Molecular Evolution of the Yap/Yorkie Proto-Oncogene and Elucidation of Its Core Transcriptional Program

Aissam Ikmi, Bjoern Gaertner, Christopher Seidel, Mansi Srivastava, Julia Zeitlinger, and Matthew C. Gibson*

1Stowers Institute for Medical Research, Kansas City, MO
2Whitehead Institute for Biomedical Research, Cambridge, MA
3Department of Pathology and Laboratory Medicine, University of Kansas School of Medicine, Kansas City, KS
4Department of Anatomy and Cell Biology, University of Kansas School of Medicine, Kansas City, KS

*Corresponding author: E-mail: MG2@stowers.org.

Abstract

Throughout Metazoa, developmental processes are controlled by a surprisingly limited number of conserved signaling pathways. Precisely how these signaling cassettes were assembled in early animal evolution remains poorly understood, as do the molecular transitions that potentiated the acquisition of their myriad developmental functions. Here we analyze the molecular evolution of the proto-oncogene yes-associated protein (Yap)/Yorkie, a key effector of the Hippo signaling pathway that controls organ size in both Drosophila and mammals. Based on heterologous functional analysis of evolutionarily distant Yap/Yorkie orthologs, we demonstrate that a structurally distinct interaction interface between Yap/Yorkie and its partner TEAD/Scalloped became fixed in the eumetazoan common ancestor. We then combine transcriptional profiling of tissues expressing phylogenetically diverse forms of Yap/Yorkie with ChIP-seq validation to identify a common downstream gene expression program underlying the control of tissue growth in Drosophila. Intriguingly, a subset of the newly identified Yorkie target genes are also induced by Yap in mammalian tissues, thus revealing a conserved Yap-dependent gene expression signature likely to mediate organ size control throughout bilaterian animals. Combined, these experiments provide new mechanistic insights while revealing the ancient evolutionary history of Hippo signaling.

Key words: Yap, growth control, target genes, Metazoa, evolution.

Introduction

Recent advances in comparative genomics have potentiated new insights into the evolution of animal multicellularity (Putnam et al. 2007; King et al. 2008; Srivastava et al. 2008, 2010), with a primary focus on molecules that maintain stable cell–cell interactions and cell differentiation (King et al. 2003; Nichols et al. 2006; Abedin and King 2008; Larroux et al. 2008; Sebe-Pedros et al. 2010). Although these fundamental processes are essential to form an epithelium composed of different cell types, the evolution of animal complexity must have also required the incorporation of novel tissue growth-regulatory mechanisms. In both insects and mammals, the Hippo tumor suppressor pathway serves such a function by antagonizing the growth-promoting activity of a transcriptional coactivator known as Yorkie (Yki) in Drosophila and yes-associated protein (Yap) in mammals (Dong et al. 2007; Oh and Irvine 2010; Zhao et al. 2010). Importantly, consistent with its function in control of tissue growth, Yap is a candidate oncogene in human disease (Overholtzer et al. 2006; Zender et al. 2006). In addition, several lines of evidence suggest that Yap also plays a critical role in other biological processes, including cell fate determination, stem cell proliferation, and regeneration (Zhao et al. 2011; Liu et al. 2012).

At the molecular level, Yap contains multiple functional domains, including TEAD-binding (TBD), WW, coiled-coil, and PDZ-binding motifs (Wang et al. 2009). To promote growth, Yap interacts with Scalloped/TEAD and other DNA-binding partners to drive the expression of cell cycle regulators and cell death inhibitors (Huang et al. 2005; Goulev et al. 2008; Wu et al. 2008; Zhang, Ren, et al. 2008; Zhao et al. 2008; Peng et al. 2009; Oh and Irvine 2011). These interactions require Yap’s TBD and WW motifs (Zhao et al. 2009; Zhang, Milton, et al. 2009). The growth-promoting activity of Yap is in turn constrained through phosphorylation by Warts/Lats (Huang et al. 2005; Dong et al. 2007), a member of an ancient eukaryotic kinase cassette including Hippo/Mst (Sebe-Pedros et al. 2012). Yap phosphorylation induces cytoplasmic retention by recruiting 14-3-3 proteins (Camargo et al. 2007; Dong et al. 2007; Zhao et al. 2007), which then limit the ability of Yki/Yap to complex with its DNA-binding partners.

Recently, the identification and functional analysis of Yap and TEAD from the amoeba Capsaspora owczarzaki suggests that the capacity to control tissue growth may have emerged through co-option of a preexisting Hippo-Yap regulatory architecture. However, unlike Human Yap, the Capsaspora ortholog alone is not sufficient to induce tissue overgrowth in Drosophila (Dong et al. 2007; Sebe-Pedros et al. 2012).
This raises the question of how and when the Yap-TBD changed during evolution. Here, we compare the structure of the TBD from phylogenetically informative lineages including multiple early branching metazoan species, as well as the closest unicellular relatives of Metazoa. We then use a heterologous expression assay to 1) directly compare the growth-promoting activity of divergent Yap orthologs and 2) identify a downstream transcriptional profile induced by select variants in the Drosophila eye disc. Combined, these results demonstrate that the Yap-TEAD interaction interface became stabilized sometime after the divergence of sponges from the eumetazoan common ancestor. In addition, coupled with Chip-seq validation of Yki/Scalloped binding sites, our comparative analysis identifies multiple novel Yap/TEAD targets in Drosophila while hinting at the existence of a conserved bilaterian gene expression program downstream of Yap/TEAD.

Results

Evolutionary Changes in Yap/Yki Protein Architecture

To determine the extent to which the structural features of Yap are conserved between animals and their unicellular relatives, we performed a detailed domain composition analysis of Yap orthologs from species occupying key phylogenetic positions (fig. 1A). In addition to Drosophila and Homo, these included the cnidarian Nematostella vectensis, the placozoan Trichoplax adhaerens and the demosponge Amphimedon queenslandica, which are modern representatives of the earliest branching Metazoa (Putnam et al. 2007; Srivastava et al. 2008, 2010). We also analyzed the domain structure of Yap orthologs from the genome of three nonmetazoan species, the amoeba Monosiga brevicollis and Salpingoeca rosetta (King et al. 2008; Suga et al. 2013). Consistent with prior findings (Sebe-Pedros et al. 2012), our phylogenetic analyses showed a well-supported monophyletic group that included the known bilaterian Yap protein together with a single putative Yap protein from each analyzed genome (supplementary fig. S1, Supplementary Material online). This contrasts with vertebrate genomes, which contain a paralogous copy of Yap (Taz), proposed to have originated from a gene duplication event early in vertebrate evolution (Hilman and Gat 2011).

Interestingly, the most critical Lats/Warts phosphorylation site in Yap (corresponding to Ser127 in Homo) was conserved in all metazoan species analyzed (fig. 1B). It was also present in the Monosiga, Salpingoeca, and Capsaspora Yap proteins, although the associated regulatory 14-3-3 binding motif was incomplete (fig. 1B). These observations are consistent with conservation of the Yap-Lats regulatory interaction throughout animal evolution.

As expected, the Yap orthologs we identified shared many additional features. However, they also displayed some critical structural variations. Most prominently, these included alterations in the architecture of the TBD (fig. 1A and supplementary fig. S2, Supplementary Material online). Although the TBD α1-loop-α2 secondary structure (Chen et al. 2010; Li et al. 2010) was conserved in Eumetazoa, this protein—protein interaction domain diverged in Amphimedon, exhibiting an additional α3 helix motif (fig. 1C and D). Beyond metazoans, the most conspicuous conserved motifs were the Yap WW and coiled-coil domains (fig. 1A), indicating that this structural combination was found in the unicellular ancestors of animals. Intriguingly, we manually identified α1 and α2 helix motifs as a cryptic TBD in the N-terminus of Capsaspora Yap (fig. 1C and D). This may represent a transitional state that existed before evolution of the complete α1-loop-α2 structure in Eumetazoa. In addition, a highly divergent Yap TBD containing only the α1 helix motif was found in Monosiga (fig. 1D). Combined, these results suggest a remarkable structural plasticity of the TBD during early Yap evolution, followed by fixation of the α1-loop-α2 structure throughout Eumetazoa.

Heterologous Functional Analysis of Yap/Yki Orthologs in Drosophila

We next set out to determine the functional implications of the evolutionary changes in Yap protein architecture. Because experimental tools are a limiting factor in most nonmodel organisms, we took advantage of Drosophila genetics to directly compare the activities of representative Yap orthologs in a uniform in vivo system. We first generated transgenic flies carrying inducible forms of yap from Monosiga, Amphimedon, Trichoplax, and Nematostella, as well as inducible forms of Drosophila yki and Homo yap for experimental controls. The phiC31-attP-attB site-specific integration technique was employed to insert all transgenes into a specific genomic site, ensuring identical transgene expression levels (Groth et al. 2004; Bischof et al. 2007). Because Yap activity is regulated by Lats/Warts phosphorylation, we also generated hypothetically nonphosphorylatable forms of each Yap ortholog by modifying the appropriate serine residues to create constitutively active Yap variants (fig. 2A). Although the mutated serine for each metazoan Yap matched the position of the critical phosphorylation site of Yki/Yap (Ser168/127), the only candidate residue within an optimal 14-3-3 binding motif for the Monosiga protein was Ser48.

Yki/Yap misexpression is sufficient to induce tissue overgrowth in Drosophila (Huang et al. 2005; Dong et al. 2007). We therefore performed a heterologous overexpression assay to compare the activity of each Yap ortholog in the Drosophila eye, using Glass Multiple Reporter-Gal4 (GMR-Gal4) to drive tissue-specific expression. Contrasting with the effects of Monosiga Yap, the Amphimedon, Trichoplax, and Nematostella variants elicited distinct degrees of overgrowth (fig. 2A and B). Importantly, each of these orthologs exhibited roughly equivalent protein stability in Drosophila S2 cells, indicating that the observed phenotypic variability arose from protein-intrinsic properties (fig. 2C). Trichoplax Yap produced an exceptionally severe overgrowth phenotype with enlarged, folded adult eyes. This effect was even further enhanced in animals overexpressing Trichoplax YapS81A (fig. 2A and B). We also found evidence for phosphorylation of Yap from Amphimedon and Nematostella; both showed similar
increases in eye size that were enhanced by mutation of the conserved phosphorylation sites (fig. 2A and B). Together, these results not only show that basal metazoan versions of Yap possess potent growth-promoting activity in Drosophila but also that they are regulated by phosphorylation via similar mechanisms to those of Drosophila Yki and human Yap. This implies that the Hippo/Wts cassette may function similar to phosphoregulate Yap in early branching metazoans. Consistent with this, we cloned a Warts/Lats ortholog from Nematostella and found that it was sufficient to rescue the corresponding Drosophila mutant, presumably through phosphorylation of Drosophila Yki (supplementary fig. S3, Supplementary Material online).

Surprisingly, in contrast to the metazoan Yaps, overexpression of Monosiga Yap or its phosphomutant form resulted in significantly reduced eye size (fig. 2A and B). Although the proximal cause of this reduced eye phenotype is still unknown, cell clones overexpressing Monosiga Yap exhibited normal growth and morphology (data not shown). These findings exclude the possibility that Monosiga Yap acts as a dominant negative to inhibit cell proliferation through effects on endogenous Yki activity.
Cellular and Molecular Basis for Yap/Yki Ortholog-Induced Overgrowth

To determine the cellular basis for eye size increases induced by each Yap ortholog, we next performed EdU incorporation assays to directly measure cell proliferation. As expected from our analysis of adult eye size, Trichoplax YapS81A induced the most extensive cell proliferation in the GMR-Gal4 domain (fig. 3A). As a consequence of this dramatic overgrowth, the eye disc epithelium was extensively folded and disorganized (fig. 3D and E). Nematostella YapS83A induced significant but moderate ectopic cell proliferation when compared with Drosophila YkiS127A and human YapS127A (fig. 3A), resulting in extra bristles and interommatidial cells (IOCs) in the adult eyes (fig. 3D and E). Consistent with a Yki-like transcriptional output of Trichoplax and Nematostella Yap, both induced expression of Cyclin E and the cell death inhibitor diap1 (fig. 3B, C, and C'). These findings show that Nematostella and Trichoplax Yap are able to induce cell proliferation and survival, behaving like their Drosophila and human counterparts.

Contrasting with their eumetazoan counterparts, Amphimedon YapS95A and Monosiga YapS48A did not induce Cyclin E, diap1, or cell proliferation as indicated by EdU incorporation (fig. 3A, B, C, and C'). Further, overgrowth...
FIG. 3. Cellular and molecular mechanisms underlying Yap ortholog-induced overgrowth in Drosophila. (A) Edu labeling in eye discs overexpressing hypothetically nonphosphorylatable forms of Yap from the indicated species under the control of GMR-Gal4. Note the dramatic induction of proliferation by Trichoplax yap and the absence of ectopic proliferation caused by the Monosiga and Amphimedon orthologs. (B) Immunostaining of Cyclin E in eye discs of the same genotypes indicated above. Arrowheads indicate the position of the morphogenetic furrow.
was not induced following misexpression in other imaginal discs (data not shown). Intriguingly, pupal retinae of GMR-Gal4 $\rightarrow$ Amphimedon Yap$_{S95A}$ animals exhibited an elevated number of IOCs, which are normally eliminated by developmentally programmed apoptosis (fig. 3D and F) (Wolff and Ready 1991). Following expression of Amphimedon Yap$_{S95A}$, these cells failed to undergo normal differentiation of the lens cuticle and retained a pupal-like appearance in adult animals (fig. 3E). More pronounced defects in the differentiation and patterning of ommatidial cells were observed in retinae overexpressing Monosiga Yap, as the normal hexagonal topology of ommatidial subunits was abolished (fig. 3D and E). These results suggest a defect in retinal differentiation and are consistent with severe reductions of eye pigmentation observed in flies overexpressing Monosiga and Amphimedon proteins (fig. 2A).

The Amphimedon and Monosiga Yap proteins both possessed an incomplete $\alpha$1-loop-$\alpha$2 TBD (fig. 1C), and thus their inability to drive proliferative growth was most likely due to a failure to recognize the endogenous Drosophila TEAD (Scalloped, Sd). To test this hypothesis, we first constructed chimeric forms of Amphimedon and Monosiga Yap bearing the 50-amino acid human TBD. Strikingly, in both cases, this single change in protein architecture was sufficient to induce extensive ectopic proliferation in the eye disc (fig. 3G and H). Second, we found that Amphimedon Yap was able to form a functional protein complex with Amphimedon TEAD in Drosophila S2 cells (supplementary fig. S4, Supplementary Material online). In parallel, as expected from our in vivo heterologous expression assay, we did not detect a physical interaction between Amphimedon Yap and Drosophila Sd (supplementary fig. S4, Supplementary Material online), despite the fact that TEAD proteins exhibit few differences in their Yap-binding interfaces (supplementary fig. S5, Supplementary Material online). These findings suggest a critical functional divergence in Yap-TEAD pairwise interactions during early animal evolution. Consistent with this scenario, Capsaspora Yap cannot induce tissue overgrowth in Drosophila unless it is co-overexpressed with the endogenous Capsaspora TEAD (Sebe-Pedros et al. 2012). In contrast to the Amphimedon Yap, Nematostella Yap was able to physically interact with both Nematostella TEAD and Drosophila Sd (supplementary fig. S4A, Supplementary Material online). Thus, these findings not only show the deep evolutionary origins of Yap-TEAD activity but also reveal that the interaction interface between Yap and TEAD changed during early metazoan evolution and ultimately became fixed in eumetazoan Yap proteins. Although the evolutionary advantages of this modification remain unknown, we speculate that it may have served as an important adaptation of Yap to critical new roles in the growth control in the multicellular context.

The Transcriptional Output of Yap/Yki Orthologs in Drosophila

In vivo, Yap acts through modulation of target gene expression (Overholtzer et al. 2006; Dong et al. 2007; Hao et al. 2008; Lu et al. 2010). To leverage the evolutionary diversity of selected Yki/Yap orthologs and identify novel transcriptional targets of the Hippo pathway in Drosophila, we performed RNA sequencing analysis (RNA-seq) on total RNA isolated from GMR-Gal4 $\rightarrow$ Yap eye discs (fig. 4A and supplementary fig. S6A–E, Supplementary Material online). We first determined the endogenous targets of GMR-Gal4-driven Drosophila Yki and then utilized the highly active Trichoplax Yap and Monosiga Yap containing the human TBD (Yap$_{\text{TBD}}$) to define core target genes activated in common. In addition, we used Monosiga Yap, which was not able to induce ectopic proliferation, as a negative control to remove the transcriptional background resulting from protein overexpression. For each of these conditions, the transcriptional profile was compared with that of a GMR-Gal4 $\rightarrow$ UAS-GFP control.

RNA-seq analysis identified a robust gene expression signature common to eumetazoan Yaps as well as the form of Monosiga Yap$_{\text{TBD}}$ (fig. 4B; supplementary table S1, Supplementary Material online). Validating our approach, this signature comprised 213 commonly upregulated targets that included almost all previously known Yki target genes (cycin E [Tapon et al. 2002], expanded [Hamaratoglu et al. 2006], e2f1 [Goulev et al. 2008], wingless [Cho et al. 2006], dally [Bena-Lopez et al. 2008], kibra [Genevet et al. 2010], and vein [Zhang, Ji, et al. 2009]), along with 258 commonly downregulated factors (fig. 4B). Interestingly, most of these genes were conserved in the Trichoplax and Monosiga genomes (supplementary table S2, Supplementary Material online), perhaps representing an ancient Yap-dependent gene expression signature. Among the 213 commonly upregulated targets, 16 genes were subsequently validated by real-time quantitative PCR analysis (supplementary fig. S7, Supplementary Material online). Although we observed a highly significant overlap between eumetazoan Yap target genes in Drosophila, only 4% and 11% of these genes were correspondingly up- and downregulated by Monosiga Yap, respectively (fig. 4B). This
FIG. 4. The transcriptional program downstream of Yap ortholog induction. (A) Experimental design for RNA-seq experiment. *Drosophila* Yki (YkiS168A), *Trichoplax* Yap (TaYapS81A), chimeric Monosiga Yap with the human TBD (MbYap+TBD), and Monosiga Yap (MbYap) were co-overexpressed with UAS-GFP using GMR-Gal4. Control is GMR-Gal4 > UAS-GFP. For each experimental condition, total RNA was extracted from surgically dissected GMR expression domains (GFP+). (B) Four-way Venn diagrams of differentially upregulated (left) and downregulated (right) genes in each overexpression condition. The number of genes up- and downregulated is indicated under each transgene name. The numbers of commonly up- and downregulated genes in YkiS168A, TaYapS81A, and MbYap+TBD are indicated in green boxes. (C, D) Matrices of gene ontology of biological processes analysis for genes upregulated (Up) and downregulated (Dn) in each overexpression condition.
indicates that the transcriptional outputs of Yap with and without the α1-loop-α2 TBD are fundamentally distinct. One notable exception was that expanded (ex), a well-defined target of Yki in Drosophila, was 1.4-fold upregulated following Monosiga Yap expression compared with more than 3-fold induction by metazoan Yap variants (supplementary table S1, Supplementary Material online).

In agreement with the phenotypes induced by Yki/Yap expression (figs. 2A and 3A), GO term analysis indicated that a significant number of commonly upregulated genes were involved in DNA replication, cell cycle, and growth regulation processes (fig. 4C). Interestingly, we found that the average expression level of the top 100 commonly upregulated genes induced by Trichoplax Yap and Monosiga Yap TBD was higher than that induced by Drosophila Yki (supplementary fig. S6, Supplementary Material online). These quantitative differences in target gene expression level may account for the more extensive cell proliferation induced by these Yap variants (fig. 3A). By contrast to the enrichment of upregulated genes for cell proliferation processes, downregulated genes were enriched for the process of nervous system development (fig. 4D). This is consistent with the observed delays in cell cycle exit and retinal determination/differentiation as a consequence of Yap expression in the eye (fig. 3D and E).

Except in the case of Monosiga Yap TBD, we did not identify significant Yap ortholog-specific enrichment for biological processes (supplementary fig. S8, Supplementary Material online).

**Genome-Wide Distribution of Yki and Sd on Chromatin**

In principle, the Yki/Yap dependent gene expression profiles described above could be the result of primary, secondary, or tertiary transcriptional events. To extend the results of our comparative studies and determine which induced genes were most likely direct targets, we performed a ChIP-Seq experiment to identify the genome-wide occupancy of Drosophila Yki in proliferating cells. For chromatin immunoprecipitation, we generated a novel antibody that specifically detects Yki protein (fig. 5A, B-B″, and C-C″) and analyzed dissected eye discs of the genotype GMR-Gal4 > UAS-Yki 1-loop-A. In parallel, we performed a ChIP-Seq experiment using Polymerase II (Pol II) antibody as a control. Similar to recent ChIP-Seq analyses of wild-type Yki in wing and eye discs (Oh et al. 2013; Slattery et al. 2013), Yki was enriched with high confidence at a large number of sites throughout the genome (using Model-based Analysis for ChIP-Seq [MACS], P < 0.001; supplementary fig. S9, Supplementary Material online). Binding peaks were found in proximity to 5,732 genes (supplementary table S1, Supplementary Material online) and a large number of them were localized within promoter regions (supplementary fig. S10A, Supplementary Material online).

To focus our analysis on bona fide targets of the Drosophila Yki/Sd complex, we also generated a novel antibody directed against Sd and performed ChIP-seq analysis under the same conditions (fig. 5D, E-E″, and F-F″). The Sd binding peaks were specific because they were highly enriched for the Sd motif (supplementary fig. S10B, Supplementary Material online). Among the 1,238 genes bound by Sd, 97% were also bound by Yki (supplementary fig. S10C and D, Supplementary Material online), and this high-confidence subset included all previously known target genes (supplementary fig. S11, Supplementary Material online).

To identify novel Yki target genes, we next focused on the overlap between our RNA-seq and ChIP-seq data sets (supplementary fig. S10C and D, Supplementary Material online). We found that Yki and Yki/Sd peaks were highly enriched near upregulated genes (the common set from figure 4B, P < 3 × 10−16; Fisher test) but not near downregulated genes (P = 0.3 for Yki and P = 0.08 for Yki/Sd). Interestingly, the genes induced in common by different Yap orthologs showed a higher enrichment of Yki and Yki/Sd peaks (78% and 32%, respectively) compared with the analysis of solely Yki-induced genes (64% for Yki peaks and 24% for Yki/Sd peaks). These results corroborate the predominant function of Yki/Yap as a transcriptional activator and led us to the identification of several novel putative targets. Most prominently, the Hippo pathway components warts (wts), fat (ft), and dachsous (ds) were commonly upregulated targets and exhibited both Yki and Sd peaks, revealing the existence of novel negative feedback loops controlling Yki activity (fig. 6A). Another class of putative direct target genes included the signaling ligands decapentaplegic (dpp), wnt6, spatzle 3 (spz3), vascular endothelial growth factor 2 (pgf2), thrombospondin (tsp), tissue inhibitor of metalloproteinase (Timp), and neuropeptide-like precursor 1 (nplp1) (fig. 6B). Our analysis also highlighted the amino acid transporter pathic (path) and two insulin-signaling components as potential growth-regulatory factors targeted by Yki/Sd activity (insulin receptor (Inr) and forkhead box protein O (foxo); fig. 6B). In addition to these Yki/Sd target genes, our ChIP-seq data clearly indicate the existence of an Sd-independent downstream program that is most likely the result of Yki’s putative interactions with other DNA-binding partners, such as Homothorax, Teashirt, and Mad (Peng et al. 2009; Oh and Irvine 2011). Indeed, using a recent ChIP-Seq analysis of Homothorax in eye discs (Slattery et al. 2013), we found that 40% of these Sd-independent targets were co-bound by Yki and Homothorax (supplementary table S3, Supplementary Material online). Together, these findings reveal a complex network of factors downstream of Yap activation and directly link Hippo signaling to the regulatory architecture for a wide variety of processes required for tissue growth in vivo.

**A Yap/Yki-Induced Transcriptional Program Shared by Drosophila and Vertebrates**

In vertebrates, Yap-induced genes have been identified by microarray profiling of hepatocytes, fibroblasts, and breast epithelial cells (Overholtzer et al. 2006; Dong et al. 2007; Hao et al. 2008; Lu et al. 2010). To interpret our transcriptional profiling results at an evolutionary level and investigate the extent to which Yap’s transcriptional output is conserved, we took advantage of the OrthoDB catalog of eukaryotic orthologs (Waterhouse et al. 2011) to test for orthologous
Validation of anti-Yki and anti-Sd antibodies. (A) Western blot analysis shows that anti-Yki antibody detects a strong signal at the expected molecular weight (arrowhead, 45 kDa). (B-B') Control wing disc stained with Hoechst (blue) and anti-Yki (red). The posterior compartment is marked by the expression of hh-Gal4 > UAS-GFP (green). Anti-Yki detects ubiquitous expression of Yki. (C-C') Wing disc overexpressing UAS-yki-RNAi under the control of hh-Gal4. The clear reduction of anti-Yki staining in the posterior compartment (inset box) confirms that our antibody recognizes Drosophila Yki. (D) Anti-Sd detects a specific band at the expected molecular weight (arrowhead, 50 kDa). (E-E') Control wing disc stained with Hoechst (blue) and anti-Sd (red). Endogenous sd expression is elevated in the wing pouch and margin, which is consistent with our Anti-Sd staining. (F-F') A wing disc expressing UAS-sd-RNAi under the control of hh-Gal4 shows a strong reduction of Sd staining in the posterior compartment (inset box), confirming the specificity of Anti-Sd.
relationships between genes induced by Yap in mouse liver (Dong et al. 2007) and those in our Drosophila RNA-seq data. Strikingly, despite the cell type differences and the substantial evolutionary distance between these two organisms, 74 Yap-induced genes from the mouse experiments were also significantly upregulated in Drosophila (supplementary table S4, Supplementary Material online). At least 63 and 31 of these 74 genes show Yki/Yap binding in Drosophila and

![Diagram](Image)

**Fig. 6.** Chromatin localization of Yki and Sd on novel target genes. Plots of Yki (orange), Sd (blue), and PolII (black) occupancy peaks in selected commonly upregulated genes from the RNA-seq data. (A) Target genes belonging to novel negative feedback loops. (B) Novel target genes. Transcriptional units of target genes are highlighted in black and their neighbor genes are in gray. Pink and gray bars indicate significant Yki/Sd and Yki peaks, respectively.

Ikmi et al. doi:10.1093/molbev/msu071 MBE

Downloaded from https://academic.oup.com/mbe/article-abstract/31/6/1375/1009129 by guest on 04 February 2019
Our results also define the Yap-TEAD interaction as a new model system for understanding the coevolution of protein complexes during the emergence of animal multicellularity. Interestingly, a similar evolutionary scenario was recently described for another central growth regulator, the Myc-Max complex. However, cross-species interactions between Monosiga and human Myc and Max were not detected (Young et al. 2011). Thus, we propose that the structural changes in these protein complexes (Yap-TEAD and Myc-Max) may have served as critical adaptations for multicellular growth control during their co-option into an ancient metazoan gene regulatory architecture.

The presence of the most critical Warts/Lats phosphorylation site in all Yap orthologs (fig. 1B) suggests the conservation of Lats-Yap phosphorylation. Consistent with this, we observed an enhancement of Yap activity following the modification of this critical site in the metazoan forms of Yap (fig. 2A and 8). In addition, it has been reported that the activity of Capaspsora Yap is regulated through phosphorylation, pushing back the origin of Lats-Yap phosphorylation to the unicellular ancestors of animals (Sebe-Pedros et al. 2012). Because it is thought that the modern Hippo/Lats pathway responds to extracellular cues to regulate tissue growth, it remains unclear what function this pathway may have served in a hypothetical unicellular ancestor of animals. One possibility is that the pathway regulated growth in multicellular aggregates or in response to local density of specific cell types. Analyses that examine the downstream transcriptional output of Yap TEAD signaling in close metazoan relatives could shed light on this important issue.

Beyond the evolutionary implications of these results, our functional phylogenetic approach also provided mechanistic insights into the contemporary Hippo pathway itself. Although recent studies have analyzed the genome-wide occupancy of Yki during normal development (Oh et al. 2013; Slattery et al. 2013), here we used a combination of RNA-seq and ChiP-seq experiments to identify novel Yki and Sd target genes induced during cell proliferation. Indeed, we report the existence of a core gene expression signature underlying the control of tissue growth in Drosophila (fig. 4B and supplementary table S1, Supplementary Material online). This signature includes novel target genes that could mediate cross talk with key signaling pathways, as well as multiple feedback loops controlling Yap activity (fig. 7). Further experimental analyses would be required to know when and where Yki regulates these novel targets during normal development.

On a broader note, although it is widely accepted that the incredible diversity of animal development is directed by a limited number of conserved signaling pathways (Pires-daSilva and Sommer 2003), it remains unclear whether these pathways act, at least partially, through conserved downstream genetic programs. By comparing the transcriptional output of Yap in Drosophila and mouse, we identified a conserved set of target genes commonly induced in these evolutionarily distant species (supplementary table S4, Supplementary Material online). These targets may represent an ancestral gene-expression signature of Yap and constitute candidate effectors of Yap-TEAD activity in other metazoans.
Components (red).

Known target genes  Novel target genes  Negative feedback loops
cycE, ed21, wg, daily, vein, diap1

Fig. 7. Novel downstream target genes of Yki in Drosophila. Schematic representation of the Fat-Hippo pathway in Drosophila. In response to dachshous (Ds) binding, fat (Ft) protocadherin activates the Hippo pathway, potentially through expanded and warts (dashed line). The Expanded-Merlin-Kibra complex (Ex-Mer-Kbr) also activates the kinase cascade leading from the Hippo/Warts (Hpo/Wts) kinases and their scaffolding proteins Salvador (Sav) and Mod as tumor suppressor (Mats) to Yki and its transcriptional cofactors (TF). Listed below are examples of putative target genes that met the dual criteria of Yki/Sp chromatin association and were upregulated in common following Yki\(^{S168A}\), YoYap\(^{S161A}\), and MtYap\(^{TBD}\) overexpression. Yap ortholog-induced genes were divided into three classes: known target genes, novel target genes (green), and candidate negative feedback loop components (red).

Additional analyses would be required to characterize these potentially conserved targets in depth. Combined, these results illustrate the power of comparative studies to provide both evolutionary and mechanistic insight into fundamental biological processes.

Materials and Methods

Bioinformatics

We identified Yap genes using the basic alignment sequence tool (Blast: TBLASTN, TBLASTX, and BlastP) with human Yap as a query. The genomes of N. vectensis, A. queenslandica, and M. brevicollis are available in http://genome.jgi-psf.org/Nemve1, http://genome.jgi-psf.org/Triad1 (last accessed February 23, 2014) (spongezome.metazome.net), and http://genome.jgi-psf.org/Monbr1 (last accessed February 23, 2014). Capsaspora owczarzaki and S. rosetta genome assemblies were examined on the Broad Institute web site (http://www.broadinstitute.org/, last accessed February 23, 2014). Reciprocal best Blast searches and protein domain structure analyses (Pfam: http://pfamsanger.ac.uk/search [last accessed February 23, 2014] and SMART: http://smart.embl-heidelberg.de [last accessed February 23, 2014]) were used to screen for positive hits. To identify the TBD, we conducted a protein structure homology analysis using the SWISS-MODEL automated comparative protein-modeling server (http://swissmodel.expasy.org, last accessed February 23, 2014). Yap orthologs were aligned using MUSCLE (Edgar 2004) with default settings. For phylogenetic analyses, the alignment was manually curated to only retain the two well-conserved WW domains. Neighbor-joining trees were generated using PHYLIP with default settings and 10,000 bootstrap replicates. Maximum likelihood analyses were run with 1,000 bootstrap replicates using PhyML with the Whelan and Goldman (WAG) model of amino acid substitution, four substitution rate categories, and the proportion of invariable sites estimated from the data set. Bayesian inference methods were implemented using MrBayes v3.1.2 with a mixed amino acid model prior and a variable rate prior.

Cloning

The full-length coding sequence of Nematostella Yap was amplified from larval cDNA using the following primers: Nematostella-YapF: 5' - TTCAATGACGAAAGAAAC A-3'; Nematostella-YapR: 5' - TCGGACCTAACAAGAACTGTTA A-3'. The amplified cDNA was cloned into the pCRTM4-TOPO vector (Invitrogen). The coding sequences of Yap from Monosiga, Amphimedon, Trichoplax, Drosophila, and Human were synthesized by GenScript (Piscataway, NJ). Amphimedon Yap was also cloned from larval cDNA into the pCRTM4-TOPO vector using the following primers: Amphimedon-YapF: 5' - ATGCATGATTATCTAACTACAGGAATCCT TCC-3'; Amphimedon-YapR: 5'-CACCCAAGATTACTACCACCATATTCC-3'. To generate the hypothetical nonphosphorylatable form of each protein, serine to alanine mutations were introduced by primer-mediated site-directed mutagenesis. Chimeric constructs of Monosiga Yap and Amphimedon Yap with the human TBD were synthesized by GenScript. All GenScript constructs were cloned into the pUC57 plasmid EcoRV site.

For phiC31-mediated site-specific transformation, all constructs were cloned into the pUAST-att8 vector using BglII-NotI or NotI-XbaI sites. Nematostella warTs (Nwats) was amplified from larval cDNA and cloned into the pCRTM4-TOPO vector using the following primers: Nematostella warTF: 5'-TGGCCCCTACACTACCAAGGAGTAA G-3'; Nematostella warTR: 5'-AAGAATACTGCTCTGCTCGA CTGTTT-3'. We then cloned Nwats into NotI-digested pCaSpeR-hs.

Protein Stability of Yap/Yki Orthologs

To generate C-terminal human influenza hemagglutinin (HA) fusion proteins of Yap orthologs, the coding sequences of each protein were cloned in frame with the HA coding sequences in pHWH (Drosophila Genomics Resource Center) using Gateway cloning (Invitrogen). Yap-HA constructs were transfected into Drosophila S2 cells using Effectene Transfection Reagent Kit (Qiagen) following the manufacturer’s instructions. Transfected S2 cells were incubated for 24 h and heat shocked for 1 h to induce Yap-HA expression. Cell lysates were collected 3, 6, and 12 h postheat shock and analyzed by Western blotting with Anti-HA (Sigma-Aldrich,
Coimmunoprecipitation

We generated N-terminal HA and C-terminal Flag fusion proteins of TEAD and Yap, respectively. The full coding sequences of TEAD/Sd from Amphimedon, Nematostella, and Drosophila were amplified from their corresponding adult cDNAs and cloned in frame with the HA coding sequences in pAHW (Drosophila Genomics Resource Center) using the Gibson assembly kit (NEB). Using the same approach, we also cloned the coding sequences of the phosphomutant forms of Yap/Yki in frame with the coding sequence of Flag in pAWF (Drosophila Genomics Resource Center). Drosophila S2 cells were transiently transfected with these constructs as indicated above. After 3 days, cells were lysed (lysat buffer: 50 mM Tris pH7.4, 150 mM NaCl, 5 mM MgCl2, 5% glycerol, 0.5% Triton X-100, and 1× protease inhibitor cocktail [Roche]) and centrifuged at 16,000 × g for 10 min. Coimmunoprecipitations were performed using Dynabeads Protein G immunoprecipitation kit (Life Technologies) following the manufacturer’s instructions.

Drosophila Stocks

All transgenic flies carrying UAS-attB transgenes were created with phiC31-mediated site-specific transformation using the attP2 site (Groth et al. 2004; Bischof et al. 2007). These transgenes were overexpressed using either GMR-Gal4 or GMR-Gal4 with UAS-EGFP. The expression of diap1 was monitored using thmNC8 (diap1-lacZ) (Hay et al. 1995). To test the specificity of anti-Yki and anti-Sd antibodies in vivo, UAS-yki-RNAi (Bloomington, 34067) and UAS-sd-RNAi (Bloomington, 35481) were overexpressed using hsp-Gal4. For the Drosophila warts rescue experiment, we crossed wY, eyeless-FLP; FRT82B/TM6 Tb (Bloomington, 5620) to w, hs-Nvwarts/hs-Nvwarts; wts+/ FRT82B/TM6 Sb Tb. Expression of hs-Nvwarts was induced by heat shock for 1 h every day from the second instar larval stage until eclosion. wts +/+ is described in Xu et al. (1995).

Eye Width Measurement

For measuring adult eyes, flies were decapitated using a sharp razor blade. The heads were imaged using a Leica MZ 16F microscope. To determine eye width, two parallel lines were drawn at the edges of each eye. The distance separating these two lines was measured using ImageJ.

Edu Incorporation

EdU incorporation was detected using the Click-iT Alexa Fluor 488 imaging kit (Invitrogen). Third instar eye imaginal discs were incubated for 30 min with 300 μM EdU in Ringer’s solution. After fixation, samples were stored for at least 1 h in 100% methanol at −20 °C. All subsequent steps were conducted according to the manufacturer’s protocol.

Generation of Yki and Sd antibodies

Custom-made polyclonal rabbit anti-Yki and anti-Sd antibodies were generated and affinity purified by GenScript. They were raised against the N-terminal 243 amino acids of Yki and the full-length protein of Sd. For Western blots, anti-Yki and Anti-Sd dilutions were 1:2,000 and 1:4,000, respectively.

Immunocytochemistry

Imaginal discs and pupal retina were fixed and processed according to standard protocols. Primary antibodies used were anti-Cyclin E (Santa Cruz Biotechnology, 1:500), anti-β-galactosidase (Sigma-Aldrich, 1:1,000), anti-Armadillo (Developmental Studies Hybridoma Bank, 1:400), anti-Yki (1:1,000), and anti-Sd (1:1,000).

RNA Extraction, Library Preparation, Sequencing, and RNA-Seq Analysis

Total RNA was recovered from surgically isolated GMR expression domains (GFP+) from 25 to 30 third instar eye discs using a RNeasy kit (Qiagen). For each genotype, total RNA extraction was conducted in triplicate. Total RNA (400 ng) was enriched for poly(A)+ RNA by oligo(dt) selection. The Poly(A)+ RNA was then fragmented, and first-strand cDNA synthesis was performed using random hexamer priming in the Stowers Institute Molecular Biology Core facility, where all subsequent steps were conducted. Following second-strand synthesis, the ends were cleaned up, a nontemplated 3’ adenosine was added, and Illumina indexed adapters were ligated to the ends. The libraries were enriched by 15 rounds of polymerase chain reaction (PCR). The purified libraries were quantified with the high-sensitivity DNA assay on an Agilent 2100 Bioanalyzer. Equal molarities of individual libraries were pooled together (five libraries per pool) for multiplex sequencing. Pooled libraries were sent to Tufts (TUCF) for single read sequencing (50 nt) on a HiSeq 2000, and fastq files were returned.

For analysis, sequence reads in fastq format were mapped to the Drosophila genome using tophat-1.0.14 (Trapnell et al. 2009) against Flybase 5.29 (dm3 compatible). Flybase transcripts (v5.29) were quantified and compared between control and each overexpression condition using cuffdiff-1.0.2 (Trapnell et al. 2010). Genes with adjusted P values of less than 0.05 were selected for functional annotation based on Gene Ontology (GO Consortium 2006). Enrichment analysis was performed using Giotools (http://www.giotools.org, last accessed February 23, 2014) to identify biological processes that were enriched among up- or downregulated genes.

Quantitative Real-Time PCR

To independently validate the RNA-seq results, total RNA was isolated as described above. Five independent RNA extractions were performed for each genotype. Following cDNA generation, each real-time quantitative PCR (qPCR) reaction contained 0.33 ng of cDNA and a PCR master mix including 0.5 μM of each primer and 1× PerfeCTa SYBR Green FastMix from Quanta Biosciences (cat. no. 95072-250) in 10 μl total reaction using a CAS-4200 qPCR loading robot from Corbett.
Life Science. qPCR reactions were performed in 384-well format on a 7900HT Real-Time PCR Detection System from Applied Biosystems. Results were analyzed using qBase Plus software from Biogazelle. actin, GAPDH, and tbp genes were used as endogenous controls and the calibrated normalized relative quantity (CNRQ) values were calculated for each tested gene. Primers are available on request.

ChIP-Seq
Chromatin immunoprecipitation from eye discs was performed using a modified protocol from Gaertner et al. (2012). First, third instar larvae were dissected in phosphate buffered saline (PBS) (pH 7.4) such that only eye discs and brain remained attached to the mouth hooks. The dissected material was subsequently fixed in 1 ml fixation buffer (50 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid [HEPES], pH 7.5; 1 mM ethylenediaminetetraacetic acid [EDTA]; 0.5 mM ethylene glycol tetraacetic acid [EGTA]; 100 mM NaCl; 2% formaldehyde) for 30 min at room temperature. After four washes (PBS, pH 7.4; 0.1% Triton X-100; 0.1% Tween-20), eye discs were hand-dissected and combined into pools of 200 discs in 300 μl buffer A2 (15 mM HEPES, pH 7.5; 140 mM NaCl; 1 mM EDTA; 0.5 mM EGTA; 1% Triton X-100; 0.1% sodium deoxycholate; 0.1% sodium dodecyl sulfate [SDS]; 0.5% N-lauroylsarcosine; 1× Roche complete protease inhibitor cocktail, cat. no. 5056489001). Sonication was performed in a Bioruptor sonicator for 30 min (30 s on/off cycle at the “high” setting). Following centrifugation (16,000 × g; 10 min at 4°C), the supernatant containing soluble chromatin was transferred to fresh tubes, and 50 μl was set aside as whole cell extract (WCE; input).

Per ChIP, 10 μg antibodies were added to 450 μl chromatin (corresponding to approximately 300 discs) and incubated overnight at 4°C with rotation. We used the following antibodies: anti-Pol II (Covance 8WG16, cat. no. MMS-126R; mouse monoclonal antibody), anti-Sd, and anti-Yki. Immunocomplexes were purified by adding 50 μl prewashed Dynabeads coated with protein A/protein G (Life Technologies, cat. no. 10002D and 10004D) for 4 h, rotating at 4°C. The beads were washed three times in radioimmunoprecipitation assay buffer (RIPA) buffer (50 mM HEPES, pH 7.5; 1 mM EDTA; 0.7% sodium deoxycholate; 1% NP-40; 500 mM LiCl) and once in TE. Immunoprecipitated DNA was eluted twice in 75 μl elution buffer (50 mM Tris, pH 8.0; 10 mM EDTA; 1% SDS) at 65°C to maximize yields. Crosslinks of ChIP and WCE DNA were reversed overnight at 65°C. DNA was purified by RNase A (Sigma, cat. no. R6513; 0.2 μg/μl; 1 h at 37°C) and proteinase K (Life Technologies, cat. no. AM2546; 0.2 μg/μl; 2 h at 55°C) treatment followed by phenol/phenol–chloroform–isoamylalcohol extractions and ethanol precipitation. The precipitated DNA was resuspended in 30 μl 10 mM Tris buffer (pH 8).

ChIP-seq library preparation, 30 μl ChIP DNA and 100 ng WCE DNA were used to construct ChIP-Seq libraries with the NEBNext ChIP-Seq Library Prep Master Mix Set (cat. no. E6200L) and the NEBNext Multiplex Oligos (cat. no. E7335S and E7500S) for Illumina, following the manufacturer’s instructions. Concentration and size distribution of the libraries were assessed on an Agilent 2100 Bioanalyzer (high-sensitivity DNA assay chip). Sequencing was performed on an Illumina HiSeq2500 instrument, with 50 bp single reads in the high-output mode.

All sequence reads were filtered to include only those passing the standard Illumina quality filter and then aligned to the Drosophila melanogaster reference genome (UCSC dm3 release) using Bowtie version 0.12.9, with the following parameters: -v 2 -k 1 -m 3 --best --strata. Peaks were called with MACS v2.0.10.20120913 (Zhang, Liu, et al. 2008), using an adjusted P value of 0.001 and 0.01 for Yki and Sd, respectively. To assign a peak to its nearest gene, the following criteria were used: If the peak overlapped a gene, it was assigned to that gene regardless of where the overlap occurred; otherwise, it was assigned to the gene if the peak summit occurred within 1,500 bp upstream of the transcription start site.

ChIP-seq and RNA-seq data are available under Gene Expression Omnibus (GEO) accession number GSE54603.

Supplementary Material
Supplementary figures S1–S11 and tables S1–S4 are available at Molecular Biology and Evolution online (http://www.mbe. oxfordjournals.org/).

Acknowledgments
The authors thank B. Degnan, L. Grice, C. Conaco and K. Kosik for providing the Amphimedon cDNA; A. Peak and K. Zueckert-Gaudenz for RNA-seq library preparation and sequencing; K. Walton and A. Perera for ChIP-seq library sequencing; B. Fleharty and W. McDowell for real-time quantitative PCR analysis; J. Johnston for assistance with bioinformatic analysis; T. Akiyama for his suggestions on the co-immunoprecipitation experiments; and L. Gutchewsky for administrative support. Financial support was provided by the Stowers Institute for Medical Research and from a Burroughs Wellcome Fund Career Award in Biomedical Sciences to M.C.G.

References


Genevet A, Wehr MC, Brain R, Thompson BJ, Tapon N. 2010. Kibra is a pathway component TAZ.


OrthoDB: the hierarchical catalog of eukaryotic orthologs in 2011.


