Adhesion to fibronectin regulates Hippo signaling via the FAK–Src–PI3K pathway

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The Hippo pathway is involved in the regulation of contact inhibition of proliferation and responses to various physical and chemical stimuli. Recently, several upstream negative regulators of Hippo signaling, including epidermal growth factor receptor ligands and lysophosphatidic acid, have been identified. We show that fibronectin adhesion stimulation of focal adhesion kinase (FAK)-Src signaling is another upstream negative regulator of the Hippo pathway. Inhibition of FAK or Src in MCF-10A cells plated at low cell density prevented the activation of Yes-associated protein (YAP) in a large tumor suppressor homologue (Lats)-dependent manner. Attachment of serum-starved MCF-10A cells to fibronectin, but not poly-d-lysine or laminin, induced YAP nuclear accumulation via the FAK–Src–phosphatidylinositol 4,5-bisphosphate 3-kinase (PI3K) signaling pathway. Attenuation of FAK, Src, PI3K, or PDK1 activity blocked YAP nuclear accumulation stimulated by adhesion to fibronectin. This negative regulation of the Hippo pathway by fibronectin adhesion signaling can, at least in part, explain the effects of cell spreading on YAP nuclear localization and represents a Lats-dependent component of the response to cell adhesion.

Introduction

Contact inhibition of proliferation (CIP) was originally defined as inhibition of cell division when cells reach their stationary density despite periodic nutrient renewal (McClatchey and Yap, 2012). In a dynamic tissue microenvironment, however, CIP is determined not only by postconfluent cell density but also by the quantitative interplay between cell–cell contacts, mitogens, and ECM. Increased cell–cell contact elevates the threshold level of EGF to overcome CIP (Kim et al., 2009). In addition, matrix stiffening dramatically reduces the threshold for responding to EGF (Kim and Asthagiri, 2011). The balance among these environmental cues is crucial in development, tissue regeneration, and organ size control.

The Hippo pathway has been implicated in the regulation of CIP (Gumbiner and Kim, 2014; Johnson and Halder, 2014). This growth inhibitory signaling pathway consists of a highly conserved kinase cascade leading to the activation of Lats (large tumor suppressor homologue) kinases, which control the nuclear exclusion and inactivation of transcriptional co-activator YAP (Yes-associated protein) and its paralogue TAZ (transcriptional coactivator with PDZ-binding motif). When YAP/TAZ are translocated into the nucleus, they interact with TEAD (TEA domain family member) DNA-binding transcription factors to transcribe growth-promoting and antiapoptotic genes (Zhao et al., 2008). YAP/TAZ are also known to interact with other transcription factors including p73, ErbB4, Smads, and FBJ murine osteosarcoma viral oncogene homologue to activate various target genes (Basu et al., 2003; Komuro et al., 2003; Varelas et al., 2010; Shao et al., 2014). Several physiological upstream regulators created by cell–cell contact (cadherin–catenin complex, polarity proteins, and tight junction proteins) are known to positively regulate the Hippo pathway (Kim et al., 2011; Gumbiner and Kim, 2014). The physical properties of cells, such as cell shape, ECM elasticity, and cytoskeletal tension, also play a role in controlling the Hippo pathway (Halder et al., 2012; Gumbiner and Kim, 2014). This mechanotransduction pathway may regulate YAP/TAZ activity independently of the Lats kinases, but through Rho–RhoC–dependent actomyosin contractility (Dupont et al., 2011; Aragona et al., 2013; Calvo et al., 2013; Low et al., 2014).

Recently, mitogens including insulin, EGF, lysophosphatidic acid (LPA), and sphingosine 1-phosphate as well as proteases such as thrombin have been identified as negative regulators of the Hippo pathway leading to YAP/TAZ nuclear activity (Miller et al., 2012; Mo et al., 2012; Straßburger et al., 2012; Yu et al., 2012; Fan et al., 2013). We previously reported that treatment with EGF, LPA, or serum inhibits Hippo signaling through the activation of the PI3K (phosphatidyli-
nositol 4,5-bisphosphate 3-kinase)–PDK1 (3-phosphoinositide–dependent protein kinase 1) pathway (Fan et al., 2013). PDK1 forms a complex with the Hippo signaling core complex, and EGF signaling blocks the complex formation in a PI3K–PDK1-dependent manner, leading to the activation of YAP by dephosphorylation and nuclear accumulation. We wondered whether other classes of upstream regulators of PI3K–PDK1 signaling could similarly regulate the Hippo pathway. In this study, we identified the stimulation of FAK–Src–PI3K by adhesion to fibronectin as an upstream regulatory branch of the Hippo pathway, which controls the activity and subcellular localization of YAP in a Lats-dependent manner.

Results

PI3K, PDK1, and Src control YAP subcellular localization

In our previous study, we found that PI3K–PDK1 signaling in response to growth factors inhibits the Hippo pathway (Fan et al., 2013). PI3K and PDK1 inhibitors prevented growth factor–stimulated YAP nuclear localization in confluent MCF-10A cells at low concentrations expected for specific effects on these enzymes (Fan et al., 2013). In subconfluent MCF-10A cells, YAP is also localized in the nucleus even under starvation conditions without any growth factors, which is enhanced by the depletion

Figure 1. PI3K, PDK1, and Src regulation of nuclear YAP via Lats in serum-starved, subconfluent cells. [A] PI3K and PDK1 inhibitors relative to Lats. MCF-10A cells transfected with control, Nf2, or Lats1/2 siRNAs were serum starved and treated with DMSO (solvent control), 10 µM wortmannin (PI3K inhibitor), or 5 µM BX-795 (PDK1 inhibitor) for 30 min. On-target plus nontargeting pool was used as a control siRNA. Localization of endogenous YAP was identified by immunofluorescence staining. [B] SFK inhibitors and YAP localization. Serum-starved, low cell density MCF-10A cells were incubated with SFK inhibitors (10 µM each of PP2, dasatinib, SKI-1, and SU6656) for 30 min. 10 µM each of PP3 and imatinib were used as controls. YAP subcellular localization was determined by immunofluorescence staining. Alexa Fluor 594 secondary antibody was used for SU6656, which has high background green fluorescence. (C) Biochemical effects of Src inhibition. PP3- or PP2-treated MCF-10A cells were analyzed by Western blot using anti-YAP and anti–phospho-YAP (S127) antibodies. Phosphorylated YAP was detected by mobility shift on Phos-tag SDS-PAGE. (D) SFK inhibitors relative to Lats. MCF-10A cells transfected with control, Nf2, or Lats1/2 siRNA were serum starved and treated with 10 µM PP3 or PP2. After 30 min, cells were fixed for immunofluorescence staining with anti-YAP antibody. (E) Depletion of individual SFK. MCF-10A cells were transfected with control, Src, Fyn, or Yes siRNA. After serum starvation, subcellular localization of endogenous YAP was identified by immunofluorescence staining and quantified based on the criteria shown under the graph. More than 120 cells from four random views were quantified. [F] Src knockdown relative to Lats. MCF-10A cells were transfected with control, Src, Lats1/2, or combined siRNA of Src and Lats1/2. Cells were serum starved for 24 h before fixation and stained with anti-YAP antibody. [A, B, and D–F] One of three independent results is presented. Bars, 25 µm.
of upstream Hippo pathway activator Nf2 (Neurofibromin 2, also known as Merlin; Fig. 1 A). Treatment of serum-starved, subconfluent MCF-10A with PI3K or PDK1 inhibitor caused the cytoplasmic localization of YAP (Fig. 1 A). This phenomenon was dependent on Lats kinases because Lats1/2 depletion blocked the effects of PI3K or PDK1 inhibitors on cytoplasmic localization of YAP (Fig. 1 A). This suggests the presence of some upstream PI3K regulators other than soluble mitogenic growth factors that negatively regulate the Hippo signaling pathway in subconfluent MCF-10A cells. To identify other potential regulators, we first tested whether inhibitors of other signaling molecules affect YAP localization in serum-starved, low cell density MCF-10A cells.

Src has been known to act as an upstream regulator of PI3K (Pleiman et al., 1994; Lu et al., 2003). Interestingly, the Src family kinase (SFK) inhibitor PP2 blocked the nuclear localization of YAP within 30 min as effectively as PI3K and PDK1 inhibitors (Fig. 1 B). Additional SFK inhibitors, including SKI-1, SU6656, and the dual Src/Ab1 kinase inhibitor dasatinib, were also tested. PP3, an analogue of PP2, and imaatinib, an Ab1 kinase inhibitor, were used as controls. Inhibition of endogenous SFK activity with these inhibitors for 30 min blocked YAP nuclear accumulation in serum-starved, low cell density MCF-10A cells (Fig. 1 B). Treatment with PP2 also increased the phosphorylation of YAP on its Lats phosphorylation site S127 as indicated by phospho-YAP (S127)–specific antibody and a decreased mobility on Phos-tag SDS-PAGE gel (Fig. 1 C; Kinoshita et al., 2006). Knockdown of Lats1/2 kinases abolished the effect of PP2 on the cytoplasmic localization of YAP (Fig. 1 D) in contrast to upstream Hippo pathway activator Nf2. These indicate that the effect of SFK activity in regulating YAP localization depends on Lats, the core kinase of the Hippo pathway.

The SFKs consist of nine family members, three of them—Src, Fyn, and Yes—being ubiquitously expressed (Thomas and Brugge, 1997). To identify which of these three ubiquitous SFK members is involved in the regulation of Hippo signaling, the expression of individual family members Src, Fyn, and Yes were knocked down by siRNA transfection (Fig. S1). Importantly, the knockdown of Src alone strongly prevented the nuclear localization of YAP in serum-starved, low cell density MCF-10A cells, whereas depletion of Yes had minimal effects (Fig. 1 E). The impact of Fyn depletion on the nuclear localization of YAP was less effective than Src knockdown, presumably because of its comparatively lower expression level in MCF-10A cells (Fig. S1 A). Similar to SFK inhibitor treatment, this phenomenon was Lats dependent because the knockdown of Lats1/2 prevented the effects of Src depletion on cytoplasmic retention of YAP (Fig. 1 F). This suggests that Src is the predominant SFK that inhibits the Hippo signaling pathway in MCF-10A cells.

**Src controls YAP activity**

For gain-of-function experiments, we generated inducible MCF-10A and HEK-293T cell lines expressing constitutively activated chicken-Src (Y527F, CA-Src)-GFP fusion protein in MCF-10A cells. Cells were treated with the indicated amount of doxycycline (Dox) for 12 h in complete medium and serum starved for an additional 24 h in the presence of doxycycline. Cells were lysed and subjected to Western blot analysis with the indicated antibodies. We performed two independent experiments. Tet, tetracycline (or doxycycline) inducible. (B) CA-Src expression and YAP localization. Doxycycline-inducible CA-Src-GFP–expressing MCF-10A cells were treated with 1 µg/ml doxycycline for 12 h and serum starved for 24 h in starvation medium containing doxycycline. Cells were fixed and immunofluorescence stained with anti-YAP antibody. Induction and membrane localization of CA-Src-GFP fusion protein were detected as green fluorescence. The results represent at least three independent experiments. (C) Reporter assay. Doxycycline-inducible HEK-293T cells expressing myr-GFP or CA-Src-GFP were transfected with HIP-flash or HOFlash reporters. Luciferase activity was measured in the absence or presence of doxycycline. Data were obtained from three independent experiments. Error bars represent standard deviation. (D) Biochemical effects of CSK depletion. MCF-10A cells transfected with control (Ctrl) or CSK siRNAs were serum starved for 24 h. Cell lysates were resolved in regular or Phos-tag SDS-PAGE gels and subjected to Western blotting with the indicated antibodies. Bar, 25 µm.

**Figure 2.** Activation of Src increases YAP nuclear activity. (A) Biochemical effects of Src expression. Doxycycline-inducible expression of constitutively activated chicken-Src (Y527F, CA-Src)-GFP fusion protein in MCF-10A cells. Cells were treated with the indicated amount of doxycycline (Dox) for 12 h in complete medium and serum starved for an additional 24 h in the presence of doxycycline. Cells were lysed and subjected to Western blot analysis with the indicated antibodies. We performed two independent experiments. Tet, tetracycline (or doxycycline) inducible. (B) CA-Src expression and YAP localization. Doxycycline-inducible CA-Src-GFP–expressing MCF-10A cells were treated with 1 µg/ml doxycycline for 12 h and serum starved for 24 h in starvation medium containing doxycycline. Cells were fixed and immunofluorescence stained with anti-YAP antibody. Induction and membrane localization of CA-Src-GFP fusion protein were detected as green fluorescence. The results represent at least three independent experiments. (C) Reporter assay. Doxycycline-inducible HEK-293T cells expressing myr-GFP or CA-Src-GFP were transfected with HIP-flash or HOFlash reporters. Luciferase activity was measured in the absence or presence of doxycycline. Data were obtained from three independent experiments. Error bars represent standard deviation. (D) Biochemical effects of CSK depletion. MCF-10A cells transfected with control (Ctrl) or CSK siRNAs were serum starved for 24 h. Cell lysates were resolved in regular or Phos-tag SDS-PAGE gels and subjected to Western blotting with the indicated antibodies. Bar, 25 µm.
the activation of Src detected by increased Y416 autophosphorylation. Depletion of CSK increased phosphorylation of C-terminal regulatory Y530 residue by C-terminal repressed through intramolecular interactions due to the phosphorylation status. In its inactive, "closed" state, Src kinase activity is tightly regulated by its conformational status. In its inactive, "closed" state, Src kinase activity is repressed through intramolecular interactions due to the phosphorylation of C-terminal regulatory Y530 residue by C-terminal Src kinase (CSK; Okada, 2012). Depletion of CSK increased the activation of Src detected by increased Y416 autophosphorylation (Fig. 2 D). CSK knockdown also increased the level of unphosphorylated YAP (Fig. 2 D). To inhibit endogenous SFK activity, we overexpressed CSK. Normally, recruitment of CSK to the plasma membrane is controlled by CSK-binding proteins such as PAG (also known as Cbp; Okada, 2012), but instead we targeted it to the membrane via a myristoyl group. Similar to treatment with SFK inhibitors (Fig. 1 B), induced expression of myristoylated CSK-GFP (myr-CSK-GFP) in serum-starved, low cell density MCF-10A cells increased the cytoplasmic localization of YAP (Fig. S3). The function of CSK depends on its kinase activity, as the expression of kinase-dead myr-CSK-K222R-GFP had no effect on the cellular localization and phosphorylation status of YAP in low cell density MCF-10A cells (Fig. S3). These data indicate that control of endogenous Src activity regulates YAP localization.

We wished to investigate the effect of Src on the Mst-dependent phosphorylation of Lats. Overexpression of Nf2 in HEK-293T cells has been shown to promote Lats1 phosphorylation by enhancing the complex formation of Lats1 with the Sav1–Mst1/2 complex on the plasma membrane (Yin et al., 2013). Transient transfection of HEK-293T cells confirmed that Lats1 hydrophobic motif (T1079) was phosphorylated by the expression of Sav1–Mst2 and was enhanced by Nf2 coexpression. Expression of CA-Src-GFP significantly inhibited the Mst2 phosphorylation on Lats1 even in the presence of Nf2 (Fig. 3 A). We also found that overexpression of Src-GFP or CA-Src-GFP decreased the binding between Sav1 and Lats1 (Fig. 3 B). These results indicate that the active Src negatively regulates the phosphorylation and activity of Lats1 as well as the Sav1–Lats1 interaction.

Regulation of Lats phosphorylation and activity depends on formation of a protein complex with Sav1 and other proteins. To determine whether Hippo pathway protein complexes are affected by Src activity, cytosolic fraction of serum-starved, low cell density MCF-10A cells that express TurboRFP or myr-CSK-GFP–expressing MCF-10A cells in low cell density (30%) were treated with 2 µg/ml doxycycline for 12 h in complete medium and starved for 24 h before harvest. Cytosolic proteins were fractionated by HPLC gel filtration chromatography on a Superose 12 column. Each fraction and corresponding input samples (Inp) were analyzed by SDS-PAGE and Western blotting with the indicated antibodies. IB, immunoblot; IP, immunoprecipitation; WCL, whole cell lysate.

Figure 3. Regulation of Lats1 and the Hippo complex by Src activity. (A) Inhibition of Mst2-dependent phosphorylation of Lats1 by active Src. HEK-293T cells expressing the indicated constructs were harvested at 24 h after transfection. Exogenous Flag-Lats1 protein was immunoprecipitated using anti-flag affinity gels and subjected to Western blotting with the indicated antibodies. Blots represent three independent results. (B) Src disrupts Sav1 binding to Lats1. Exogenous protein-expressing HEK-293T cells were harvested in NP-40 buffer. Sav1–Lats1 complex was coimmunoprecipitated using an anti-HA antibody. One of three independent results is presented. (C) Size-exclusion chromatography. Doxycycline-inducible TurboRFP- or myr-CSK-GFP–expressing MCF-10A cells were analyzed by SDS-PAGE and Western blotting with the indicated antibodies. IB, immunoblot; IP, immunoprecipitation; WCL, whole cell lysate.

a HIP/HOP-flash reporter assay, HIP (Hippo-YAP signaling incompetent promoter; negative control)-flash and HOP (Hippo-YAP signaling optimal promoter)-flash reporters (Fig. S2 A), and confirmed the specificity of reporter gene expression (Fig. S2, B–D). As shown in Fig. 2 C, induced expression of CA-Src-GFP, but not membrane-targeted GFP, selectively activated HOP-flash activity in HEK-293T cells. Collectively, these results show that catalytically active Src promotes the transcriptional activity of YAP through dephosphorylation of S127, its target site, and its nuclear accumulation.

SRC kinase activity is tightly regulated by its conformational status. In its inactive, “closed” state, SRC kinase activity is repressed through intramolecular interactions due to the phosphorylation of C-terminal regulatory Y530 residue by C-terminal SRC kinase (CSK; Okada, 2012). Depletion of CSK increased the activation of SRC detected by increased Y416 autophosphorylation (Fig. 2 D). CSK knockdown also increased the level of unphosphorylated YAP (Fig. 2 D). To inhibit endogenous SFK activity, we overexpressed CSK. Normally, recruitment of CSK to the plasma membrane is controlled by CSK-binding proteins such as PAG (also known as Cbp; Okada, 2012), but instead we targeted it to the membrane via a myristoyl group. Similar to treatment with SFK inhibitors (Fig. 1 B), induced expression of myristoylated CSK-GFP (myr-CSK-GFP) in serum-starved, low cell density MCF-10A cells increased the cytoplasmic localization of YAP (Fig. S3). The function of CSK depends on its kinase activity, as the expression of kinase-dead myr-CSK-K222R-GFP had no effect on the cellular localization and phosphorylation status of YAP in low cell density MCF-10A cells (Fig. S3). These data indicate that control of endogenous SRC activity regulates YAP localization.

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Provenzano and Keely, 2011). The phosphorylated Y397 residue creates a high affinity binding site for SH2-containing proteins, such as Src and PI3K, which activate a downstream signaling cascade (Cabodi et al., 2010). To test whether FAK is involved in regulating the Hippo signaling pathway, serum-starved, subconfluent MCF-10A cells were incubated for 30 min at the indicated concentrations of the two FAK inhibitors. Cell lysates were separated by SDS-PAGE or Phos-tag SDS-PAGE gels and immunoblotted with the indicated antibodies. γ-Tubulin was used as a loading control. We performed two independent experiments. (B) FAK inhibitors and YAP localization. MCF-10A cells transfected with control or Lats1/2 siRNA were serum starved and treated with DMSO, 5 µM PF-573228, or 10 µM PF-562271 for 30 min. Endogenous YAP was immunofluorescence stained with anti-YAP antibody. One of three independent results is presented. (C) Biochemical effects of FRNK expression. Doxycycline-inducible FRNK-GFP-expressing MCF-10A cells were treated with 1 µg/ml doxycycline (Dox) for 12 h and serum starved for 24 h in starvation medium containing doxycycline. Cells were detached, held in suspension for 30 min (Sus), and then replated on fibronectin (Fn)-coated coverslips in starvation medium for 2 h. The result shown is one of two independent results. (D) FRNK expression and YAP localization. MCF-10A cells were either uninduced or induced to express FRNK-GFP with 1 µg/ml doxycycline for 12 h and serum starved for 24 h in starvation medium containing doxycycline. Cells were dissociated and seeded on fibronectin-coated coverslips in starvation medium. Induction of FRNK-GFP and localization to focal adhesions were detected as green fluorescence. More than 150 cells from four random views were quantified, and data represent one of three independent results. Tet, tetracycline (or doxycycline) inducible. Bars, 25 µm.

FAK, inducible FRNK (FAK-related nonkinase)-expressing MCF-10A cells were generated. FRNK is an autonomously expressed splicing variant of FAK that contains the noncatalytic C-terminal domain of FAK (Richardson et al., 1997) and is known to act as a dominant-negative regulator of endogenous FAK and Pyk2 activity (Siegl et al., 1999; Govindarajan et al., 2000). Induced expression of FRNK blocked FAK activation in response to cell attachment to fibronectin (Fig. 4 C). Similar to results in the previous study (Zhao et al., 2012), suspension of MCF-10A cells phosphorylated YAP, whereas attachment to fibronectin-coated coverslips induced dephosphorylation of YAP (Fig. 4 C). Expression of FRNK inhibited dephosphorylation and nuclear localization of YAP in MCF-10A cells attached to fibronectin (Fig. 4, C and D). These results suggest that FAK plays an important role in the regulation of YAP phosphorylation and localization in a Lats-dependent manner.

To determine a functional order for FAK, Src, and PI3K–PDK1 in the regulation of the Hippo signaling pathway, inhib-
itors were added to the CA-Src–expressing MCF-10A cells (Fig. 5). Induced expression of CA-Src led to the nuclear localization of YAP (Fig. 2 B and Fig. 5). As before, treatment with Src, PI3K, or PDK1 inhibitors completely prevented the effect of CA-Src on nuclear YAP. The FAK inhibitor, however, did not block the effect of CA-Src even though it completely prevented the nuclear localization of YAP in MCF-10A cells without CA-Src expression (Fig. 5). These results imply that FAK acts as an upstream effector of Src in the regulation of YAP activity, and PI3K and PDK1 act as downstream effectors. Thus, the FAK–Src–PI3K–PDK1 pathway is responsible for YAP nuclear activity in serum-starved, subconfluent MCF-10A cells.

PI3K–PDK1 also mediate nuclear localization of YAP in confluent MCF-10A cells in response to mitogens such as EGF and LPA (Fan et al., 2013), and we therefore wished to test the roles of Src and FAK in their effects. The effect of mitogens on YAP localization can also be observed when added to serum-starved, subconfluent MCF-10A cells (Fig. 6 A). Pretreatment of cells with FAK inhibitor did not prevent YAP nuclear accumulation stimulated by EGF or LPA, suggesting that FAK is not required for receptor tyrosine kinase and G protein–coupled receptor regulation of the Hippo pathway (Fig. 6 A). Inhibition of Src kinase by siRNA-mediated depletion, myr-CSK-GFP expression, or dasatinib treatment prevented the EGF effect on nuclear localization of YAP (Fig. 6, A–C). This suggests that Src mediates EGF receptor (EGFR)–dependent regulation of the Hippo pathway, even though we previously reported that the Src inhibitor PP2 did not block the effect of EGF on YAP (Fan et al., 2013). In contrast, Src knockdown or induced expression of myr-CSK-GFP did not prevent the YAP nuclear accumulation induced by LPA signaling by G protein–coupled receptor (Fig. 6, A and B), indicating that Src does not act downstream of LPA. Because PI3K inhibitors block YAP nuclear accumulation caused by LPA treatment (Fan et al., 2013), LPA appears to activate YAP nuclear accumulation through PI3K in a Src-independent manner.

Adhesion to fibronectin activates YAP via FAK–Src–PI3K–PDK1

FAK-Src signaling is often stimulated by integrins in focal adhesions when cells spread on a substrate, as in this study. To directly test whether integrin activation of FAK controls the Hippo signaling pathway, we examined the subcellular localization of YAP in MCF-10A cells attached to fibronectin-, poly-d-lysine–, or laminin-coated surfaces. To minimize the actions of soluble factors that inactivate the Hippo pathway, it is critical to treat MCF-10A cells with serum starvation and amphiregulin-blocking antibody (Zhang et al., 2009; Fan et al., 2013) for 24 h before cells are detached to perform the adhesion assay. Suspended cells were allowed to adhere and spread on coverslips coated with fibronectin, poly-d-lysine, or laminin in starvation medium. Plating on a fibronectin-coated coverslip led to a significant increase of nuclear localization and dephosphorylation of YAP compared with plating on a poly-d-lysine– or laminin-coated coverslip (Fig. 7 A and see Fig. 8 B).

We next examined the effects of roles of signaling mediators on the subcellular localization of YAP in MCF-10A cells attached to fibronectin-coated coverslips. FAK, SFK, PI3K, and PDK1 inhibitors all decreased the nuclear localization of YAP stimulated by attachment to fibronectin-coated surfaces to levels similar to poly-d-lysine attachment (Fig. 7 B). In contrast, treatment with EGF or LPA increased the nuclear localization of YAP in MCF-10A cells attached to a poly-d-lysine–coated coverslip (Fig. 7 B). Notably, LPA treatment resulted in an additional increase of nuclear YAP in MCF-10A cells attached to a fibronectin-coated coverslip. To determine the role of SFK in the regulation of YAP cellular localization by ECM–cell binding, doxycycline-induced CA-Src-GFP–, myr-CSK-GFP–, or myr-GFP (control)–expressing MCF-10A cells were attached to fibronectin–, poly-d-lysine–, or laminin-coated coverslips. Induced expression of CA-Src-GFP led to the increase of nuclear localization and dephosphorylation of YAP in MCF-10A cells attached to poly-d-lysine– or laminin-coated coverslips (Fig. 8, A and B). In contrast, inhibition of endogenous SFK activity by myr-CSK-GFP prevented the nuclear localization and dephosphorylation of YAP in fibronectin-attached MCF-10A cells (Fig. 8, A and B). Collectively, these data suggest that fibronectin–integrin signaling controls the Hippo pathway through FAK–Src–PI3K–PDK1 (Fig. 9).

Discussion

In this study, we find that an integrin signaling pathway regulates the Hippo pathway and nuclear localization of YAP. Previous work demonstrated that PI3K–PDK1 signaling activity counters Hippo signaling and activates nuclear YAP by inhibiting Lats kinase activity in response to soluble mitogenic growth factors (Strallburger et al., 2012; Fan et al., 2013; Gumbiner and Kim, 2014). Our observation in the current study that PI3K and PDK1 inhibitors cause Lats-dependent YAP nuclear exclusion in highly spread subconfluent cells, even in the complete absence of soluble mitogens, suggested that PI3K–PDK1 might have similar roles in integrin adhesion signaling. This was confirmed...
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by the evidence for a role of FAK-Src signaling in Lats-dependent regulation of nuclear YAP, because FAK-Src activation is a major mechanism of integrin signaling, including activation of PI3K in response to adhesion to fibronectin (Tilghman and Parsons, 2008; Hynes, 2009; Schwartz, 2010). Moreover, adhesion to fibronectin was found to specifically enhance YAP nuclear localization dependent on the FAK–Src–PI3K–PDK1 signaling pathway. Thus, the well-known mitogenic integrin signaling pathway appears to operate in parallel with mitogenic growth factors in counteracting the growth inhibitory Hippo pathway.

Other studies have also provided evidence for a role of integrins in regulation of nuclear YAP. The expression of constitutively active β1-integrin mutants in membrane type 1 matrix metalloproteinase–null skeletal stem cells triggers FAK activation and increases nuclear localization of YAP/TAZ (Tang et al., 2008; Hynes, 2009; Schwartz, 2010). Moreover, adhesion to fibronectin was found to specifically enhance YAP nuclear localization dependent on the FAK–Src–PI3K–PDK1 signaling pathway. Thus, the well-known mitogenic integrin signaling pathway appears to operate in parallel with mitogenic growth factors to counteract the growth inhibitory Hippo pathway.

Figure 6. Roles of Src and FAK in the regulation of nuclear YAP by EGF or LPA. (A) Src depletion and FAK inhibition. MCF-10A cells transfected with control or Src siRNA were serum starved for 24 h and treated with EGF or LPA for 30 min. For FAK inhibitor treatment experiment, serum-starved, subconfluent MCF-10A cells were pretreated for 30 min with DMSO or 5 μM PF-573228, followed by a 30-min EGF or LPA treatment. YAP subcellular localization was determined by immunofluorescence staining. One of three independent experiments is presented. (B) CSK expression to inhibit Src activity. Paired with uninduced cells, doxycycline (Dox)-induced MCF-10A cells expressing myr-CSK-GFP fusion protein were serum starved for 24 h. Cells were treated with EGF or LPA for 30 min before fixation. Subcellular localization of YAP was determined by immunofluorescence staining. One representative out of three independent experiments is shown. Tet, tetracycline (or doxycycline) inducible. (C) Src inhibitors. Serum-starved, subconfluent MCF-10A cells were pretreated with PP3, PP2, imatinib, or dasatinib for 30 min. Cells were then treated with 20 ng/ml EGF for 30 min before fixation. Endogenous YAP was immunofluorescence stained. The result represents three independent experiments. Bars, 25 μm.

One earlier study did not detect a role for integrin-FAK signaling in the regulation of nuclear YAP in response to cell spreading, arguing instead for a distinct cell shape/actin/Rho signaling pathway (Zhao et al., 2012). In that study, FAK inhibitor treatment was not found to inhibit cell attachment–induced YAP dephosphorylation. Using their reported concentrations of the same FAK inhibitor (1 μM PF-562271), we observed a similar result regarding YAP phosphorylation status (Fig. 4 A, eighth lane). However, increased amounts of PF-562271 in doses that inhibit FAK autophosphorylation dramatically increase the phosphorylation of YAP (Fig. 4 A). Also, expression of dominant-negative FAK protein FRNK prevents fibronectin-induced dephosphorylation and nuclear accumulation of YAP, demonstrating that the Hippo pathway is controlled by the activation of FAK (Fig. 4, C and D). FAK inhibitors increased the cytoplasmic localization of YAP in a Lats-dependent manner (Fig. 4 B), indicating that they act through the Hippo pathway. In the earlier study (Zhao et al., 2012), detecting the effect of fibronectin adhesion on YAP may have been missed because it requires very careful control of cell-plating conditions to avoid residual mitogens, including blocking autocrine-secreted...
amphiregulin with neutralizing antibodies (Zhang et al., 2009; Fan et al., 2013). Under the conditions of our experiments, the effects of FAK inhibitors and cell adhesion were consistent, as were the roles of Src and PI3K signaling.

In the context of integrin-FAK signaling, we find that Src controls YAP activity in a Lats-dependent manner. Depletion of Lats1/2 kinases prevents YAP nuclear exclusion caused by SFK inhibitors (Fig. 1 D). Also, SFK inhibitor PP2 increases the phosphorylation of YAP on S127 residue (Fig. 1 C), a well-known Lats target phosphorylation site that induces 14-3-3 binding and cytoplasmic retention (Zhao et al., 2007), whereas the activation of Src, through overexpression of active Src or CSK depletion, decreases the phosphorylation of YAP (Fig. 2, A and D) and Mst-dependent phosphorylation of Lats (Fig. 3 A). In contrast, expression of membrane-targeted CSK increased the YAP nuclear exclusion (Fig. S3). Src appears to exert its effects on Lats and YAP via PI3K and PDK1 because PI3K or PDK1 inhibitor treatment prevents the effect of CA-Src on nuclear YAP. The stimulation of PDK1 by EGF and LPA signaling was previously found to dissociate the Hippo pathway complex and activate YAP (Fan et al., 2013). Our findings also indicate that Src stimulation of the PI3K-PDK1 pathway causes YAP nuclear accumulation because of its effects on the core Hippo pathway complex and Lats activity.

A recent study found that Src and Yes directly bind, phosphorylate, and regulate the activity of YAP rather than control the Hippo pathway during epithelial regeneration in inflammatory diseases (Taniguchi et al., 2015). However, we failed to detect the tyrosine phosphorylation of YAP by Src-GFP or CA-Src-GFP expression in HEK-293T cells but confirmed the reduction of YAP phosphorylation on S127 residue (Fig. S4 A). Interestingly, tyrosine residues of Mob1, Sav1, Nf2, and Lats1, but not Mst2, were phosphorylated by the expression of CA-Src-GFP in HEK-293T cells (Fig. S4, B and C). Although we cannot rule out the possibility that Src also interacts with other Hippo pathway components (Sudol, 1994; Zaidi et al., 2004; Rosenbluh et al., 2012; Enomoto and Igaki, 2013; Taniguchi et al., 2015), especially in other contexts, the sum of our findings demonstrates that Src affects YAP in MCF-10A cells through the Hippo complex and Lats activity.

MCF-10A cell attachment to fibronectin and collagen I (unpublished data), but not laminin or poly-d-lysine, induces nuclear localization of YAP (Fig. 7 A). In these experiments, substrate rigidity and mitogen starvation conditions are the
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same after attachment to different substrate-coated coverslips. Spreading on fibronectin, but not poly-d-lysine, is known to increase the phosphorylation of FAK and paxillin (Weiger et al., 2009). Laminin and fibronectin differentially activate Rho family GTPases through discrete signaling pathways involving the α3β1 and α5β1 integrins (Gu et al., 2001). Thus, our findings indicate that extracellular matrix composition delivers distinct signals to control the Hippo pathway.

The fibronectin–integrin signaling pathway identified in this study may contribute, at least in part, to the observed mechanotransduction mechanism that regulates the Hippo pathway and/or YAP nuclear activity. Cell shape and the actin cytoskeleton are critical regulators of YAP nuclear activity (Dupont et al., 2011; Sansores-Garcia et al., 2011; Wada et al., 2011; Aragona et al., 2013; Kim et al., 2013). Spread, flattened cells under tension have nuclear YAP, whereas round cells on soft substrates exhibit cytoplasmic YAP (Dupont et al., 2011; Wada et al., 2011), and loss of actin regulatory proteins leads to increased YAP activity (Sansores-Garcia et al., 2011; Aragona et al., 2013). Integrins are well-known mechanotransducers that link to the actin cytoskeleton and cause cell spreading and increased tension (Schwartz, 2010; Weber et al., 2011), and therefore it makes sense that FAK-Src–mediated integrin signaling could be involved. However, in some cases, the effect of cell shape and perturbation of the actin cytoskeleton affects YAP independent of Lats activity or the Hippo signaling cascade (Dupont et al., 2011; Aragona et al., 2013), so there may also be a shape-dependent signaling mechanism distinct from the pathway we describe. Rho signaling has been implicated in the actin/tension-dependent shape pathway (Takeichi, 2014), but the detailed mechanism of YAP regulation is not yet understood. Importantly, Lats-dependent mechanisms of cell shape/cytoskeletal regulation of nuclear YAP have also been reported (Sansores-Garcia et al., 2011; Wada et al., 2011; Halder et al., 2012; Zhao et al., 2012; Kim et al., 2013). We propose that the activation of PI3K–PDK1 by integrin-FAK-Src signaling is a component of the reported Lats-dependent cell shape/mechanotransduction perhaps in parallel with the reported Lats-independent regulation of YAP by cell shape. It is worth noting that PI3K signaling is also known to regulate Rho/Rac signaling (Welch et al., 2003; Di Paolo and De Camilli, 2006), so it is possible that there is overlap between the two mechanisms.

In summary, we demonstrate that the adhesion of cells to fibronectin functions as a negative upstream regulator of the Hippo pathway via the FAK–Src–PI3K–PDK1 pathway. Our results provide an explanation for the effects of the fibronectin–integrin interaction on YAP nuclear localization and can account, at least in part, for the Lats-dependent components of mechanotransduction.

Materials and methods

Cell culture
MCF-10A human mammary epithelial cells (a gift from Joan S. Brugge, Harvard Medical School, Boston, MA) were cultured in DMEM/F12 medium supplemented with 5% of horse serum, 0.5 µg/ml hydrocortisone, 100 ng/ml cholera toxin, 10 µg/ml insulin, and
arginine using the QuikChange II Site-Directed Mutagenesis kit (Agc-src (amino acids 1–16). Lysine 222 residue of CSK was mutated to primers containing a myristoylation signal peptide sequence of chicken c-src under a CMV promoter), were gifts from Marius Sudol (Weis Center for Reagents

siRNA oligonucleotides specific to human Src (GAGAACCCUG-GUGUCGAAAG), Fyn (GCUCUGAAAUUACCAAAUC), Yes (GAAGGACCCUGAAGAAAGA), CSK (L-003110-00-0005), N12 (L-003917-00-0005), Lats1 (L-004632-00-0005), Lats2 (L-003865-00-0005; GE Healthcare), and RNAiMax (Life Technologies) were used to knock down the expression of genes that we investigated. On-target plus nontargeting pool (D-001810-10-05; GE Healthcare) was used as a control siRNA. Antibodies used for immunofluorescence staining or Western blot include YAP (1:100 dilution for immunofluorescence staining, 1:1,000 dilution for Western blotting; mouse; no. 63.7), CSK (1:1,000 dilution; rabbit; no. C-20), Sav1 (1:1,000 dilution; mouse; no. J1-6), FAK (1:1,000 dilution; rabbit; no. C-20), GFP (1:2,000 dilution; mouse; no. B-2), c-Myc (1:1,000 dilution; mouse; no. 9E10; Santa Cruz Biotechnology, Inc.), Src (1:1,000 dilution; mouse; no. 2110), phospho-PI3K (1:1,000 dilution; rabbit; no. 4228), AKT (1:1,000 dilution; mouse; no. 2920), pAkt-T580 (1:1,000 dilution; rabbit; no. 4056), pAkt-S473 (1:1,000 dilution; rabbit; no. 3787), pSrc-Y416 (1:1,000 dilution; rabbit; no. 2113), pYAP-S127 (1:1,000 dilution; rabbit; no. 4911), pLats1-T1079 (1:1,000 dilution; rabbit; no. 8654), pLats1-S909 (1:1,000 dilution; rabbit; no. 9137), PDK1 (1:1,000; rabbit; no. 3062), Fyn (1:1,000; rabbit; no. 4023; Cell Signaling Technology). pFAK-Y397 (1:1,000 dilution; rabbit; no. F7926), Flag (1:1,000 dilution; mouse; no. F1804), β-tubulin (1:3,000 dilution; rabbit; no. 2920), pAkt-T308 (1:1,000 dilution; rabbit; no. 4043), and phospho-tyrosine (1:1,000 dilution; mouse; no. 05–321; EMD Millipore). The following inhibitors were used: SFK inhibitors SKI-1 and SU6654, Src/Abi inhibitor dasatinib monohydrate, Abl inhibitor imatinib meylate, PDK1 inhibitor BX-795, FAK inhibitors PF-573228 and PF-562271 (Santa Cruz Biotechnology, Inc.), SFK inhibitor PP2 and its inactive analogue PP3 (EMD Millipore), and PTK inhibitor wortmannin (Sigma-Aldrich). Unless indicated otherwise, all inhibitors were used at a concentration of 10 μM, except for PF-573228 and BX-795 (5 μM). Flag-YAP and Flag-Lats1, under a cytomegalovirus (CMV) promoter, were gifts from Marius Sudol (Weis Center for Research, Danville, PA). HA-mob, HA-Nf2, HA-Sav1, and Myc-Mst2 in the pcDNA3 backbone were gifts from Kun-Liang Guan (University of Edinburgh, Edinburgh, Scotland, UK).

Construction of plasmids

To generate doxycycline-inducible lentiviral vector, coding sequence of TurboRFP in TRIPZ (GE Healthcare) was replaced with GFP, and the following genes were cloned to generate GFP-fusion proteins. The constitutively active Src was from CA-Src-GFP vector. Myr-CSK was amplified from the human Orfeome library (Lamesch et al., 2007) with primers containing a myristoylation signal peptide sequence of chicken c-src (amino acids 1–16). Lysine 222 residue of CSK was mutated to arginine using the QuikChange II Site-Directed Mutagenesis kit (Agilent Technologies) to create the kinase-dead form of CSK (K222R). FRNK was amplified (amino acids 693–1,053) from the chicken FAK expression vector under a CMV promoter (a gift from James Casanova, University of Virginia, Charlottesville, VA). Induced expression of target genes was validated in HEK-293T cells by transient transfection and doxycycline treatment.

Construction of reporters

To measure the transcriptional activity of TEAD-YAP, we developed a HIP/HOP-flash luciferase reporter system. HIP-flash contains multiple copies of wild-type TEAD-binding sites with minimal promoter and a luciferase reporter gene. HIP-flash was generated as a negative control for HOP-flash activity. TEAD-binding sites in the promoter of CTGF (Zhou et al., 2008) were multimerized (Nakajima and Yaoita, 1997) using primers 5′-GGTGGGGGAATAAGCGGATGCTCCGCCGGGGAGGAA-3′ and 5′-ACAAGCTGAAAAATTCTGACATCCCTCCCACCCACCAACACGG-3′ and then were cloned into pCR-bluntII-Topo vector (Life Technologies). Mutated TEAD-binding sites were multimerized using primers 5′-GGTGGGGGAAGATCAGGAGGAGATCCTGCTGTTGTTGGGGGA-3′ and 5′-ACAAGCTGAAAAATTCTGACATCCCTCCCACCCACCAACACGG-3′. Wild-type and mutated TEAD-binding sites are underlined, and mutated sequences are marked in lowercase letters. DNA fragments with eight TEAD-binding sites or seven mutant TEAD-binding sites were ligated into pGL3-basic vector (Promega) together with minimal promoter of TOP-flash reporter (Korinek et al., 1997), creating HOP-flash and HIP-flash reporters, respectively.

Generation of doxycycline-inducible cell lines

Lentiviral doxycycline-inducible vectors were cotransfected with envelope vector pMD2.G (Addgene plasmid 12259) and packaging vector psPAX2 (Addgene plasmid 12260) into HEK-293T/17 cells using Lipofectamine 2000 (Life Technologies). After 18 h of transfection, the medium was replaced with UltraCULTURE medium (Lonza) supplemented with L-glutamine and antibiotics. Starting the next day, the lentivirus-containing supernatants were collected every 12 h for 3 days and stored at 4°C. After spin-down at 2,000 rpm for 5 min to remove cell debris, supernatants were filtered using a 0.45-μm pore size filter. About 30 ml of filtered supernatants were then transferred into ultracentrifuge tubes containing 5 ml of 20% sucrose cushion. Lentiviral particles were concentrated by centrifugation at 100,000 g for 2 h at 4°C and finally resuspended in 1× HBSS buffer. MCF-10A cells were incubated overnight with lentiviral particles and 6 μg/ml polybrene (EMD Millipore) and replaced with fresh medium. Antibiotics selection was started 3 d after transduction with 2 μg/ml puromycin and maintained on selection medium for an additional 7 d. After selection, cells were treated with 2 μg/ml doxycycline for 24 h, and MCF-10A cells with high levels of GFP were collected by FACS cell sorting. Cells were then maintained in the absence of doxycycline.

Mitogen and inhibitor treatment

For the inhibitor treatment experiments, 6 × 104 MCF-10A cells were seeded into 1 well of 24-well plates (~10% cell density). After 24 h, cells were serum starved in DMEM/F12 medium supplemented with 1 μg/ml amphiregulin-blocking antibody (AF-262; R&D Systems) for 12–24 h. The blocking antibody to amphiregulin, an EGFR ligand secreted by MCF-10A cells (Zhang et al., 2009), was added to the starvation medium to prevent non–cell-autonomous activation of EGFR signaling. After 3 h before the treatment with mitogens or inhibitors, the culture medium was replaced with fresh starvation medium to remove secreted soluble factors from cells. Inhibitors were added for 30 min. For the cotreatment experiments, cells were pretreated with the indicated inhibitors for 30 min and then treated with 20 ng/ml EGF (PeproTech) or 25 μM LPA (Sigma-Aldrich). Doxycycline-inducible
MCF-10A cell lines were induced by 1 μg/ml doxycycline for 12 h in complete medium and starved for another 24 h with 1 μg/ml amphiregulin-blocking antibody and 1 μg/ml doxycycline. We performed at least two independent experiments.

**Immunoprecipitation and immunoblotting**

For immunoprecipitation, HEK-293T cells were washed with ice-cold HBSS and lysed with radioimmunoprecipitation assay buffer (50 mM Tris, pH 7.4, 150 mM NaCl, 1% NP-40, 0.1% SDS, 0.5% sodium deoxycholate, 2 mM EDTA, 1 mM Na3VO4, 10 mM NaF, and protease inhibitors) or NP-40 buffer (150 mM NaCl, 1.0% NP-40, 50 mM Tris, pH 8.0, 1 mM Na3VO4, 10 mM NaF, and protease inhibitors) at 24 h after transfection. Cell lysates were incubated with the indicated immunoprecipitation antibodies and Sepharose 4 Fast Flow Protein A/G beads (GE Healthcare) or anti-flag M2 affinity gels (Sigma-Aldrich). For regular Western blotting, cells were washed with ice-cold HBSS and directly lysed with 2x SDS buffer (0.12 M Tris, pH 6.8, 4% SDS, 20% glycerol, and 10% 2-mercaptoethanol) after brief sonication to reduce viscosity. For regular SDS-PAGE, the lysates were separated by 5–17% gradient SDS-PAGE gel and transferred to polyvinylidene difluoride membrane (EMD Millipore) using a semidy transfer apparatus (Bio-Rad Laboratories). Phos-tag SDS-PAGE gel consisted of a stacking gel (4.5% wt/vol acrylamide, 125 mM Tris, pH 6.8, and 0.1% SDS) and a separating gel (7% wt/vol acrylamide, 380 mM Tris, pH 8.8, 15 µM Phos-tag acrylamide, 30 µM MnCl2, and 0.1% SDS). A 37.5:1 wt/vol mixture of acrylamide/bis-acrylamide (Bio-Rad Laboratories) was used as an acrylamide solution. Gel electrophoresis was performed at 10 mA with SDS running buffer (25 mM Tris, 192 mM glycine, and 0.1% wt/vol SDS) until the dye reached the bottom. After electrophoresis, acrylamide gel was equilibrated two times with transfer buffer (25 mM Tris, 192 mM glycine, 0.1% wt/vol SDS, and 10% methanol) containing 10 mM EDTA for 10 min, followed by washing two times with transfer buffer without EDTA. Wet transfer was performed to transfer proteins onto polyvinylidene difluoride membrane (EMD Millipore). The blot was sequentially incubated with primary antibody and HRP-conjugated secondary antibody. The HRP reaction was performed with chemiluminescent HRP substrate (EMD Millipore) and exposed to x-ray film (Genesee Scientific) or detected using an imaging system (LAS-3000; Fujifilm). Proteins were also detected with an infrared imaging system (Odyssey; LI-COR Biosciences) using IRDye-labeled secondary antibody (LI-COR Biosciences). Western blot images were assembled for illustration using Photoshop (Adobe). Band intensity was quantified using ImageJ (National Institutes of Health).

**HPLC gel filtration chromatography**

Doxycycline-inducible TurboRFP or myr-CSK-GFP–expressing MCF-10A cells were seeded in two 150-mm-diameter plastic culture dishes at low cell density (∼30%) and treated with 2 µg/ml doxycycline for 12 h in complete medium. Cells were then starved for 24 h in DMEM/F12 medium containing 2 µg/ml doxycycline. Cells were washed with ice-cold HBSS and resuspended in S100/P100 buffer (10 mM Hepes, pH 7.4, 1 mM EDTA, and 150 mM NaCl) supplemented with protease and phosphatase inhibitors. Cells were passed repeatedly through a 27-gauge needle and then subjected to low speed centrifugation at 21,130 g for 5 min. Supernatants were further ultracentrifuged at 100,000 g for 1 h. The supernatant’s pellets were collected and concentrated using Microcon centrifugal filters (EMD Millipore). The concentrated supernatants were fractionated by gel filtration HPLC on a Superose 12 column (GE Healthcare) as follows: 0.15 ml of the cytosolic fraction was loaded onto the column at a flow rate of 0.5 ml/min, the absorbance at 280 nm was monitored, and 0.5-ml fractions were collected. The proteins were incubated with 2 ml of chilled acetone at −20°C for 2 h and precipitated by centrifugation at 12,000 g for 5 min. The proteins were resuspended in SDS-DTT-urea buffer (6 M urea, 1% wt/vol DTT, 62.5 mM Tris-HCl, pH 6.8, 2% SDS, 10% glycerol, and 0.01% bromophenol blue) and subjected to Western blotting.

**Immunofluorescence staining and luciferase assay**

MCF-10A cells seeded in a 24-well plate or in fibronectin-, poly-d-lysine-, or laminin-coated coverslips were used in immunofluorescence staining. Cells were fixed with 4% paraformaldehyde in PBS for 15 min at room temperature (or overnight at 4°C) and washed three times for 5 min each in 100 mM glycine containing PBS, followed by permeabilization with 0.1% Triton X-100 in PBS for 10 min. After blocking with 3% nonfat dry milk in PBS for 1 h, cells were incubated with primary antibody diluted in 1% BSA/PBS overnight at 4°C. After washing with PBS, cells were incubated with Alexa Fluor 488– or 594–conjugated secondary antibodies (Life Technologies) for 1 h and washed with PBS. Cell nuclei were counterstained and mounted with a mounting medium with DAPI (Vectashield; Vector Laboratories). Immunofluorescence staining images were collected at room temperature using a 20x objective (LUCPlanFLN; NA 0.45; Olympus) on an inverted microscope (IX71; Olympus) coupled to a camera (ORCA-R2 C10600-10B; Hamamatsu Photonics). Images were acquired, normalized, analyzed by SlideBook 5.0 (Intelligent Imaging Innovations), and assembled for illustration using Photoshop (Adobe). For luciferase reporter assay, myr-GFP– or CA-Src-GFP–inducible HEK-293T cells were seeded in 24-well plates the day before transfection. HIP-flash or HOP-flash reporter and indicated plasmids were cotransfected with Lipofectamine 2000 (Life Technologies). 24 h after transfection, myr-GFP– or CA-Src-GFP–inducible HEK-293T cells were incubated in 0.1% FBS containing DMEM/F12 medium with or without 1 µg/ml doxycycline for 16 h. Luciferase activity was measured with a luciferase assay system (Promega).

**Online supplemental material**

Fig. S1 shows the validation of siRNA depletion of SFKs and the activation of downstream PI3K signaling by Src expression. Fig. S2 shows a diagram of HIP/HOP-flash reporters and the validation of the specificity of reporter gene expression in response to YAP/TAZ activity. Fig. S3 shows the cytoplasmic localization of YAP by the induced expression of myr-CSK. Fig. S4 shows the tyrosine phosphorylation of Hippo pathway components by the expression of Src kinase. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201501025/DC1.

**Attachment of cells to coated coverslips**

Coverslips were acid-washed and coated with 20 µg/ml fibronectin (Sigma-Aldrich), 50 µg/ml poly-d-lysine, or 20 µg/ml laminin. MCF-10A cells were starved for 24 h with DMEM/F12 medium containing 1 µg/ml amphiregulin-blocking antibody. Cells were dissociated with 2 ml Accutase (Life Technologies) and centrifuged at 1,000 rpm for 5 min. Cells were then washed two times with DMEM/F12 medium to remove residual Accutase. 6 x 104 MCF-10A cells were held in suspension by rotating on a tube rotator at 37°C for 30 min. Cells were then allowed to attach to coated coverslips in starvation medium and incubated at 37°C for 2–3 h. For Western blotting, the suspended cells were seeded on 25-mm-diameter fibronectin-, poly-d-lysine–, or laminin-coated coverslips in starvation medium for 2 h and directly lysed with 2x SDS buffer.
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