Phosphorylation by NLK inhibits YAP-14-3-3-interactions and induces its nuclear localization

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Abstract

Hippo signaling controls organ size by regulating cell proliferation and apoptosis. Yes-associated protein (YAP) is a key downstream effector of Hippo signaling, and LATS-mediated phosphorylation of YAP at Ser127 inhibits its nuclear localization and transcriptional activity. Here, we report that Nemo-like kinase (NLK) phosphorylates YAP at Ser128 both in vitro and in vivo, which blocks interaction with 14-3-3 and enhances its nuclear localization. Depletion of NLK increases YAP phosphorylation at Ser127 and reduces YAP-mediated reporter activity. These results suggest that YAP phosphorylation at Ser127 and at Ser128 may be mutually exclusive. We also find that with the increase in cell density, nuclear localization and the level of NLK are reduced, resulting in reduction in YAP phosphorylation at Ser128. Furthermore, knockdown of Nemo (the Drosophila NLK) in fruit fly wing imaginal discs results in reduced expression of the Yorkie (the Drosophila YAP) target genes expanded and DIAP1, while Nemo overexpression reciprocally increased the expression. Overall, our data suggest that NLK/Nemo acts as an endogenous regulator of Hippo signaling by controlling nuclear localization and activity of YAP/Yorkie.

Keywords 14-3-3; cell density; Hippo signaling; NLK; YAP

14-3-3 family and participates in a variety of biological responses, tissue homeostasis have become a long-standing topic of interest. As loss of the organ size control is linked to many human diseases, including cancer and degenerative diseases, regulation of the organ size could be an attractive therapeutic strategy. Recently, Hippo signaling has been identified as a major signaling pathway to control the organ size; dysregulation of this pathway results in aberrant growth [1].

The Hippo pathway is evolutionarily conserved from nematodes to humans and controls a variety of cellular processes, such as cell proliferation and apoptosis, culminating in the organ size regulation [2]. This signaling pathway was originally discovered via genetic screening in Drosophila to identify genes involved in the regulation of the organ size. Hippo, Warts, Salvador, and Yorkie are core components of the pathway which, when mutated, result in dramatic overgrowth of mutant tissues [3,7]. In addition, a number of novel regulatory components in the Hippo pathway have emerged [3,8,9]. In mammals, the Hippo pathway consists of the serine/threonine kinases MST1/2 (mammalian Ste20-like kinase, Hippo orthologs) and LATS1/2 (large tumor suppressor kinase, Warts orthologs), the scaffolding protein WW45 (Salvador ortholog), a transcriptional co-activator YAP (Yes-associated protein, Yorkie ortholog) and its paralogue TAZ (known as WWTR1) [10]. Activation of the Hippo signaling by extracellular stimuli, mechanical sensation, or cell density promotes cytoplasmic retention of YAP/TAZ via interaction with 14-3-3, followed by β-TrCP-mediated proteasomal degradation of YAP/TAZ [11–13]. These processes depend on LATS1/2-mediated direct phosphorylation at different sites of YAP [11,14]. YAP phosphorylation on S127 is essential to associate with 14-3-3, while phosphorylation on S397 by LATS1/2 creates a phospho-degron motif for β-TrCP binding [12]. Conversely, dephosphorylated YAP/TAZ functions through the TEAD family transcriptional factors in the nucleus to activate genes involved in cell proliferation and tissue overgrowth [15]. Therefore, YAP/TAZ phosphorylation by LATS1/2 has been accepted as a critical process that regulates YAP/TAZ transcriptional activity.

Nemo-like kinase (NLK) belongs to the atypical MAP-kinase family and participates in a variety of biological responses,
including cell motility and embryogenesis [16,17]. NLK is thought to act as a proline (P)-directed serine/threonine kinase [16]. As several transcriptional regulators have been identified as NLK substrates, NLK is known to play crucial regulatory roles in diverse signaling pathways, including Wnt/β-catenin, Notch, FOXO, and mTOR signaling [18–21]. However, the functional interaction between NLK and Hippo-YAP signaling has been unknown.

Here, we show that NLK phosphorylates YAP at Ser128, which leads to reduction in LATS-mediated phosphorylation at Ser127. NLK thus promotes YAP nuclear localization and transcriptional activity via dissociation from 14-3-3. The level of YAP phosphorylation at Ser128, as well as the nuclear localization and overall NLK levels, becomes decreased upon augmentation of the cell density. Consistently, we further show that knockdown of NLK in Drosophila reduces expression of YAP target genes in imaginal discs. Overall, our data suggest that NLK-mediated YAP phosphorylation is a bona fide mechanism for regulation of YAP activity in vivo.

Results and Discussion

NLK interacts with and phosphorylates YAP

Previous reports have shown that NLK is involved in a variety of signaling pathways, including Wnt, Notch, Foxo, and mTOR signaling. This prompted us to examine whether NLK is able to regulate Hippo signaling. To this end, we overexpressed core components of the Hippo pathway with or without NLK and examined mobility shifts, which might be induced by NLK-mediated phosphorylation. As shown in Fig EV1A, NLK led to a strong mobility shift of YAP but not other components such as MST1, WW45, and TEAD1. Meanwhile, overexpression of two forms of kinase-negative NLK constructs (K155M, later on referred as “KM”, and T286V, later on referred as “TV”) [16] did not induce a mobility shift for YAP, indicating the kinase activity of NLK was required for the mobility shift of YAP (Figs 1A and EV1B). Elimination of the mobility shift by treatment with λ-phosphatase as well as an in vitro kinase assay further confirmed that YAP was phosphorylated directly by NLK (Figs EV1C and 1B). In line with these findings, we observed that both endogenous and exogenous NLK interacted with YAP, and the two kinase-negative forms of NLK interacted with YAP much more strongly than the wild type did (Figs 1C and EV1D). Taken together, these results suggest that NLK interacts with and directly phosphorylates YAP, and this phosphorylation in turn may regulate binding of NLK and its substrate.

YAP phosphorylation at Ser128 by NLK attenuates phosphorylation at Ser127

YAP is evolutionarily conserved from nematodes to flies and to mammals, and phosphorylation of YAP at Ser127 (corresponding to human YAP) by serine/threonine kinase LATS1/2 is an important

Figure 1. NLK phosphorylates YAP.

A NLK but not kinase-negative NLK (Flag-NLK-KM, K155M) caused mobility shift of EGFP-YAP. EGFP-YAP was transfected with empty vector, Flag-NLK-WT, or Flag-NLK-KM into HEK293T cells. Mobility shift was shown by immunoblotting.

B NLK phosphorylates YAP in vitro. Flag-NLK-WT or Flag-NLK-KM was immunoprecipitated with anti-Flag antibody from cell lysates of HEK293T cells transfected with each plasmid and subjected to in vitro kinase assay with bacterially purified GST-YAP in the presence of [32P-ATP]. Phosphorylation of GST-YAP and autophosphorylation of NLK were examined by autoradiography (top panel), and Coomassie blue staining showed similar levels of GST-YAP and Flag-NLK (bottom panel).

C NLK interacts with YAP at the endogenous level. Immunoprecipitates from HEK293T cell lysates with indicated antibody were blotted with anti-YAP or anti-NLK antibody.
step for regulation of nuclear/cytoplasmic localization and stability (Fig 2A). Interestingly, YAP contains a highly conserved SP motif, a putative NLK phosphorylation site [22], immediately adjacent to Ser127 (Fig 2A). To examine whether NLK is involved in YAP phosphorylation at Ser128, HA-YAP was transiently transfected into HEK293T cells along with either Flag-NLK-WT or Flag-NLK-KM and then HA-YAP was immunoprecipitated with the anti-HA antibody. LC–MS/MS analysis revealed phospho-peaks of YAP at Ser128 when Flag-NLK-WT was transfected and phospho-peaks of YAP at Ser127 when Flag-NLK-KM was transfected (Fig EV2). This finding prompted us to explore whether or not NLK affects YAP phosphorylation at Ser127. As shown in Fig 2B, endogenous phosphorylation of YAP at Ser127 was reduced by NLK-WT but not by the NLK-KM. Interestingly, YAP phosphorylation at Ser397, another LATS1/2 consensus site in YAP that creates a phospho-degron motif for β-TrCP binding, was also reduced when NLK was overexpressed (Fig 2B). Next, we used lithium chloride (LiCl), previously used as an inhibitor of NLK and GSK3β activities [23,24], to further test whether NLK has a negative effect on YAP phosphorylation at Ser127. LiCl augmented YAP phosphorylation at Ser127 when compared to KCl (Fig EV3A). On the other hand, both the GSK3β-specific inhibitor BIO (Fig EV3A) and knockdown of GSK3β had no effect on YAP phosphorylation at Ser127 (Fig EV3B). These data suggest that LiCl-mediated induction of YAP phosphorylation at Ser127 was due to inhibition of NLK. To further show that NLK regulates YAP phosphorylation at Ser127 and Ser397, we used the CRSPR/Cas9 system to assess loss-of-function effects of NLK on YAP phosphorylation. Transient transfection of two different gRNA

![Figure 2: YAP phosphorylation at Ser128 by NLK attenuates phosphorylation at Ser127.](image-url)

A Conservation of phosphorylation sites for LATS1/2 and NLK. Amino acid sequences of different species of YAP, Yorkie (in Drosophila), and TAZ (YAP paralog) were aligned. NLK is a MAPK (mitogen-activated protein kinase)-type kinase, which prefers proline-directed motif (S/TP).

B NLK-WT but not NLK-KM reduces Ser127 phosphorylation of endogenous YAP. Cell lysates from HEK293 cells transfected with empty vector (−), Flag-NLK-WT, or Flag-NLK-KM were immunoblotted with antibodies shown in the figure (left panel). The ratio of p-YAP(S127) or p-YAP(S397)/total-YAP of three independent Western blots was quantified (right panel). Data are presented as mean ± SEM. **P < 0.01. Student’s t-test was used for statistical analysis.

C Deficiency of NLK leads to increased YAP phosphorylation at Ser127. Lysates from wild-type and two independent NLK knockout HEK293 cell lines were immunoblotted with antibodies indicated in the figure (left panel). The ratio of p-YAP(S127) or p-YAP(S397)/total-YAP of three independent Western blots was quantified (right panel). Data are presented as mean ± SEM. *P < 0.05 and **P < 0.01. Student’s t-test was used for statistical analysis.

D Ectopic expression of NLK attenuates YAP phosphorylation at Ser127 while increases phosphorylation at Ser128. Cell lysates from HEK293 cells transfected with empty vector (−), Flag-NLK-WT, or Flag-NLK-KM were immunoblotted with antibodies shown in the figure.

E Substitution of Ser127 with alanine enhances phosphorylation at Ser128, and vice versa. Cell lysates from HEK293T cells transfected with empty vector (−), EGFP-YAP, EGFP-YAP-S127A, or EGFP-YAP-S128A were immunoblotted with antibodies shown in the figure.

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sequences targeting NLK with Cas9 resulted in an efficient ablation of NLK (Fig 2C). Consistent with above results, the levels of phosphorylation at Ser127/397 were significantly increased in the NLK KO cell pool (Fig 2C).

To further confirm phosphorylation of YAP at Ser128 by NLK, we generated antibody against the phosphorylated YAP-S128 peptide and confirmed its specificity as determined by no positive signal detected upon expression of YAP-S128A unlike wild-type YAP (Fig EV3C). Marked increase in endogenous YAP phosphorylation at Ser128 by NLK-WT, but not NLK-KM, was observed (Fig EV3D). Also, increased YAP phosphorylation at Ser128 by ectopic expression of NLK was significantly blocked by the treatment of LiCl but not KCl (Fig EV3E). To further test whether NLK phosphorylates YAP-S128, in vitro kinase assay was performed. Phosphorylation at S128 of GST-YAP, but not GST-YAP-S128A, was induced by immunoprecipitated Flag-NLK, but not NLK-KM (Fig EV3F). Overall, these data suggest that our antibody is specific to pYAP-Ser128 that is phosphorylated by NLK.

Interestingly, increased phosphorylation of Ser128 by NLK showed a reverse correlation with the phosphorylation at Ser127 (Fig 2D). Phosphorylation at Ser128 was elevated in the YAP-S127A mutant, whereas the YAP-S128A mutant exhibited increased phosphorylation at Ser127 when compared to wild-type YAP (Fig 2E). To further test whether the phosphorylation of S128 by NLK inhibits phosphorylation of YAP-S127 by LATS1, in vitro kinase assay was performed. As shown in Fig EV3G, the phosphorylation at Ser127 of GST-YAP and GST-YAP-S128A by immunoprecipitated Flag-LATS1 was highly efficient, while phosphorylation at Ser127 of GST-YAP-S128D was strongly suppressed (Fig EV3G). Together with LC–MS analysis, which did not reveal a concomitant YAP phosphorylation on S127 and S128 (Fig EV2), these results suggest that phosphorylations of Ser127 and Ser128 of YAP may be mutually exclusive.

**NLK promotes nuclear localization of YAP by inhibiting its interaction with 14-3-3**

It has been well known that phosphorylation of YAP at Ser127 by LATS1/2 induces cytoplasmic retention of YAP through its interaction with 14-3-3 [11]. We showed that NLK increased YAP phosphorylation at Ser128 along with reduced phosphorylation at Ser127 (Fig 2D). Thus, we examined whether or not YAP phosphorylation at Ser128 by NLK affects the interaction between YAP and 14-3-3. Overexpression of NLK-WT, but not NLK-KM, completely blocked the interaction between YAP and 14-3-3 (Fig 3A). Consistently, NLK KO cells displayed stronger interaction between YAP and 14-3-3 compared with normal cells (Fig 3B). To further confirm this, we utilized the YAP mutants S127A, S128A, and S128D (a phosphorylation mimetic form) (Fig 3C). Consistent with the above findings (Figs 2E and 3A and B), YAP-S128A showed a much stronger interaction with 14-3-3 than YAP-WT did, whereas neither YAP-S127A nor YAP-S128D showed binding to 14-3-3 (Fig 3C).

Since interaction with 14-3-3 leads to cytoplasmic sequestration of YAP, we next explored whether or not NLK induces nuclear localization of YAP. Overexpression of NLK-WT, but not NLK-KM, strongly induced nuclear localization of EGFP-YAP (Fig EV4A). Consistently, the results from immunofluorescence and subcellular fractionation analysis clearly showed that YAP-S128A mutant localized to the cytoplasm while the phospho-mimetic YAP-S128D mutant was mainly present in the nucleus, which was similar to YAP-S127A (Figs 3D and EV4B and C). Also, normal cells exhibited strong nuclear localization of YAP at low density, but the NLK KO cell pool exhibited reduced nuclear YAP level (Fig EV4D) along with enhanced cytoplasmic localization of YAP (Fig 3E). Overall, these results indicate that NLK-mediated phosphorylation at Ser128 leads to nuclear localization of YAP by disrupting the interaction between YAP and 14-3-3.

**NLK localization and protein level are regulated in a cell density-dependent manner**

Recent studies have suggested that NLK is present in the nucleus or cytoplasm, and dimerization of NLK induced by growth factors is essential for its nuclear localization and functional activation [25]. Because NLK regulates YAP localization (Figs 3 and EV4) and nuclear localization of YAP is influenced by cell confluence [11], we investigated whether or not nuclear localization of NLK is changed by cell density. NLK mainly localized to nuclei of cells at low density, whereas it was redistributed in the cytoplasm at high cell density of MCF10A and NIH3T3 cells (Fig 4A). Interestingly, the protein level of NLK, but not its mRNA expression, decreased along with increasing cell density (Fig 4B and C). Consistently, the phosphorylation of endogenous YAP at Ser128 was severely reduced in cells at high cell density, concomitant with increased YAP phosphorylation at Ser127 and Ser397 (Fig 4B). These data indicate that localization as well as protein level of NLK is controlled in a cell density-dependent manner, which affects the status of phosphorylation and cytoplasmic localization of YAP.

**NLK regulates YAP transcriptional activity and cell migration**

YAP interacts with TEAD and induces expression of genes involved in the regulation of cell proliferation, anti-apoptosis, and cell migration [15]. Therefore, we next investigated whether or not NLK controls transcriptional activity of YAP. Knockout of NLK severely reduced the interaction between YAP and TEAD (Fig 5A). Consistently, reporter activities using a luciferase construct driven by 8xGTIIC promoter [13] were significantly reduced when NLK was knocked down (Fig EV5A). Real-time PCR analysis revealed that the levels of endogenous CTGF and ANKRDI mRNA, well-known YAP/TEAD target genes [13,15], were significantly reduced by knockdown of NLK (Fig EV5B). To investigate the role of NLK in regulation of the transcriptional activity of YAP in vivo, we tested the effects of manipulation of Nemo (Drosophila NLK) levels during fruit fly wing development on Yorkie (Drosophila YAP) target gene expression. Normal expression of expanded, as can be seen in the anterior, anti-Ci marked control regions of the Drosophila wing imaginal discs (Figs EV5C and 5D), is low within the wing pouch and high at the wing pouch periphery [26]. Nemo overproduction leads to dramatic upregulation of expanded within the wing pouch, quantified in Fig EV5C. In contrast, Nemo downregulation reduces the peripheral expression of this target (high magnifications in Fig EV5D). To test whether Nemo downregulation could suppress another Yorkie target gene, whose expression would be more uniform across the wing pouch area, we performed immunostaining against the Yorkie target DIAP1 [7], which reveals a salt-and-pepper
expression pattern within the wing pouch area [27]. This analysis reveals that Nemo downregulation strongly suppressed DIAP1 throughout the RNAi-expressed posterior domain (Fig 5C). Finally, we wished to test the epistasis between Nemo and Yorkie. For this aim, we co-expressed the nemo-targeting RNAi and a Hippo insensitive mutant form of Yorkie (yki[S168A]) in the posterior...
compartment. The resulting discs display typical yki[S168A] phenotype [28] of massive posterior overgrowth and increase in the Yorkie target gene expression (Fig 5D), indistinguishable from discs expressing yki[S168A] alone. This observation indicates that Yorkie is epistatic to Nemo in Drosophila.

Since YAP enhances cell migration and proliferation [29], we tested whether NLK regulates cell migration. Knockdown of NLK significantly reduced cell migration, which depends on YAP because its expression rescued this defect (Fig EV5E). We further tested the effect of NLK on YAP-dependent wound healing as a measure of cell proliferation and migration. Wound healing ability was affected by the phosphorylation status of Ser127 or Ser128. Ectopic expression of YAP-S127A and the phosphorylated mimetic form of YAP at Ser128 (YAP-S128D) in wild-type cells or NLK KO pool cells showed enhanced wound healing ability compared to cells expressing YAP-WT, whereas the S128A mutation produced the opposite effect (Figs 5E and EV5F). Overall, these findings demonstrate that NLK positively regulates YAP activity in physiological settings.

YAP has oncogenic properties and mainly functions as a downstream effector of the Hippo pathway [30]. LATS1/2 phosphorylates YAP at five serine/threonine residues, and phosphorylation of these sites acts as a fundamental input for YAP regulation [11]. Of these phosphorylation sites, Ser127 and S397 are the most functional residues regulating YAP activity. Phosphorylation at Ser127 by LATS1/2 creates a binding consensus site for 14-3-3, resulting in cytoplasmic retention, while phosphorylation on S397 induces β-TrCP-mediated proteasomal degradation of YAP. In addition to LATS1/2 kinases, Akt is also reported to phosphorylate YAP at serine 127, thereby leading to association with 14-3-3 and regulating apoptosis following the cellular damage [31]. More recently, NDR1/2 kinases were identified as YAP-S127 kinases and might function as tumor suppressors upstream of YAP in colon cancer [32]. Conversely, YAP phosphorylation can be reversed by phosphatase, leading to YAP reactivation. Regarding the physiological importance of YAP-S127 phosphorylation, one interesting study has addressed it by introducing a Yap-S112A (corresponding site for the human homologue in mouse) knock-in mutation in the endogenous Yap locus. Surprisingly, although nuclear localization of Yap protein was observed in tissues and cultured cells harboring mutant Yap-S112A, knock-in mice were phenotypically normal probably due to compensatory decrease in YAP protein levels [33]. These findings suggest a homeostatic regulation that maintains physiological YAP activity by yet unidentified mechanisms.

In addition to serine phosphorylation, phosphorylation of YAP at tyrosine residues by Src or c-Abi kinase modulates YAP transcriptional activity [34,35]. Tyrosine phosphatase PTPN14 is able to inhibit YAP transcriptional activity by inducing cell density-dependent translocation of YAP from the nucleus to cytoplasm [36]. A more recent report showed that monomethylation of YAP at K494 by Set7, a SET domain-containing lysine (K) methyltransferase, promotes cytoplasmic retention and inhibition of YAP-dependent

![Figure 4. Subcellular localization and protein level of NLK are regulated by cell density.](image-url)
growth [37]. Taken together, these observations suggest that control of YAP subcellular localization is the sum of multiple regulatory layers and is critical for transactivation. Thus, identifying the molecular regulatory mechanisms of YAP localization will be important to fully understand how the organ size and occurrence of tumors are controlled.
NLK regulates activities of several transcription factors, including TCF/LEF1, NICD, and FOXO, which are key players in diverse signaling pathways [18–20]. Consistently, NLK knockout mice show varying degrees of morphological abnormalities, such as growth retardation, neurological abnormalities, and hematopoietic defects [38]. Here, we identified NLK as a novel kinase for YAP. Our results show that NLK phosphorylates Ser128 of YAP and plays a crucial role in regulating YAP localization and activation. The phosphomimetic form of YAP-S128 showed strong nuclear localization and led to enhanced wound healing ability, whereas substitution of Ser128 with alanine showed the opposite effects. NLK depletion led to significant reduction in the YAP-dependent reporter activity. In the accompanying paper, Dr. Guan and colleagues report similar findings that NLK mediates YAP phosphorylation at Ser128 and nuclear localization by inhibiting association with 14-3-3 upon osmotic stress [39]. Together with their results, our findings suggest that NLK is required for YAP nuclear localization and transcriptional activity in various cellular processes.

We expanded the role of NLK in the regulation of YAP activity in vivo. Knockdown of Nemo (Drosophila NLK) in the fruit fly wing imaginal discs reduced expression of the Yorkie (Drosophila YAP) target genes expanded and DIAP1, while Nemo overproduction produced the opposite effect. The localization of NLK was changed from nuclei to cytoplasm, and the level was reduced along with increasing cell density. It has been shown that dimerization and nuclear localization of NLK are necessary for its activity [25]. Further work is required to explore how the cell density affects NLK nuclear localization and level. It is possible that regulation of subcellular localization and level of NLK at different cell densities could be another mechanism to secure YAP localization and regulation of its transcriptional activity; nuclear NLK phosphorylates Ser128 of YAP and prevents phosphorylation of Ser127/Ser397 by LATS1/2 at LATs1/2, which allows YAP to act as a transcriptional activator in nuclei of low-density cells.

Using phospho-specific antibody and in vitro kinase assay, we observed that NLK phosphorylated YAP at Ser128, which is adjacent to the LATs1/2-mediated phosphorylation site Ser127. More importantly, LATs1/2-mediated phosphorylation at Ser127/Ser397 was inhibited by YAP phosphorylation at Ser128. As shown in Fig 2A, Ser128 is evolutionarily conserved from fruit flies to humans. A previous report examined potential phosphorylation sites within Drosophila Yorkie and observed reduced phosphorylation at Ser168, corresponding to Ser127 of human YAP, in a double-mutant harboring S169D and S172D [28]. This study suggested that phosphorylation at Ser169/Ser172 could reduce Ser168 phosphorylation, thereby promoting Yorkie activity. However, they did not identify endogenous kinase(s) responsible for phosphorylation at Ser169 and Ser172. Although it is unclear whether or not NLK affects phosphorylation at Ser131 (Ser172 of Drosophila Yorkie), our findings provide strong evidence that NLK is an endogenous kinase for phosphorylation at Ser128 that promotes YAP nuclear localization and transcriptional activity. Therefore, we speculate that NLK and LATs1/2 might compete for adjacent phosphorylation residues. Currently, we do not know how the phosphorylation of Ser397 is regulated by NLK, but we think that decrease in phosphorylation on S397 might be due to enhanced nuclear localization of YAP by NLK (Figs 3D and E, and EV4A), which sequesters YAP from phosphorylation by LATs1/2. Overall, our study uncovers a previously unknown kinase involved in regulation of YAP activity and sheds new light on the evolutionarily conserved regulatory mechanism of the Hippo signaling pathway.

Materials and Methods

Plasmids, siRNAs, and CRISPR

Full-length human YAP was PCR-amplified using cDNA of HEK293T cells and inserted into pCMV4-FLAG (Sigma), pEGFP (Clontech), pCS2-HA3 (constructed in our laboratory), and pGEX-4T (GE Healthcare) expression vectors. Point mutation constructs were generated by a PCR-based site-directed mutagenesis method using Pfu polymerase (Promega) and KOD hot start DNA polymerase (Novagen). MST1, WW45, and TEAD1 coding sequences obtained by PCR were subcloned into pEGFP-C1. FLAG-tagged NLK and its mutant constructs were kindly provided by Dr. Ishitani (Kyushu University, Japan). HA-tagged 14-3-3 was a gift from Dr. Hong (Korea University, Korea). CTGF-luciferase was generously provided from Dr. Guan (University of California San Diego, USA). 8xGTTCC-luciferase was a gift from Dr. Lim (Korea Advanced Institute of Science and Technology, Korea). ON-TARGET plus siRNAs targeting human and mouse NLK were purchased by Dharmacon. siGSK3β for human duplexes was purchased by Samchully and its sequence is as follows: sense: 5’-CCCAATGTCACAATCACA-3’ antisense: 5’-TGGTAGTGTGACATTGGG-3’. To generate a guide RNA Vector for NLK knockout using CRISPR-Cas9 system, we inserted two different 20-mer guide RNA sequences (#1 5’-AAAAATGATGCGCGGTCCAAAC-3’ and 5’-TGTGAAGGCCGCTACATTCTTT-3’, #2 5’-ACCACTTCTATCTCCGCGTGGT-3’ and 5’-ACCCGGATGAAGATGGTGT-3’) into pSpCas9(BB)-2A-Puro (a gift from Feng Zhang, Addgene plasmid # 62988).

Cell culture and transfection

HEK293, HEK293T, HeLa, and NIH3T3 cells were maintained in Dulbecco’s modified Eagles’ medium (DMEM) supplemented with 10% fetal bovine serum (FBS, Gibco) and 1× antibiotic-antimycotic (Gibco) and maintained at 37°C in a humidified 5% CO₂ incubator. MCF10A cells were maintained in DMEM/F-12 medium supplemented with 5% horse serum, 20 ng/ml of epidermal growth factor, 0.5 μg/ml of hydrocortisone, 10 μg/ml insulin, 100 ng/ml of cholaer toxin (Sigma), 1× penicillin, and streptomycin (Gibco). Cell transfection was performed using Lipofectamine 2000 and LTX (Invitrogen), Turbofect (Fermentas) or calcium phosphate method.

Immunoprecipitation and immunoblotting

Cell lysates were prepared in lysis buffer (20 mM Tris, pH 7.5, 100 mM NaCl, 1 mM EDTA, 2 mM EGTA, 50 mM β-glycerophosphate, 50 mM NaF, 1 mM sodium vanadate, 2 mM dithiothreitol, 1 mM phenylmethylsulfonyl fluoride, 1 μg/ml of leupeptin, and 1% Triton X-100). Lysates were centrifuged at 14,200 g for 10 min at 4°C, and the supernatant was collected and used for immunoprecipitation and immunoblotting. For immunoprecipitation, 400–800 μg of cell lysate was incubated with antibodies indicated in the figures
and protein A/G plus agarose beads (Santa Cruz Biotechnology). Beads were washed five times with lysis buffer and centrifuged at 30,000 g for 3 min between each wash. Equal amounts of protein were boiled with Laemmli sample buffer and resolved on 6, 8, or 10% SDS–PAGE and transferred to PVDF membrane (Pall). Phos-tag\textsuperscript{TM} acrylamide was prepared from Wako Pure Chemical Industries, and Phos-tag\textsuperscript{TM} SDS–PAGE gel was prepared according to the manufacturer’s protocol.

Mouse monoclonal anti-GST (B-14), anti-YAP (63.7), anti-GFP (B-2), anti-NLK (B-5), pan 14-3-3 (H-8), anti-HA (F-7), and goat polyclonal lamin B (M-20) antibodies were obtained from Santa Cruz. Rabbit polyclonal anti-YAP phospho-128 antibody was generated using recombinant YAP peptide PQHVRAHS(pS)PASLQLG and affinity purification (AbClon). Rabbit polyclonal anti-HA (#A190-108A) antibody was purchased from Bethyl Laboratories. Mouse monoclonal anti-FLAG M2 antibody was obtained from Stratagene. Mouse monoclonal anti-β-catenin (#610154), anti-GSK3β (#610201), and anti-Tead1 (#610923) antibodies were purchased from BD Transduction Laboratories. Rabbit polyclonal anti-β-tubulin (#GTX101715) antibody was obtained from GeneTex. Rabbit polyclonal anti-YAP, anti-phospho-YAP (Ser127), and anti-phospho-YAP (Ser397) antibody were purchased from Cell Signaling Technology. Mouse monoclonal anti-β-actin and rabbit polyclonal anti-NLK (#N8288) antibodies were purchased from Sigma. Mouse monoclonal anti-active-β-catenin (anti-ABC) (#05-665) antibody was obtained from Millipore.

**Immunofluorescence analysis**

HEK293T, NIH3T3, HeLa, and MCF10A cells were seeded onto glass coverslips in 6-well plates, fixed for 20 min in 4% paraformaldehyde in PBS, and then permeabilized for 15 min with 0.1% Triton X-100 in PBS at room temperature. Samples were washed three times with PBS, blocking with 5% BSA (Bio basic) in PBS for 1 h, and then incubated with the indicated primary antibodies for overnight at 4°C. Samples were then rinsed three times with 1% BSA in PBS and incubated with the secondary antibodies Alexa-Fluor antibodies 488 or 633 (Invitrogen) for 2 h. For nuclei staining, Hoechst 33342 solution or DAPI were added to each sample and incubated at 5 min at room temperature. Localization of proteins was analyzed using fluorescence microscopy [Cell Observer (Zeiss)] and confocal microscopy [TCS SP8 (Leica)]. The values for the histogram were calculated using ImageJ software.

**Cellular fractionation**

To separate cytoplasmic and nuclear proteins, cells were washed in phosphate-buffered saline (PBS) and suspended in hypertonic buffer (50 mM HEPES (pH 7.5), 10 mM KCl, 1.5 mM MgCl\textsubscript{2}, 1 mM EDTA, 1 mM Na\textsubscript{2}VO\textsubscript{4}, 50 mM NaF, 1 mM PMSF, 1 μg/ml of leupeptin). After being incubated on ice for 30 min, cells were sheared by being passed 25 times through a 26-gauge needle. The lysates were centrifuged at 800 g for 5 min at 4°C to obtain the cytosolic and membrane fraction (supernatant) and nuclear fraction (pellet). The nuclear pellet was washed three times with hypotonic buffer and resuspended in hypotonic buffer containing 1% Triton X-100 and 150 mM NaCl for 30 min on ice. The supernatant of lysed pellet, which has nuclear proteins, was isolated after centrifugation at 12,000 g for 20 min at 4°C.

**Preparation of recombinant proteins and in vitro kinase assay**

pGEX-YAP was transformed into E. coli BL21 (DE3). For induction of GST fusion protein, 0.2 mM IPTG (Calbiochem) was added to the YTA medium. Purification of GST fusion proteins was performed according to standard procedures (GE Healthcare). For in vitro kinase assays, WT or KM FLAG-NLK was transfected into HEK293T cells, followed by immunoprecipitation using anti-FLAG monoclonal antibody coupled to protein A/G plus agarose beads. Immunoprecipitates were incubated in kinase buffer containing GST-fused proteins, 10 μCi of [γ\textsuperscript{32}P]ATP, 10 mM HEPES (pH 7.4), 1 mM DTT, and 5 mM MgCl\textsubscript{2} at 25°C for 5 min. Samples were resolved by SDS–PAGE, and phosphorylated proteins were detected by autoradiography. In vitro kinase assays without using radioisotope were performed as described previously [25].

**Analysis by nano-LC-ESI-MS/MS**

The protein separated by SDS–PAGE was excised from the gel and digested with endoproteinase chymotrypsin (Roche) in incubation buffer (100 mM Tris–HCl, 10 mM CaCl\textsubscript{2}, pH 7.6) at 25°C for overnight. Following digestion, chymotryptic peptides were extracted with 5% formic acid in 50% acetonitrile solution at room temperature for 20 min. The supernatants were dried and desalted using C18 ZipTips (Millipore, MA) before LC-ESI-MS/MS analysis. Nano-LC (EksigentnanoLC Ultra 2D, Eksigent Technologies, Dublin, CA) separation was conducted under a linear gradient from 3 to 40% solvent B (0.1% formic acid in 100% acetonitrile) with a flow rate of 250 nl/min for 60 min. The column was directly connected to LTQ linear ion-trap mass spectrometer (Thermo Fisher Scientific, San Jose, CA) equipped with a nano-electrospray ion source. All spectra were acquired in data-dependent scan mode. Each full MS scan was followed by five MS/MS scan corresponding from the most intense to the fifth intense peaks of full MS scan. The acquired LC-ESI-MS/MS fragment spectra were searched in the BioWorksBrowserTM (version Rev. 3.3.1 SP1, Thermo Fisher Scientific, San Jose, CA) with the SEQUEST search engines against the data in FASTA format generated from YAP (NCBI accession number NM_001282101.1) in National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/).

**Luciferase assay**

HEK293T cells were seeded in 12-well plates 1 day before transfection. A mixture of 8xGTTC promoter-luciferase reporter, pRL-TK-Renilla, and the indicated plasmids was co-transfected into HEK293T cells. Twenty-four hours after transfection, cells were lysed and luciferase activity was measured using a dual-luciferase reporter assay system (#E1960; Promega) according to the manufacturer’s instructions. Luciferase activity was measured by a GLOMAX 20/20 luminometer (Promega). Transfection efficiency was normalized to thymidine kinase promoter-driven Renilla luciferase (pRL-TK) activity as the internal control.

**Phosphatase treatment**

Cell lysates were prepared for PPase lysis buffer (25 mM Tris–HCl, pH 8.0, 150 mM NaCl, 10% NP-40, 10% glycerol, 0.25%
deoxycylic acid, 1 mM phenylmethylsulfonyl fluoride, and 1 μg/ml of leupeptin). Lysates were incubated with or without λ-PPase (New England Biolabs) in PPase buffer at 30°C for 30 min.

**Real-time PCR analysis**

For quantitative real-time PCR, total RNA was isolated using TRIzol reagent (Invitrogen) according to the manufacturer’s protocol. cDNA was synthesized from total RNA using ImProm-II™ Reverse Transcriptase (Promega) with random primer. Real-time PCR was carried out by an ABI prism 7000 Sequence Detector (Applied Biosystem) with SYBR Green PCR Master Mix (Applied Biosystem). PCR was carried out by an ABI prism 7000 Sequence Detector (Applied Biosystem) with SYBR Green PCR Master Mix (Applied Biosystem). Reverse Transcriptase (Promega) with random primer. Real-time PCR products were designed to amplify specific mRNA fragments and as a displayed unique dissociation (melting) curve. PCR conditions were 95°C (10 min) and 40 cycles at 95°C (30 s) and 60°C (1 min). The threshold cycle (Ct) value for each gene was normalized to the Ct value for β-actin or GAPDH. Relative mRNA expression was calculated using the ΔΔCT method. Primers used for human cell lines: ANKK1, 5′-CACCTTCTAGCCACCCCTGTA-3′ and 5′-CCACAGGTCTCGTAGATGTTT-3′; CTGF, 5′-CCTAGCAACGCCCTCTGT-3′ and 5′-GAGTCTTCTCCGGTCAACCA-3′; NLK, 5′-CCAGT GACCTTTGGACCTGTC-3′ and 5′-GATGGCTGAGCAACCAGTG-3′; and β-ACTIN, 5′-GGGGAATTCGTCGTGACATT-3′ and 5′-GATGAGTGAAGTTCTTTGCG-3′.

Primers used for mouse cell lines: Amotl2, 5′-AGAGATTGGAATCGCCAAAC-3′ and 5′-TCTCTCCTTCTCTTGGTCTG-3′; Ctgf, 5′-CAAGGCCAGCACAGCTTTG-3′ and 5′-AGAACGGCCCTCCACTG-3′; Cty61, 5′-ATGATGATCCAGTCCTGCAA-3′ and 5′-ATGATGATCCAGTCCTGCAA-3′; and 5′-TTCTCCTGTTCCTGTTGCTG-3′; and 5′-GATGAGTGAAGTTCTTTGCG-3′.

**Migration assay**

HeLa cells (3 × 10⁴ cells) were plated in the upper chamber of cell culture inserts (SPL, 8-μm membrane pore size) with serum-free media. Cell growth medium containing 5% serum was added to the lower well. After 24-h incubation, cells were fixed for 2 min in 3.7% formaldehyde in PBS and then permeabilized for 20 min with 100% methanol at room temperature. After removal of non-migratory cells, migrated cells were stained with 0.5% crystal violet. Migrated cells were counted using an inverted microscope, and the values for histogram were calculated using ImageJ software.

**Wound healing assay**

HEK293 and NLK KO pool cells were transfected with control or YAP mutants in 24-well plates and cultured until full confluence. Cells were scratched by 200-μl pipette tip and then washed with media to remove cell debris, and the media was replaced. Photographs of the wound region were taken at 0 and 24 h. Cell free area was measured using TSratch software [40].

**Experiments with Drosophila**

Analysis of the effects of manipulations of Nemo levels on Yorkie target gene expression was performed following [41], whereas the posterior-specific hedgehog-Gal4 (hh-Gal4) or en-gaughter-Gal4 (en-Gal4) drivers were used to induce overexpression or downregulation of nemo using the UAS-nmo [17] or UAS-nmo-RNAi lines (104885KK or 3002GD [42]), respectively. UAS-yki[S168A] was used as described [28]. In third instar larval wing imaginal discs, these posterior-specific manipulations were compared to the unaffected anterior halves of the discs, which served as internal controls. expanded-LacZ [43] and DIAP1 were used to monitor Yorkie target gene expression. Mouse anti-DIAP1 [27], and anti-Ci, anti-β-gal, and anti-Hh antibodies were used as described [41]. Quantification of fluorescent intensity after confocal microscopy was performed with ImageJ.

**Statistical analysis**

All data analysis in this study was carried out using GraphPad Prism7 (San Diego, CA, http://www.graphpad.com) for Mac OS. Quantifications were done from at least three independent experiment groups. Statistical analysis between groups was assessed by two-tailed Student’s t-test to determine significance. P-values < 0.05 were considered significant. Error bars on all graphs are presented as standard error of the mean (SEM) unless otherwise indicated.

**References**


