiment repeated six times. (d) MST1/2 phosphorylation and quantitative PCR of Yap target genes. See uncropped figure scan in Supplementary figures. Both lines of mice also carry a p53 homozygous floxed allele. (e) Overexpression of LATS1, LATS2 and MOB1 (MLL) in LKB1-knockdown HEK293T cells can restore STBS reporter activity. MST1/2 phosphorylation and quantitative PCR of Yap target genes. See uncropped figure scan in Supplementary figures. Data represent mean ± s.d. (f) MST1/2 phosphorylation in the full-length and cleaved forms of MST1 and increase in levels of cleaved active MST1 peptide. Representative blots are shown; experiment repeated three times. Also see uncropped figure scan in Supplementary figures. (g) Activity of LATS1/2 is increased on LKB1 activation as measured by phosphorylation at Thr 1079 LATS1/2. Representative blots are shown; experiment repeated three times. (h) Ad-Cre-infected livers from Lkb1−/− mice exhibit an increase in liver size; error bars represent mean ± s.d. from n = 6 mice per group. Scale bar, 1 cm. (i) Lkb1−/− mice treated with Ad-Cre also leads to an increase in YAP target expression. Data represent mean ± s.e.m., n = 6 mice treated with Ad-Cre. Experiment was repeated in two additional mice with similar results. (j) Western blot analysis performed on liver lysates derived from Ad-Cre-infected Lkb1+/− or Lkb1−/− mice 3 months post infection leads to an overall decrease in cleaved activated Mst1 and Thr183/Thr180 Mst1/2 phosphorylation and quantitative PCR of Yap target genes. See uncropped figure scan in Supplementary figures. Both lines of mice also carried a p53 homozygous floxed allele. (k) Lkb1−/− mice treated with Ad-Cre also leads to an increase in YAP target expression. Data represent mean ± s.e.m., n = 6 mice treated with Ad-Cre. Experiment was repeated in two additional mice with similar results. (l) Overexpression of LATS1, LATS2 and MOB1 (MLL) in LKB1-knockdown HEK293T cells can restore STBS reporter activity. LATS1/2 phosphorylation compared with Lkb1 wild-type livers, n = 6 mice per group, 20 fields of view (FOV) counted for each sample in a group. Error bars represent mean ± s.d. (j) Western blot analysis performed on liver lysates derived from Ad-Cre-infected Lkb1+/− or Lkb1−/− mice 3 months post infection leads to an overall decrease in cleaved activated Mst1 and Thr183/Thr180 Mst1/2 phosphorylation and quantitative PCR of Yap target genes. See uncropped figure scan in Supplementary figures. Both lines of mice also carried a p53 homozygous floxed allele. (l) Lkb1−/− mice treated with Ad-Cre also leads to an increase in YAP target expression. Data represent mean ± s.e.m., n = 6 mice treated with Ad-Cre. Experiment was repeated in two additional mice with similar results. (m) Knockdown of Mst1/2 and Lats1/2 in Mst1/2-deficient HEK293T cells also leads to an increase in YAP target expression. Data represent mean ± s.e.m., n = 6 mice treated with Mst1/2 knockdown LATS1/2 siRNA. Scale bar, 1 cm. (n) Endogenous co-immunoprecipitation experiments using HEK293T cells demonstrate physical association of LKB1 with LATS1 and MST1. Immunoblots represent one of three independent experiments.
siRNA oligonucleotides for 710 kinase genes in a HEK293T cell line stably carrying the reporter (Fig. 1c). Initial hits were identified by a statistical Z-score cutoff of 2 in addition to a >4-fold change of mean fluorescence intensity compared with scrambled siRNA controls (Fig. 1d). Our high-stringency statistical analysis revealed 21 kinases whose silencing resulted in enhanced STBS reporter activity (Fig. 1d and Supplementary Table 1). Through a secondary screen using a different commercial source of siRNAs to control for off-target effects, we confirmed that knockdown of 16 of these kinases robustly induced STBS reporter activity (Fig. 1e). Loss of 13 of these kinases also led to YAP nuclear accumulation even in high-density conditions where Hippo signalling is typically activated (Fig. 1f and Supplementary Fig. 1a). To further characterize these hits, we evaluated their effects on YAP phosphorylation at Ser 127, as this is a highly conserved direct-substrate site for LATS1/2 and is one of the best characterized biochemical markers for Hippo-mediated YAP inactivation\textsuperscript{14}. Silencing of 8 of the 16 kinases resulted in decreases in YAP\textsuperscript{S127} phosphorylation (Fig. 1g and Supplementary Fig. 1b), indicating that some of these molecules regulate YAP activity independently of Hippo.

Interestingly, four of the validated kinase hits (MAP2K7, MAP3K9, MAP4K4, MAP4K5) are part of an activating network of the c-Jun amino-terminal kinase (JNK) branch of the mitogen-activated kinase (MAP) pathway, a stress-activated cascade implicated in compensatory growth and tumorigenesis\textsuperscript{15}. Silencing of these kinases does not lead to a reduction in YAP Ser 127 phosphorylation, indicating an alternative mode of YAP regulation (Supplementary Fig. 1b). A targeted analysis using RNA-interference (RNAi) and small-molecule manipulation confirmed that only the JNK arm of the MAP kinase pathway controlled YAP/TEAD reporter activity (Supplementary Fig. 1c,d). Although the role of JNK signalling in cancer is complex, our data support emerging findings suggesting that JNK activators are tumour suppressors, and implicate Hippo–YAP signalling as a downstream mechanism\textsuperscript{16,17}. The ephrin receptor EPHA7 (Fig. 1d–g), implicated in providing cell-positioning cues during development and mutated in lung cancer and lymphomas\textsuperscript{18,19}, also regulates YAP activity. Intriguingly, other ephrin-type A receptors (EPHA4, EPHA5 and EPHA8; Supplementary Table 2) are also found to enhance STBS activity, indicating an important crosstalk between ephrin signalling and Hippo. We also identify MAGI1 (Fig. 1e–g and Supplementary Table 1), a growth
We were particularly interested by the fact that YAP phosphorylation was significantly repressed by STK11 knockdown (Fig. 1e). Whereas YAP is predominantly nuclear at low cell densities, STK11 knockdown also resulted in loss of LATS1/2 Thr 1079 phosphorylation (Fig. 2g). Our results are consistent with a recent report indicating YAP activation in LKB1-mutant cell lines26.

LKB1 regulates YAP through MST/LATS

We were particularly interested by the fact that YAP phosphorylation was significantly repressed by STK11 knockdown (Fig. 1e–g). STK11, also known as LKB1, is a well-established human tumour suppressor that controls, among other things, cellular metabolism, proliferation and polarity24. The effect of LKB1 knockdown on YAP phosphorylation and localization was reproduced with multiple oligonucleotides and cell lines (Supplementary Fig. 2a–d). LKB1 knockdown also resulted in the upregulation of known YAP target genes, such as Amotl2 and Cyr61 (ref 6 and Fig 2a). This transcriptional response was entirely YAP-dependent, as endogenous YAP and reporter responses were suppressed in YAP/LKB1 double-knockdown cells (Fig. 2a,b and Supplementary Fig. 2e). To further demonstrate a regulatory role of LKB1 upstream of YAP we used an engineered intestinal epithelial cell line (W4) in which LKB1 activity could be induced following treatment with doxycycline (Dox). Dox-dependent LKB1-activity is evidenced by polarization and actin cytoskeleton rearrangements (Fig. 2c,d). Whereas YAP is predominantly nuclear at low cell densities, stimulation of LKB1 activity induced a striking and significant shift of YAP localization into the cytoplasm and actin cap of polarized cells (Fig. 2c,d). Consistent with this, we observed a significant reduction of YAP/TEAD transcriptional activity in Dox-treated cells (Supplementary Fig. 2f). Our results are consistent with a recent report indicating YAP activation in LKB1-mutant cell lines26.

We next determined whether LKB1 acts through the canonical Hippo kinases to regulate YAP. We observed increased MST1 activity, as measured by phosphorylation and the presence of a cleaved MST1 catalytic fragment following LKB1 activation in W4 cells (Fig. 2e). Similarly, LKB1 activation led to a marked increase in phosphorylation of Thr 1079 in LATS1/2 (Fig. 2f). This residue marks LATS1/2 activation by MST1/2 and its co-activator SAV1 (ref. 14). Correspondingly, LKB1 silencing led to loss of LATS1/2 Thr 1079 phosphorylation (Fig. 2g).

To confirm that LKB1 is important for MST1/2 activation, we used a mouse model in which Lkb1 was deleted in the liver using Ad-Cre. In agreement with MST1/2 loss-of-function phenotypes, Lkb1 deletion resulted in hepatomegaly and increased hepatocyte proliferation (Supplementary Fig. 2g). As predicted, we also observed a significant decrease in the amount of cleaved and phosphorylated MST1 peptide in Lkb1-deficient livers (Fig. 2h and Supplementary Fig. 2h) and upregulation of YAP target genes (Fig. 2i). Supporting our findings that LKB1 acts upstream of the Hippo kinases, we find that expression of LATS1/2 and its co-activator MOB1 rescues the increase in YAP/TEAD transcriptional activity following knockdown of LKB1 (Fig. 2j). Furthermore, knockdown of MST1/2 or LATS1/2 in Dox-treated W4 cells significantly suppresses the LKB1-mediated shift in YAP subcellular localization.

**Figure 4** Scribble acts downstream of LKB1 to regulate Hippo–YAP. (a) Confocal immunofluorescent and Z-stack analysis for Scribble (SCRIB, green), F-actin (red) and nuclei (blue) in LKB1 and MARK knockdown MCF7 cells. Note mislocalization of SCRIB following LKB1 or MARK silencing. Representative images from 4 independent experiments. (b) Immunofluorescence in W4 cells demonstrates that LKB1 knockdown leads to SCRIB re-localization to the cell membrane and actin cap and that this requires MARKs activity. (c) Knockdown of SCRIB in HEK293T cells reduces Ser 127 YAP phosphorylation.
Figure 5 Yap activity is enhanced in Lkb1-deficient tumours. (a) Immunohistochemistry for Yap on grade I–II lung adenocarcinomas derived from Kras-G12D mutant (K) and Kras-G12D/Lkb1fl/fl (KL) mice treated with intranasal Ad-Cre. Representative picture shown; n = 5 for each genotype. (b) Gene set enrichment analysis demonstrates significant enrichment of a transcriptional Hippo signature in KL versus K murine lung tumours. (c) Immunoblot analysis shows reduced active phosphorylated and cleaved forms of Mst1/2 in individual KL lung tumour nodules. Similarly, Ser 127 Yap phosphorylation is reduced (n = 3 mice). (d) Immunohistochemistry for Scribble localization in K and KL lung adenocarcinomas, (n = 5 mice). (e) Immunohistochemistry for Yap in pancreas from control pancreas (WT) or Lkb1-deficient tissue. (f) Yap localization assessed by immunohistochemistry in human intestinal tissue and PJS intestinal polyps. Representative data, n = 3 patients. (g) Immunohistochemistry for Yap localization and expression in normal ductal tissue compared with ductal breast adenocarcinoma (BC), and in normal human liver compared with metastatic liver adenocarcinoma derived from a PJS patient (h). Scale bars, 500 µm.

(Fig. 2k and Supplementary Fig. 2i–j). Supporting a regulatory role, we find that endogenous and overexpressed LKB1 can strongly interact with both LATS1 and MST1 in co-immunoprecipitation experiments (Fig. 2l and Supplementary Fig. 2k–l).

LKB1 acts upstream of MARKs to regulate YAP

To shed light on a possible mechanism for regulation, we performed in vitro kinase assays and mass spectrometry analyses to determine whether MST1 or LATS2 could be direct targets of LKB1. Our results found no evidence for LKB1-mediated phosphorylation at potential consensus sites in either MST1 or LATS2, thus suggesting that the LKB1 effect on these kinases was indirect. We then performed a siRNA mini-screen evaluating most known downstream targets of LKB1 (ref.27), including AMPK and mTOR, commonly implicated in growth suppression by LKB1, for their ability to regulate the STBS reporter. This screen revealed that three members of the MARK family (MARK1, 3 and 4; hereafter referred to as MARKs) were able to modulate TEAD-reporter activity (Fig. 3a). These kinases also hits in our primary kinome screen if lower hit thresholds are selected (Supplementary Table 2). The effect of MARK knockdown was reproduced across several cell types and with multiple oligonucleotides (Fig. 3b and Supplementary Fig. 3a), and its effect on TEAD-reporter activity was also suppressed with concomitant knockdown of YAP (Supplementary Fig. 3b). Loss of MARK4 also results in enhanced YAP nuclear localization (Fig. 3c), and a decrease in LATS and YAP phosphorylation (Fig. 3d). Suggesting that MARKs also act upstream of the Hippo kinases, overexpression of LATS and MOB1 can fully suppress the MARK4 knockdown effect on TEAD-reporter activity (Fig. 3e and Supplementary Fig. 3c). To ascertain whether MARKs were functionally downstream of LKB1, we knocked down MARKs in LKB1-induced W4 cells. Dox addition to W4 cells leads to MARK1 activation27 (Supplementary Fig. 3d), and silencing of MARKs in this context resulted in a significant loss of cytoplasmic YAP translocation (Fig. 3f and Supplementary Fig. 3e–f). Combined, these data demonstrate that LKB1 is exerting its effects on the Hippo pathway through its direct substrate, the MARKs.
MARKs regulate SCRIB localization and Hippo kinase activity

MARKs are also known as the PAR-1 family of proteins and have been implicated in the regulation of cell polarity and microtubule dynamics through different mechanisms\(^3\). In *Drosophila*, the PAR-1 orthologue has been shown to phosphorylate and regulate localization of Discs large\(^2\) (DLG), a member of the basolateral polarity complex also consisting of Lethal giant larvae (LGL) and Scribble\(^3\) (SCRIB). Proper localization of SCRIB is required for Hippo pathway activity in both *Drosophila* and mammalian cells\(^2\)–\(^4\). Thus, we posited that LKB1 could be regulating Hippo–YAP activity through regulation of the basolateral polarity complex by the MARKs. Indeed, we find that MARKs knockdown results in mislocalization of SCRIB (Fig. 4a and Supplementary Fig. 4a), and reduction of SCRIB protein (Supplementary Fig. 4b–c). Demonstrating a direct role for LKB1 and MARKs in the localization of SCRIB, Dox-mediated activation of LKB1 in W4 cells results in SCRIB recruitment to the cellular membrane and the actin cap (Fig. 4b). Knockdown of MARKs in this context reduces the sub-cellular localization shift of SCRIB (Fig. 4b). As predicted, SCRIB knockdown also leads to an increase in TEAD-reporter activity and a decrease in YAP phosphorylation (Fig. 4c and Supplementary Fig. 4d). Importantly, knockdown of SCRIB in LKB1-activated W4 cells significantly rescues the shift of YAP localization to the cytoplasm and actin cap (Fig. 4d and Supplementary Fig. 4f–h), indicating that SCRIB is critical for LKB1-mediated regulation of YAP. Moreover, co-immunoprecipitation experiments demonstrate that endogenous MARK1 or overexpressed MARK4 can be detected in a complex with LKB1, MST1, LATS1 and SCRIB (Fig. 4e and Supplementary Fig. 4i), indicating the existence of a Hippo regulatory protein complex. It has been proposed that association of SCRIB with MST1/2 is important for the activation of the Hippo cascade\(^4\). We find that this association is highly dependent on MARKs (Fig. 4f and Supplementary Fig. 4j), as their loss impairs the interaction of both MST1/2 and LATS1/2 with SCRIB.

YAP activation is a hallmark of LKB1-mutant tumours

Lkb1 germline mutations are associated with Peutz–Jeghers syndrome (PJS), an inherited disorder in which patients develop intestinal polyps and are at higher risk for developing multiple malignancies\(^5\). Lkb1 alterations are also present in many types of sporadic epithelial cancer, particularly lung and pancreatic carcinomas\(^6\). Loss of Lkb1 in mice is associated with more aggressive and metastatic potential of lung tumours\(^7\). To corroborate our *in vitro* observations, we evaluated the status of Hippo signalling in lung tumours derived from mice carrying an activating K-Ras mutation (K) or the K-Ras transgene and concomitant Lkb1 deletion (KL). Strikingly, we find that stage-matched KL adenocarcinomas were strongly positive for nuclear YAP in contrast to K tumours, which exhibit predominantly cytoplasmic and diffuse YAP localization (Fig. 5a). To further assess the extent of YAP transcriptional activity in *Lkb1*-null tumours, we carried out gene set enrichment analysis to examine the enrichment of a YAP transcriptional signature derived in our laboratory (Supplementary Fig. 5a). Gene set enrichment analysis demonstrates...
**Figure 7** YAP is essential for the growth of Lkb1-mutant tumours and tissue. (a) Four-week soft-agar assay using the Lkb1-deficient lung adenocarcinoma line A549. These cells expressed a Dox-inducible shRNA against YAP (iYAP shRNA). Scale bar 100 µm. (b) Representative images of metastatic lesions following intravenous injection of parental or iYAP shRNA A549 cells. Dox treatment of hosts was carried for 2 months at which time lung tissue was collected. Scale bars, 200 µm. (c,d) Ad-Cre-mediated deletion of a conditional allele of Yap1 following Ad-Cre intravenous administration leads to significant suppression of hepatomegaly and hepatocyte hyperplasia. Scale bar, 1 cm. Animals received Ad-Cre at 1 month of age and tissues were collected 2.5 months later. (e) PCNA immunohistochemistry on Ad-Cre-treated mouse livers from wild-type, Lkb1 and Lkb1/Yap1 mutant mice. Scale bars, 500 µm. n = 5 mice per genotype. (f) Quantification of the average number of PCNA-positive cells per field of view from e. n = 5 mice per genotype, 20 fields of view. Error bars represent ±s.d. from n = 5 mice. *P < 0.05, two-tailed t-test.

YAP is functionally important downstream of LKB1

We next investigated functionally whether YAP acted downstream of LKB1 in tumour suppression. Using W4 cells, we found that inducible LKB1 activation has a powerful growth suppressive function in vitro (Fig. 6a and Supplementary Fig. 6a), and in xenografts (Fig. 6b and Supplementary Fig. 6b). However, expression of a YAP-S217A mutant protein is able to significantly overcome all of LKB1 tumour suppressive effects (Fig. 6a,b and Supplementary Fig. 6a,b). Silencing of either LATS2 or SCRIB also rescues growth suppression by LKB1 activation (Fig. 6d–f and Supplementary Fig. 6c,d). To determine whether we could reverse the effects of LKB1 loss by manipulating YAP-expression levels, we developed a Dox-inducible YAP short hairpin RNA (shRNA) A549 cell line (Supplementary Fig. 7a). A549 is a lung cancer cell line mutant for LKB1 widely used in tumour growth and metastasis assays. In both a soft-agar colony-formation assay, and in vivo metastatic assays, we find that YAP depletion following Dox-treatment reduces the number and/or size of colonies and tumours (Fig. 7a,b and Supplementary Fig. 7a–c). Lung adenocarcinoma cell lines that are wild type for LKB1 and expressed lower levels of YAP were...
insensitive to YAP modulation (Supplementary Fig. 7d–f). Finally, we used Ad-Cre infection in mice to demonstrate that conditional deletion of Yapi suppresses the liver overgrowth phenotype (Fig. 7c,d) and hepatocyte hyperplasia observed following acute deletion of Lkb1 (Fig. 7e,f and Supplementary Fig. 7g). Together these data provide multiple lines of evidence that Hippo–YAP is a functionally critical pathway downstream of LKB1.

**DISCUSSION**

One important question in the Hippo–YAP field relates to the upstream signals that regulate the Hippo kinases. Our studies here have identified many molecules and pathways that might impinge on Hippo activity and growth control. As many of these kinases are also mutated in human cancer, their identification as regulators of YAP might provide a molecular explanation for the observations that YAP is highly active in numerous epithelial tumours, where mutations in the canonical Hippo components are not found.

The tumour suppressive function of LKB1 has primarily been linked to its ability to regulate cellular metabolism through AMPK activation37. LKB1 is linked to mTOR through the sequential activation of AMPK and the tumour suppressor TSC2, whose activation leads to suppression of mTOR activity38. It has been shown that polyps from FAP patients show upregulated mTOR activity, as do pancreata, cardiomyocytes and endometria of Lkb1-deficient mice. Treatment of endometrial LKB1-mutant adenocarcinomas with rapamycin and mTOR inhibitor, leads to regression of these tumours, supporting a functional role for mTOR downstream of LKB1 (ref. 38). Our studies here suggest that LKB1 can also exert its tumour suppressive effects through activation of a PAR-1-mediated polarity axis that controls the Hippo signalling pathway. Our data demonstrating that YAP loss could completely rescue growth phenotypes mediated by LKB1 loss in vivo suggest that this might a central mechanism. On this note, it has been shown that YAP can lead to mTOR activity through transcriptional activation of miR-29. Thus, YAP activation due to LKB1 alterations could also lead to mTORC1 activation.

Our data provide insight into a signalling axis downstream of LKB1 and PAR-1 kinases that regulates the interaction of the Hippo kinases with SCRIB and perhaps other components of the basolateral polarity complex. MARKs can also lead to changes in polarity by antagonizing the PAR-3/PAR-6 polarity complex39. This complex is localized apically whereas PAR-3 lacking PAR-1 phosphorylation results in ectopic lateral mislocalization. Under normal conditions, the lateral exclusion of PAR-3/PAR-6 by PAR-1 also cooperates with Crumbs to restrict PAR-3 localization, and loss of both pathways disrupts epithelial polarity39.

The literature supports that the Hippo pathway is indeed regulated by these polarity complexes42–48. Whether PAR-3, PAR-6, Crumbs and other substrates of Par-1/MARKs are also involved in controlling SCRIB remains to be investigated. Similarly, a connection between Hippo–YAP signalling and the actin cytoskeleton has recently been demonstrated49. Considering that LKB1 and SCRIB have effects on the actin cytoskeleton42,43, it is possible that actin fibre regulation could be an additional mechanism by which LKB1 modulates YAP activity. LKB1 is then a candidate upstream regulator of the multiple inputs that impinge on YAP activity. Collectively, these data suggest that manipulation of the Hippo signalling pathway should now be evaluated for the treatment of LKB1 mutant cancers.

**METHODS**

Methods and any associated references are available in the online version of the paper.

**Note: Supplementary Information is available in the online version of the paper.**

**ACKNOWLEDGEMENTS**

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**AUTHOR CONTRIBUTIONS**


**COMPETING FINANCIAL INTERESTS**

The authors declare no competing financial interests.

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METHODS

RNAi screen. An RNAi library against all known kinases in the human genome was used (Ambion Silencer Select Human Kinase siRNA Library, catalogue # 4397918). Reverse transfection using 0.1 µl of transfection reagent (RNAi max, Invitrogen) and siRNA of a final concentration of 5 nM was performed on approximately 10,000 HEK293T TBS-m-Cherry cells per well of a 96-well plate. We estimate that close to 98% of HEK293T cells are transfected using this method. The screen was performed in triplicate, with 3 oligonucleotides for each gene (Fig. 1d). To ensure limited edge effects, outer rows and columns were not used and instead were occupied by cell media. At 96 h post-transfection, HEK293T cells in negative control wells are confluent. Plates are subsequently trypsinized in 20 µl of trypsin/EDTA and re-plated in 30 µl of DMEM/10% FBS and analysed by Flow cytometry using Texas red and GFP (HTS, BD LSRii) to obtain the mean fluorescent intensity of each cell. Data were collected and analysed using FACS Diva 6.0 (BD Biosciences). For follow-up work, individual oligonucleotides targeting YAP (523066), LATSI (17392), LATSI2 (523560), MST1 (13570), MST2 (523667), SCRB2 (523970), MARK1 #58512, MARK1 #53718 and NF2 (194647), all from Ambion, were used.

Hit selection. Positive hits for each gene were identified as follows. Z-scores and fold changes were calculated for each oligonucleotide when compared with the negative control for each individual 96-well plate. Rigorous hit selection was performed by eliminating data that did not reproduce in at least two out of the three experiments for each oligonucleotide. Subsequently, these data were further filtered to identify oligonucleotides that reproducibly have a Z-score > 2 and fold change > 4. The final hit selection is based on the mean Z-score values and mean fold changes of each oligonucleotide for each gene, and if two or more oligonucleotides for each gene meet these thresholds.

Immunofluorescence. Cells were seeded in a 24-well plate on sterilized glass coverslips overnight and then fixed in 4% paraformaldehyde/PBS for 10 min at room temperature, followed by three washes in PBS. Cells are permeabilized in 0.01% Triton/PBS for 1 min, followed by three washes in 0.01% Tween/PBS. Cells were then incubated in blocking buffer (0.5% FBS/PBS/0.01%Tween) for 1 h at room temperature and then incubated overnight at 4°C in primary antibody (YAP 1:1,000, Cell Signaling #4912; Scribble 1:1,000, Santa Cruz #1049; GFP 1:500, Abcam #ab290) in blocking buffer. Coverslips were washed three times in 0.01%PBS Tween and then incubated in secondary antibody and/or Alexa-fluor Phalloidin of each oligonucleotide for each gene, and if two or more oligonucleotides for each gene meet these thresholds.

Immunohistochemistry. p53/p16, p53/Lkb1 mouse livers and KrasG12D/Lkb1 mice were collected and fixed in neutral buffered formalin (4%; pH 7.0; 16-24 h, 20°C), and then switched to 70% ethanol after 18-24 h. Tissues were embedded in paraffin and 5 µm sections were mounted on positively charged slides. Tissues were deparaffinized and antigen retrieval was performed in pH 6.0 citrate buffer for 30 min at 95°C. ABC tissue staining was performed by using a modified protocol from Vector Laboratories VectorStain Elite ABC kit (#PK-6101, #PK-2200). Briefly, the endogenous peroxidase was blocked by incubation in 0.3% hydrogen peroxide for 1 min. Sections were blocked in rabbit sera followed by primary antibody incubation overnight at 4°C (YAP 1:1000, Cell Signaling #4912; Scribble 1:400, Santa Cruz #1049; K67 1:50, DAKO MIB-5). Sections were washed and then incubated with HRP-conjugated secondary antibody (30 min; 20°C), and developed with 3,3′-diaminobenzidine tetrachloride (DAB)/H2O2. Counterstaining was done with haematoxylin and samples were washed, dehydrated, and mounted with VECTaMount (Vector Labs #H-5000). Data obtained for immunohistochemistry analyses were repeated using sections from the same tissue and/or from amongst the same genotype group. Representative images shown for each gene exhibit the least signal/noise background and stains that represent most of the tissues that were analysed; however, there are some tissues that were stained that exhibited either more intense or weaker staining, which are not shown as they represented the minority of tissues examined.

Immunoblotting. Cell lines and tissues were collected, and processed for western blotting by solubilizing extracts in lysis buffer (50 mM Tris, 100 mM NaCl, protease inhibitor cocktail (Roche #04693159001) and phosphoSTOP (Roche, # 04906837001)). For standard immunodetection of proteins, 20 µg of protein was used. For detection of phospho-proteins, 30 µg of total protein lysate was used. Protein lysates were then resolved by PAGE under reducing conditions (4-12% SDS-PAGE Bis-Tris gels; MOPS buffer system; Invitrogen; NuPAGE-MOPS system). The gels were blotted onto PVDF or nitrocellulose papers and blocked in either milk for standard antibodies or BSA for phospho-antibodies (phospho-antibodies blocked in PBS, 5% w/v BSA, 0.1% Tween-20 at dilutions: pYAP 1:1000; Cell Signaling #4911; pMST1/2 1:1000; Cell Signaling #3681; pLATS1/2 1:1000; Cell Signaling #1513; pMARK1/2 1:500; Cell Signaling #2575; Cell Signaling #4834; standard antibodies blocked in TBS-T, 5% w/v milk at dilutions: MST1 1:500, Cell Signaling #3682; Lats1 1:1000, Cell Signaling #3347; MARK1 1:1000 Cell Signaling #4834; LKB1 1:1000 Santa Cruz sc-32245; Scribble 1:500, Santa Cruz sc-11049; MARK1 1:100 Cell Signaling #3319; AMPK 1:1200, Cell Signaling #2650) for 1 h at room temperature, followed by incubation in primary antibodies diluted in blocking buffer. Unless otherwise stated, all primary antibody incubation steps were performed overnight at 4°C. After washing in TBS-T, antibodies were detected using HRP-conjugated secondary antibodies (1:2000 in TBS-T; Thermo #52340, #32460, Santa Cruz sc-2020), and visualized using enhanced chemiluminescence (Thermo, #34096). Immunoblots shown are representative of experiments that were reproduced at least three independent times. For some challenging experiments and antibodies, the representative blots are ones that show the least nonspecific background and have a low signal-to-noise ratio.

Cell lines. LS174T, HEK293T, DLD1, MCF7 and HcaCt cells were cultured in DMEM + 10% FBS in 5% CO2, >95% humidity. AS49 cells were cultured in RPMI-1640 supplemented with 2 mM l-glutamine and 10% FBS. The LS174T–W4 clone cell line was a gift from H. Clevers (Utrecht Institute, Netherlands). Other cell lines were acquired from the American Type Culture Collection (Manassas).

Soft-agar colony-formation assay. The base agar consisted of low-melting-point 0.6% agar dissolved in RPMI-1640 (Life Technologies, # 31800-022) or DMEM (Life Technologies #12100046), 10% FBS and 1% penicillin/streptomycin. Base agar was allowed to set for at least 1 h before plating of the top agar. The top agar consisted of approximately 250 cells resuspended in 0.3% low-melting-point agar dissolved in RPMI or DMEM, 10% FBS, 1X penicillin/streptomycin in a well of a 6-well plate. Samples were incubated for 4 weeks following seeding and then stained with 0.1% crystal violet in 10% ethanol for 20 min. Wells were destained in distilled water or 5 times or until the decanted water ran clear before imaging.

In vitro proliferation assay. A colorimetric MTS assay (Promega, # G5430, Madison) was used to determine the proliferation rate for different cell lines. Experiments were done following the manufacturer’s instructions. Briefly, 1,000 cells per well were cultured in triplicate in 96-well plates and incubated for 0–7 days. At the time of culture and each day for a total of 7 days, a plate was analysed by colorimetric reading (absorption of light at 450 nm).

Mouse models. Animal work was approved by the institutional committee at Boston Children’s Hospital. Animals were housed in specific pathogen-free facilities at the hospital. The mouse models p53fl/flox (JAX Labs, B6.129P2-Tp53null/J), Lkb1fl/flox (ref. 36), LSL-KRASG12D (JAX Labs, 129S/Sv-KrasG12D1Crt), Yapfl/fl (ref. 11), PDX-Cre have been previously described44. For all experiments involving Ad-Cre-mediated deletion, female mice of approximately 5–6 weeks of age were used. Ad-Cre administration was performed between 2–4 weeks of age. Xenograft assays were performed in 5-week-old male Nu/ mice (JAX Labs, B6.Cg-Fsgrtm1WLe/J), using 1 × 106 cells 100 µl-1 volume of Matrigel (BD Biosciences). Metastasis assays were performed in 5-week-old male NOD/SCID mice (JAX Labs, NOD.CB17-Prkdcscid/J), using 1 × 106 cells 100 µl-1 volume of PBS injected intravenously. For induction in the liver, 100 µl of Ad-Cre was introduced intravenously at 1 × 106 pfu per mouse (AdCMV-Cre, University of Iowa, GTVC). No statistical method was used to predetermine sample size for treatment groups. There was also no requirement for animal randomization during the course of the animal studies.

Tissue samples. We studied formalin-fixed paraffin-embedded tissue-biopsy sections of diagnosed PIS and juvenile polyposis patients with confirmed mutations in LKB1, SMAD4 and PTEN respectively. Studies with patients’ samples at Boston Children’s Hospital were covered under IRB-CR409-12-0660.

Small-molecule inhibitors. Metformin (10 mM) was from EMD Millipore, and AICAR (2 mM) and rapamycin (1 mM) were from TOCRIS Bioscience. MAP Kinase
METHODS

Signaling Pathway Inhibitor Panel was purchased from EMD Millipore (#444189) and incubated according to the manufacturer’s recommended concentrations (FR180204: IC50 = 510 nM, JNK II: IC50 = 40 nM for JNK-1 and JNK-2 and 90 nM for JNK-3, JNK IX: pIC50 = 6.5, MEK1/2: IC50 = 220 nM, MK2a: IC50 = 60 nM, Inhibitor 5: IC50 = 40 nM, PD98059: IC50 = 10 µM, Inhibitor IV: IC50 = 10 nM, SR203580: IC50 = 600 nM, TPL2 Inhibitor: IC50 = 50 nM, ZM336372: IC50 = 70 nM).

Luciferase assays. For stable cell lines expressing STBS–luciferase, cell lysates were analysed using Dual-glo luciferase assay (Promega, #E2940). Alternatively, cell lines not containing the reporter were transiently transfected with the STBS-firefly-luciferase and firefly–Renilla constructs using Lipofectamine 2000 (Life Technologies) and assayed 48 h following plasmid transfection (Berthold Technologies).

Microarray and gene set enrichment analysis. Reverse transfection of RNAi oligonucleotides for NF2, LATS2, YAP, TAZ and Scrambled RNAi was performed in HEK293T, HaCaT and DLD1 cells with RNAiMAX (Invitrogen) according to the manufacturer’s instructions. Duplicate samples were prepared for each condition. Four days after transfection, confluent cell culture was collected for RNA extraction with the Trizol reagent (Invitrogen). RNA quality assessment, cDNA synthesis, probe generation, array hybridization and scanning were carried out by Boston Children’s Hospital Molecular Genetics Microarray Core Facility. Data sets were analysed with the online microarray analysis software GenePattern (Broad Institute) with default settings. Differentially expressed genes were defined as those with at least a twofold change in NF2/LATS2 double-knockdown cells and a P value smaller than 0.05. To generate a generic Hippo target gene signature, genes upregulated in all three NF2/LATS2 double-knockdown cell lines were combined, and those without a gene symbol were eliminated from the list. For gene set enrichment analysis, we used a data set from published gene expression profiles of lung adenocarcinomas developed in KrasG12D, KrasG12DLkb1f/- or KrasG12DLkb1f/f mouse models46. The enrichment analysis was performed in the GSEA software available from the Broad Institute with the default settings.

Microarray accession numbers. Microarray data generated for this study have been deposited in the GEO database under accession number GSE49384. The published microarray data set46 re-analysed in this study is available from the GEO database under accession number GSE6135.

Statistical analyses. No statistical method was used to predetermine sample size. For experiments using LS174T-W4 cells, we always included a −/+ Dox control to detect the level of activity of LKB1 in inducing cell polarity. Experiments were discarded in which Dox administration had less than 50% affect on cell polarization. For biochemical experiments, we performed experiments at least three independent times to be confident in the experimental reproducibility. In some cases, experiments were repeated more than 3 times to ensure validity of our hypotheses. Investigators were not blinded to allocation during experiments and outcome assessment. All statistical analyses were performed by SAS program (Version 9.1, SAS Institute). All P values were two-sided and statistical significance was set at P = 0.05. For categorical data, the χ2 test was performed. For the experiments shown, the variance was similar between groups that were being statistically compared.

A genetic screen identifies an LKB1–MARK signalling axis controlling the Hippo–YAP pathway

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In the version of this Article originally published, the name ‘Kwok-Kin Wong’ was spelled incorrectly in the author list. This has now been corrected in all online versions of the Article.
Supplementary Figure 1 Identification of kinases that can modulate the Hippo Signaling Pathway in vitro. (a) Immunofluorescence for YAP localization in confluent HaCaT cells following siRNA transfection of selected kinase hits. Scale bars, 200μm. Experiment was repeated 3 times independently (b) Western Blot for YAP S127 phosphorylation, and total YAP in 293T cells four days following siRNA transfection of selected kinase hits. Representative blot is shown. This experiment was repeated 3 independent times (c) Mitogen activated kinase pathway siRNA mini-screen using the STBS-luciferase reporter in 293T cells compared to scrambled control (CTR). Data shown is from technical triplicates. The experiment was repeated in three independent experiments. 5 bD) Selective JNK small molecule screen using the STBS-luciferase reporter in 293T cells. Schematic demonstrates where the inhibitors function. Fold changes calculated are compared to untreated controls.
**Supplementary Figure 2** LKB1 acts upstream of the Hippo Signaling Pathway in vitro and in vivo. (a) Knockdown of LKB1 using multiple different siRNA oligos increases STBS-luciferase reporter activity in 293T cells. n= 5 biological replicates ± SD. (b) Knockdown of LKB1 reproducibly increases STBS-luciferase reporter activity across various cell lines. n=3 biological replicates. Error bars represent mean ± SD. (c) Knockdown of LKB1 reproducibly decreases YAP S127 phosphorylation across various cell lines by western blot analysis. (d) Knockdown of LKB1 in DLD1 cells promotes YAP nuclear localization at confluent cell densities. Experiment has been performed independently three times. Scale bar, 200 μm (e) qPCR validation of LKB1 and YAP siRNA knockdown in 293T cells. n=3 biological replicates, ± SD. (f) LKB1 activation in W4 cells repressed TEAD-reporter activation. n= 3 replicates per biological triplicate (g) Decreased activity of MST1/2 in LKB1 deficient livers. Ratio of cleaved MST1 versus full length MST is shown for an average from n=4 mice. (h) Quantification of YAP localization in W4 cells following MST1/2 or LATS1/2 knockdown. Data are derived from three independent experiments where at least 300 cells where scored. N = 3, error bars represent mean ± SD. (i) Validation of MST1/2 and LATS2 downstream of LKB1 using two additional sets of siRNAs. Representative figure from three independent experiments is shown. Scale bars, 20 μm (j, k) Immunoprecipitation of overexpressed LKB1, LATS1, and MST1 in 293T cells. Representative blot is shown. Experiment has been performed three times independently. Error bars represent mean ± SD from triplicate samples. **, P≤ 0.01, two-tailed t-test.
**Supplementary Figure 3** Yap1 activity and localization is dependent on the LKB1 substrate, MARKs. (a) qPCR validation of siRNA knockdown of MARK1 and MARK4 and expression of Yap-target genes, CTGF and CYR61. n=3 biological replicates (b) STBS reporter activity following knockdown of MARK4 and Yap in 293T cells. n=3 biological replicates (c) Western blot showing expression level of Mob1/LATS1/LATS2 in MARK4 knockdown cells. Representative blot is shown from three independent experiments (d) Activation of LKB1 in W4 cells promotes activation of MARKs as measured by Thr215 phosphorylation (MARK1) in the kinase activation loop. Representative blot is shown from three independent experiments (e) Immunofluorescence for Yap localization (green) and cell polarization (red) in doxycycline-untreated W4 cells following knockdown of MARK4. N= 3 biological replicates. Each experiment was performed with technical triplicates. Scale bar, 20\(\mu\)m. (f) Yap localization in W4 cells following LKB1 activation and knockdown of MARK4 using 2 independent siRNA’s. Three independent experiments were performed. Scale bars, 20\(\mu\)m. (g) Biochemical analysis of phospho-Yap localization by subcellular fractionation of W4 cells treated with doxycycline and MARK knockdown. Representative blot from three independent experiments is shown. Error bars represent mean ± SD.
Supplementary Figure 4 Scribble expression and correct localization is required for Yap1 activity. (a) Immunofluorescence for Scribble (SCRIB) localization (green) in DLD1 cells following LKB1 and MARK siRNA knockdown. Experiment was repeated three times. Scale bar, 200 μm. (b-c) Immunoblot and qPCR for SCRIB expression following LKB1 and MARK1 knockdown in MCF7 cells. n=3 biological triplicates (d) siRNA knockdown of SCRIB in 293T cells induces TEAD-reporter activity. n = 3 independent experiments. (e) qPCR validation of SCRIB knockdown using 3 independent siRNA’s. n= 3 independent experiments. (f) Immunofluorescence for YAP localization (green) and cell polarization (red) in doxycycline-untreated W4 cells following knockdown of SCRIB. Biological replicates were performed three times. Scale bar, 20 μm. (g) Quantification of YAP localization in W4 cells following scribble knockdown. Data are derived from three independent experiments where at least 300 cells were scored. (h) Immunofluorescence for YAP localization following activation of LKB1 and knockdown of scribble using 2 independent siRNA’s. Scale bar 20 μm. (i) Immunoprecipitation of overexpressed MARK4 with SCRIB, LKB1, MST1, and LATS1. Experiment was repeated three times. (j) Immunoprecipitated SCRIB in MARK1-knockdown cells show decreased interaction with MST and LATS. Experiment was repeated three times. Error bars represent mean ± SD.
**Supplementary Figure 5** Generation of a Hippo Gene Expression Signature and localization of Yap in human LKB1-mutant tissues. (a) Microarray analysis on three different cancer cell lines reveal a core set of genes whose expression changes following siRNA knockdown of NF2 + LATS1/2, and for which this response is dependent on YAP/TAZ. Differentially expressed genes were defined as those with at least 2 fold change in NF2/LATS2 double knockdown cells and a p value smaller than 0.05. (b) YAP immunohistochemistry on human SMAD4 juvenile polyposis (JP) intestinal polyp compared to a human LKB1 mutant Peutz-Jeghers (PJ) intestinal polyp. Scale bar, 500 μm.
**Supplementary Figure 6** Yap1 acts downstream of LKB1 and can overcome LKB1 tumor suppressive function. (a) Quantification of triplicate samples of number and size of W4, W4TetOYAP, +/- doxycycline colonies in soft agar assays. n= 3 biological triplicates (b) Tumor weights of W4, W4TetOYAP, +/- doxycycline after seven weeks. Representative images of nude mice carrying xenografts with W4, W4TetOYAP tumors. N = 7 mice per each of the four genotypes. (c) qPCR validation of shRNA knockdown of LATS2 and activation of YAP-target genes in W4 cells. N = 3 biological replicates. (d) qPCR of Scribble following Scribble siRNA knockdown in the presence of LKB1 activation. n= 3 biological replicates. Error bars represent mean ± SD.
Supplementary Figure 7 Loss of Yap1 in LKB1 mutant xenograft show tumor growth inhibition. (a) Quantification of triplicate samples of number and size of W4, W4TetOYAP, +/- doxycycline colonies. n=3 biological replicates (b-c) Average tumor size and number per mouse following intravenous injection of 1x10^6 A549/YAPshRNA cells (-/+ doxycycline) for 6 weeks. n=3 mice per group (d-f) Two cell lines that are low in YAP target gene expression were infected with shYAP1 hairpins and assessed for proliferation using MTS assays (e) and soft agar colony formation assay (f) n = 3 biological replicates. Scale bar, 500 μm. (g) Expression levels of LKB1 and YAP following Ad-cre administration in Lkb1-floxed and Lkb1/YAP floxed livers. n=4 mice per genotype. Error bars represent mean ± SD.
Supplementary Figure 8 Full scans