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Eaf1 and Eaf2 negatively regulate canonical Wnt/β-catenin signaling

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SUMMARY

Eaf factors play a crucial role in tumor suppression and embryogenesis. To investigate the potential mechanism of Eaf activity, we performed loss- and gain-of-function assays in zebrafish using morpholino and mRNA injections, respectively. We found that eaf1 and eaf2 inhibit Wnt/β-catenin signaling, thereby modulating mesodermal and neural patterning in the embryo. Moreover, ectopic expression of eaf1 and eaf2 in embryos and cultured cells blocked β-catenin reporter activity. By immunoprecipitation, we also observed that Eaf1 and Eaf2 bound to the Armadillo repeat region and C-terminus of β -catenin, as well as to other β -catenin transcription complex proteins, such as c-Jun, Tcf and Axin, suggesting the formation of a novel complex. In addition, the N-terminus of Eaf1 and Eaf2 bound to β-catenin and exhibited dominant-negative activity, whereas the C-terminus appeared to either harbor a suppression domain or to recruit a repressor. Both the N- and C-terminus must be intact for Eaf1 and Eaf2 suppressive activity. Lastly, we demonstrate a conservation of biological activities for Eaf family proteins across species. In summary, our evidence points to a novel role for Eaf1 and Eaf2 in inhibiting canonical Wnt/β-catenin signaling, which might form the mechanistic basis for Eaf1 and Eaf2 tumor suppressor activity.

KEY WORDS: Eaf1, Eaf2, Wnt, β-catenin, Tumor suppressor

INTRODUCTION

Eaf1 and Eaf2 were originally identified as partners of ELL (elevennineteen lysine-rich leukemia), a fusion protein frequently associated with myeloid leukemia (Simone et al., 2001; Simone et al., 2003). Studies suggest that Eaf2 serves as a novel tumor suppressor, especially in prostate cancer (Xiao et al., 2003; Xiao et al., 2008; Xiao et al., 2009); however, the molecular mechanisms underlying this tumor suppression activity remain largely unknown.

The Wnt/β-catenin pathway plays crucial roles in embryonic development and tissue regeneration (Klaus and Birchmeier, 2008), and dysregulation of the pathway can result in cancer. Several groups have shown that inhibition of Wnt/β-catenin signaling constitutes a common mechanism for tumor suppressor activity (Morin et al., 1997; Sadot et al., 2001; Major et al., 2007; Meani et al., 2009). Furthermore, Wnt/β-catenin signaling participates in the formation of embryonic axes and neuroectoderm in vertebrates, and perturbation of this pathway often leads to defects in neuroectodermal and mesodermal patterning. Overexpression of chick Wnt8c in the mouse results in anterior neuroectoderm truncation (Pöpperl et al., 1997). Consistently, mice lacking functional Axin or Dkk1, both negative regulators of Wnt signaling, exhibit anterior neuroectoderm truncation (Zeng et al., 1997; Glinka et al., 1998; Mukhopadhyay et al., 2001). In zebrafish, enhanced Wnt signaling also leads to loss of rostral neural domains. Forebrain defects and small eyes are observed in two zebrafish mutants: masterblind (mbl), which carries a mutation in axin1 with high Wnt/β-catenin activities (Heisenberg et al., 2001), and headless (hdl), which carries a mutation abolishing the repressor function of tcf3 (tcf7l1a – Zebrafish Information Network) on Wnt/β-catenin signaling (Kim et al., 2000). By contrast, zebrafish wnt8 mutants and morphants display significant expansion of forebrain markers (Lekven et al., 2001).

In mesoderm patterning, maternal Wnt/β-catenin signaling is essential for the induction of sqt (ndr1), boz (dharma) and gsc during Nieuwkoop center formation, which then induces formation of the dorsal mesodermal organizer. After the mid-blastula transition, zygotic Wnt/β-catenin signaling is activated by Wnt ligands to antagonize the organizer, and is involved in ventralposterior mesoderm and neuroectoderm induction by directly regulating ventral mesoderm tbx6 (Szeto and Kimelman, 2004) and posterior ectoderm/mesoderm cdx4, respectively (Pilon et al., 2006). Zebrafish wnt8 mutants show a significant expansion of the organizer and almost absent expression of ventrolateral mesoderm markers, similar to the phenotypes observed in *wnt8* morphants (Lekven et al., 2001).

Multiple molecules at several different points in the Wnt/β-catenin signaling pathway can modulate the output of the pathway. Extracellular proteins, including members of the Frzb or Dickkopf families, inhibit Wnt signaling by competitively binding to Wnt ligands (Leyns et al., 1997) or the LDL receptor-related protein 5/6 (Mao et al., 2001; Mao et al., 2002), respectively. By contrast, BIO, a bona fide drug that inhibits Gsk3, enhances Wnt/β-catenin signaling (Notani et al., 2010). In the cytoplasm, several proteins, including Gsk3, Apc and Axin, form a destruction complex that phosphorylates β-catenin, leading to its degradation (Barker and Clevers, 2000). As such, apc mutant fish have constitutively active Wnt/β-catenin signaling (Hurlstone et al., 2003; Haramis et al., 2006; North et al., 2007; Goessling et al., 2008). In the nucleus, tcf3 is essential for forebrain formation by repressing Wnt caudal target genes (Kim et al., 2000). The dominant-negative form of Tcf3, dn-Tcf, which lacks the β-catenin-binding domain and acts as a repressor, can efficiently promote anterior brain formation (Kim et al., 2000).

Given the potential association between cancer and development, we have explored eaf1 and eaf2 activity in vertebrate

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embryogenesis, using the zebrafish model. Our previous studies demonstrate that both *eaf1* and *eaf2* mediate effective convergence and extension movements through maintaining expression of the non-canonical Wnt/β-catenin signaling ligands *wnt5* and *wnt11* (Liu et al., 2009). In addition, *eaf1* and *eaf2* regulate the expression of another non-canonical Wnt/β-catenin signaling ligand, *wnt4*, in a negative-feedback loop (Wan et al., 2010). We also observed that embryos with morpholino-mediated *eaf1* and *eaf2* knockdown display forebrain defects (Liu et al., 2009; Wan et al., 2010). Because upregulated Wnt/β-catenin signaling leads to loss of rostral neural domains of the forebrain in zebrafish (Kim et al., 2002), we investigated whether *eaf1* and *eaf2* also modulate canonical Wnt/β-catenin signaling.

In this study, we used zebrafish as an *in vivo* model together with cell culture to investigate the role of Eaf1 and Eaf2 in the regulation of Wnt/ β -catenin signaling. Our results not only reveal a novel function of Eaf1 and Eaf2 during embryogenesis but also suggest a potential mechanism of Eaf1 and Eaf2 in tumor suppression.

MATERIALS AND METHODS

Fish stocks

Wild-type (AB line) zebrafish (*Danio rerio*) maintenance, breeding and staging were performed as described previously (Liu et al., 2009). *apc*^{mer} heterozygous mutants were kindly provided by Dr Wolfram Goessling (Brigham and Women's Hospital, Dana-Farber Cancer Institute). Genotyping for *apc* mutants was performed by sequencing the mutant allele fragment amplified from genomic DNA extracted from embryos stained by whole-mount *in situ* hybridization for *opl* (*zic1* – Zebrafish Information Network) (Hurlstone et al., 2003). Primers are listed in supplementary material Table S1. The *hs:dkk-GFP* transgenic line was kindly provided by Dr Leonard Zon (Harvard University) (North et al., 2007).

Drug exposure

Embryos developed to 50% epiboly were exposed to BIO (Sigma-Aldrich) at 5 μ M.

Heat-shock modulation

Embryonic heat-shock experiments were conducted at 38°C for 20 minutes. Genotype was determined by the presence of GFP fluorescence at 3 hours post-heat shock; the non-fluorescent (wild-type) siblings were sorted and used as controls.

Morpholino (MO) and mRNA synthesis

The translation-blocking MOs (ATG targeted) Eaf1-MO1 and Eaf2-MO1 have been described previously (Liu et al., 2009). The splicing MOs Eaf1-MO3 and Eaf2-MO3 were designed based on the junction sequence between exon 1 and intron 1 of *eaf1* and *eaf2*, respectively. Eaf1-MO3 and Eaf2-MO3 specificity was validated by RT-PCR (primers listed in supplementary material Table S1). β -catenin 1 and β -catenin 2 antisense MOs (β -catenin1-MO and β -catenin2-MO) have been described previously (Bellipanni et al., 2006). All MOs were purchased from Gene Tools and their sequences are listed in supplementary material Table S2.

Capped mRNAs were synthesized using the Ampticap SP6 High Yield Message Maker Kit (Epicenter Biotechnologies). The plasmids for zebrafish eaf1 and eaf2 and human EAF1 and EAF2 mRNA were described previously (Liu et al., 2009). The full-length genes, exons 1-3 and exons 4-6 of zebrafish eaf1 and eaf2 were subcloned into PSC2+VP16 and PSC2+EnR (provided by Yonghua Sun, Institute of Hydrobiology, CAS, Wuhan, China) to form fusion protein expression vectors. In addition, exons 1-3 and exons 4-6 of zebrafish eaf1 and eaf2 were subcloned into PSP64 (Promega) for synthesizing mRNAs. The synthesized mRNAs were diluted to different concentrations and injected into one-cell stage embryos.

Whole-mount in situ hybridization

Probes for zebrafish six3b, opl and cdx4 were amplified from cDNA pools using the appropriate sets of primers (supplementary material Table S1). Probes for tbx6 were a generous gift from Dr Yang Wang (Institute of

Hydrobiology, Chinese Academy of Sciences, Wuhan, China). Probes for *eaf1*, *eaf2*, *gsc* and *chd* have been described previously (Liu et al., 2009).

Plasmid construction

The full-length zebrafish β -catenin 2 (ctnnb2) vector was a generous gift from Dr Gianfranco Bellipanni (Temple University, Philadelphia, USA). Full-length zebrafish β -catenin 1 (ctnnb1) and c-jun were amplified from cDNA pools using the appropriate sets of primers (supplementary material Table S1). Zebrafish ctnnb1, ctnnb2, c-jun, eaf1 and eaf2 were subcloned into pCGN-HAM (provided by William Tansey, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY, USA), pCMV-Myc, pEGFP-N1, PM-RFP or PM vectors (Clontech). Sequences encoding the N-terminus, Armadillo repeats and C-terminus of zebrafish ctnnb1 and ctnnb2 were subcloned into pCGN-HAM, pCMV-Myc and PM vectors. Sequences encoding the N-terminus (exons 1-3) and C-terminus (exons 4-6) of zebrafish eaf1 and eaf2 were subcloned into pCMV-Myc. All plasmids were verified by sequencing. Human Myc-EAF1 and Myc-EAF2 were described previously (Zhou et al., 2009). Human HA-AXIN and HA-c-JUN were generous gifts from Dr Lin Li (Institute of Biochemistry and Cell Biology, Chinese Academy of Science, Shanghai, China). Full-length human βcatenin and TCF4 (TCF7L2 – Human Gene Nomenclature Committee) were amplified from cDNA pools and subcloned into pCGN-HAM.

Luciferase reporter assay

Human embryonic kidney (HEK) 293 cells were cultured in Dulbecco's modified Eagle's medium (DMEM) containing 10% fetal bovine serum (HyClone). The luciferase reporter assay was performed as described previously (Zhou et al., 2009). The reporter vectors 8xTopFlash and FopFlash were generous gifts from Dr Randall Moon (University of Washington, Seattle, USA). The pER-luc vector was purchased from Stratagene. A c-Myc promoter luciferase reporter was amplified by PCR based on the human *c-MYC* promoter region described previously (Chung et al., 1986) and cloned into pGL3-Basic (Promega). Statistical analysis of luciferase assay data was performed using GraphPad Prism 5.0.

Fluorescence microscopy

Cos-7 cells were transfected with different combinations of zebrafish plasmids. After 24-30 hours, cells were directly observed under a Nikon T-2000 Eclipse inverted fluorescence microscope.

Western blots and immunoprecipitation assays

Western blots were performed as described (Zhou et al., 2009). HEK 293T cells and mouse L cells (with *Wnt3a* expression) were cultured in DMEM. Immunoprecipitation assays were performed as described (Xiao et al., 2009). The following antibodies were used in the assays: anti-Myc (Santa Cruz), anti-HA (Covance), anti-β-catenin (Santa Cruz), anti-β-catenin-ABC (Santa Cruz), anti-GAPDH (Abcam), anti-PCNA (Epitomics), anti-α-tubulin (Epitomics) and anti-H3 (Cell Signaling Technology).

Nuclear and cytoplasmic separation

Cell fractionation followed the method of Suzuki et al. (Suzuki et al., 2010) with the following modifications. Briefly, the embryos were collected in 24-well plates, Pronase E was added (1 mg/ml; Solarbio, Beijing, China) and incubated for 15-30 minutes at 37°C. The chorion was discarded and the embryos washed three times with PBS, then transferred to 1.5-ml Eppendorf tubes, 1 ml PBS added, and the embryos disrupted using a micropipette tip. The embryos were then spun for 5 minutes at 1100 rpm to separate the yolks. Embryos were collected at the gastrula stage and HEK 293T cells were harvested 18-26 hours after transfection.

RESULTS

Patterning of anterior neuroectoderm and mesoderm in zebrafish embryos requires *eaf1* and *eaf2*

In our previous studies, we observed ubiquitous expression of zebrafish *eaf1* and *eaf2* during early embryogenesis (supplementary material Fig. S1) (Liu et al., 2009). Furthermore, *eaf1* and *eaf2* morphants displayed forebrain defects (Liu et al., 2009). In addition

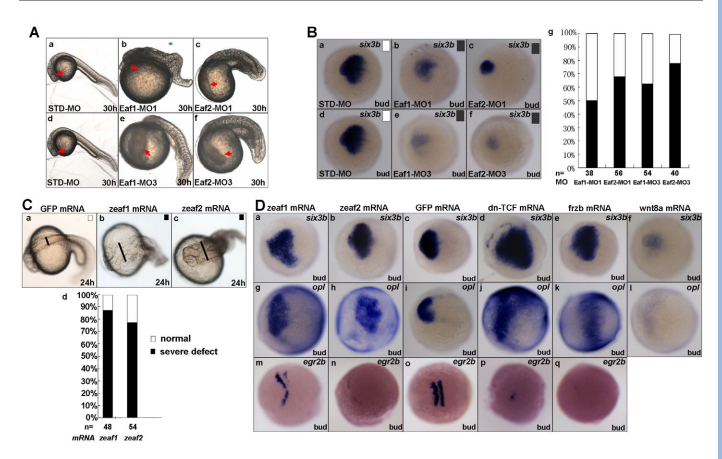


Fig. 1. Effects of eaf1 and eaf2 knockdown and overexpression on zebrafish embryos. (A) Morphology of representative embryos injected with the indicated MOs (8 ng/embryo) at 30 hpf. STD-MO, standard MO. (B) (a-f) In situ hybridization for the anterior brain marker six3b in the presence of the indicated MOs. (q) Scoring of six3b expression: black, reduced expression; white, normal expression. (C) (a-c) Morphology of representative embryos injected with the indicated mRNAs (200-500 pg/embryo), zeaf1 and zeaf2 refer to zebrafish eaf1 and eaf2. (d) Morphology was scored as normal (white) or severely defective (black) in eaf1- and eaf2-overexpression embryos. The expanded dorsal part is indicated by the black line. (D) eaf1 and eaf2 function as forebrain inducers in embryos. Embryos injected with eaf1 and eaf2 mRNA displayed increased expression of six3b (a,b) and opl (g,h), similar to embryos injected with dn-Tcf (d,j) or frzb (e,k) mRNA, but in contrast to embryos injected with wnt8a mRNA (f,l); egr2b exhibited reduced expression in embryos injected with eaf1 and eaf2 mRNA (m,n), similar to embryos injected with dn-Tcf (p) or frzb (q) mRNA. Ba-f,Da-q, dorsal views, anterior to the left.

to the translation-blocking MOs, Eaf1-MO1 and Eaf2-MO1 (Liu et al., 2009), we included two splicing-blocking MOs, Eaf1-MO3 and Eaf2-MO3. The efficiency and specificity of the splicing-blocking MOs were confirmed by RT-PCR assays (supplementary material Fig. S2). Knockdown of eaf1 or eaf2 resulted in forebrain defects in embryos (Fig. 1Ab,c,e,f), consistent with our previous observations (Liu et al., 2009).

In order to further characterize the observed phenotypes, we evaluated the expression of the anterior neuroectoderm markers six3b, foxg1 and opl, as well as the hindbrain marker egr2b, as a means to assess defects in neuroectoderm anterior-posterior patterning. We also evaluated the dorsal mesoderm markers sqt, gsc, dkk1, frzb, the posterior ectoderm/mesoderm marker cdx4, and the ventral mesoderm marker tbx6, to assess defects in dorsal-ventral

In the Eaf morphants at the bud stage, expression of six3b was dramatically reduced (Fig. 1Ba-f) and expression of foxg1 was almost completely abolished (supplementary material Fig. S3A). Moreover, quantitative analysis indicated that at least half of the embryos with Eaf knockdown had reduced six3b expression (Fig. 1Bg). We also detected defects in dorsal-ventral mesoderm patterning as indicated by enhanced expression of sqt and gsc, the main direct targets of maternal Wnt/β-catenin signaling (Kelly et al., 1995; Shimizu et al., 2000), at the blastula stage (supplementary material Fig. S3B-D), as well as increased expression of cdx4 and tbx6, two zygotic Wnt direct targets (Szeto and Kimelman, 2004; Pilon et al., 2006), at the gastrula stage (supplementary material Fig. S3E-H). The observations suggested that Wnt/β-catenin activities increased in Eaf morphants.

As expected, overexpression of eaf1 and eaf2 by mRNA injection (200-500 pg/embryo) caused phenotypes opposite to those of Eaf morphants. Of the embryos injected with zebrafish eafl or eaf2 mRNA (Fig. 1Cb,c), 75-85% displayed obvious dorsalized phenotypes with expansion and widening of the dorsal part (Fig. 1C, black lines). Similarly, overexpression of eaf1 or eaf2 produced opposite effects on marker gene levels to the morphants. The expression of six3b and opl increased dramatically in embryos with ectopic expression of eaf1 or eaf2 (Fig. 1Da,b,g,h). These resembled embryos with ectopic expression of dn-Tcf (Fig. 1Dd,j) or frzb (Fig. 1De,k) (Kim et al., 2002; Momoi et al., 2003), but not wnt8a (Fig. 1Df,l) (Kelly et al., 1995; Kim et al., 2002). Injection with eaf1 or eaf2 mRNA reduced the expression of egr2b in most embryos (Fig. 1Dm,n), similar to ectopic expression of dn-*Tcf* (Fig. 1Dp) or frzb (Fig. 1Dq). eaf1 or eaf2 mRNA also significantly increased the expression of sqt and gsc, as revealed by both whole-mount in situ hybridization and RT-PCR analysis at the gastrula stage

(supplementary material Fig. S4A), again similar to embryos with ectopic expression of dn-*Tcf* (supplementary material Fig. S4Bb) or *frzb* (supplementary material Fig. S4Bc).

eaf1 and eaf2 function in anterior neuroectoderm and mesoderm patterning by antagonizing Wnt/β-catenin signaling

It has been reported that hyperactive Wnt/ β -catenin signaling suppresses the rostral neural domains of the forebrain in zebrafish by promoting the posterior midbrain and hindbrain domains (Kim et al., 2002). Here, we observed that expression of the direct Wnt targets sqt, gsc, cdx4 and tbx6 increased in Eaf morphants at blastula and gastrula stages (supplementary material Fig. S3). To confirm these results, we examined the expression of axin2, a bona fide direct target of Wnt/ β -catenin signaling at bud stage. As expected, we observed an obvious increase in axin2 expression in the midbrain of Eaf morphants at bud stage (supplementary material Fig. S5A, red arrows). These observations suggested that Wnt/ β -catenin signaling was upregulated in Eaf morphants during early embryogenesis.

The above results, which were based on detecting the expression of Wnt signaling direct target genes through loss- and gain-of-function assays, implied that eaf1 and eaf2 might serve as antagonists of Wnt/ β -catenin signaling to affect neuroectoderm and mesoderm patterning. If this were the case, then ectopic expression of eaf1 or eaf2 should counteract the effects of enhanced Wnt/ β -catenin signaling induced by ectopic expression of wnt8a. To test

this, we performed rescue experiments by monitoring *opl* expression. As expected, co-injection of *eaf1* or *eaf2* mRNA with *wnt8a* mRNA rescued *opl* expression dramatically (Fig. 2A; scored based on the expression level of *opl*). In addition, *eaf1* or *eaf2* also partially counteracted the strong inhibitory effect of BIO on *opl* expression (Fig. 2B).

We also performed rescue experiments using apc mutant zebrafish embryos, which have enhanced Wnt/β-catenin activity. As expected, in the offspring of $apc^{+/-} \times apc^{+/-}$, the ratio of strongly reduced to reduced to normal opl was close to 1:2:1, correlating with the predicted genotype ratio (Fig. 2Cb, column 1 from top to bottom). Further genotyping of these offspring not only confirmed the genotype ratio of 1:2:1 (data not shown) but also showed that the phenotypes (indicated by opl expression) mirrored the genotypes very well (supplementary material Fig. S5B). After injection of eaf1 or eaf2 mRNA, wild-type embryos $(apc^{+/+})$ exhibited increased opl expression, whereas the heterozygous embryos $(apc^{+/-})$ exhibited normal or reduced opl, and most of the homozygous embryos $(apc^{-/-})$ exhibited reduced *opl* (Fig. 2C). Therefore, ectopic expression of eaf1 or eaf2 could increase opl levels in embryos of all three genetic backgrounds (Fig. 2Cb). These results further suggest that eaf1 and eaf2 are involved in Wnt/ β -catenin signaling and might function downstream of apc.

If eaf1 and eaf2 truly act as antagonists of Wnt/ β -catenin signaling, ectopic expression of Wnt/ β -catenin signaling inhibitors or knockdown of β -catenin 1 and 2 should counteract the effects seen with Eaf knockdown. To test this, we performed rescue

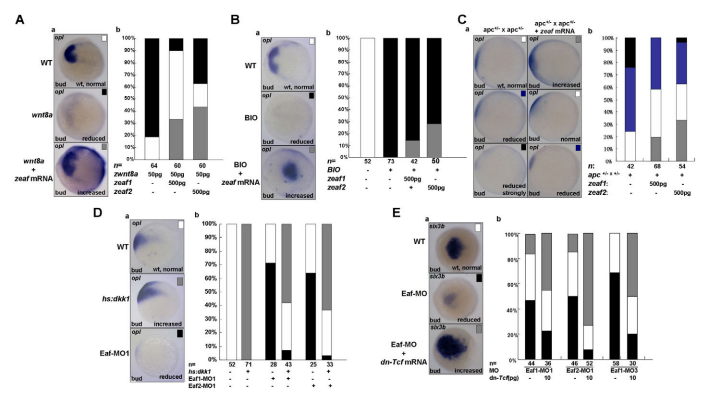


Fig. 2. Eafs antagonize Wnt/β-catenin signaling in anterior neuroectoderm patterning. (A) *eaf1* and *eaf2* counteract *wnt8a*-mediated forebrain defects. (a) *In situ* hybridization for the anterior neuroectoderm marker *opl* in wild type (WT) and in the presence of *wnt8a* mRNA alone or together with *eaf1* or *eaf2* mRNA. (b) Scoring of *opl* expression: black, reduced; white, normal; gray, expanded. (**B**) *eaf1* or *eaf2* partially counteracts the strong reduction in *opl* expression that occurs after exposure to the Gsk3 inhibitor BlO, which enhances Wnt/β-catenin signaling. *opl* expression is scored as in A. (**C**) *eaf1* and *eaf2* rescue forebrain defects in *apc* mutants. *opl* expression is scored as: black, strongly reduced; blue, reduced; white, normal; gray, increased. (**D**) Heat shock-induced expression of *dkk1* at 50% epiboly rescues forebrain defects in Eaf morphants. *opl* expression is scored as in A. (**E**) dn-*Tcf* partially rescues the anterior brain defects in Eaf morphants. Embryos were scored by *six3b* expression: black, reduced; white, normal; gray, increased. Aa,Ba,Ea, dorsal views, anterior to the left; Ca,Da, lateral views, anterior to the left.

experiments using hs:dkk-GFP transgenic zebrafish. Heat shock of hs:dkk-GFP embryos at 50% epiboly resulted in all of the embryos displaying enhanced opl expression (Fig. 2Db). In Eaf morphants at 50% epiboly, dkk1 induced by heat shock efficiently rescued the reduced *opl* expression seen with Eaf knockdown (Fig. 2D). Some morphants even displayed enhanced opl expression (Fig. 2Db), suggesting that dkk1 might be more efficient than Eaf in inhibiting Wnt/β-catenin signaling. Similarly, co-injection of dn-*Tcf* mRNA effectively restored six3b expression in Eaf morphants (Fig. 2E). Subsequently, we evaluated morphants after simultaneous knockdown of β -catenin 1/2 and eaf1/2. Co-injection of β -catenin1-MO with the Eaf MOs effectively restored six3b expression (supplementary material Fig. S5C). Surprisingly, co-injection of βcatenin2-MO with the Eaf MOs resulted in a further reduction in six3b expression compared with embryos injected with the Eaf MOs alone (supplementary material Fig. S5C).

For mesoderm patterning, only β-catenin2-MO rescued gsc expression in Eaf1-MO morphants at 30% epiboly (supplementary material Fig. S5Da-h), whereas both β-catenin1-MO and βcatenin2-MO partially rescued gsc expression in Eaf1-MO morphants at the sphere stage (supplementary material Fig. S5Dim), although β-catenin2-MO was more effective (supplementary material Fig. S5Dm).

To gain a more complete picture of eaf1 and eaf2 in Wnt/βcatenin signaling, we also determined whether β-catenin could modulate eaf1/2 expression. Embryos injected with the active form of β-catenin mRNA displayed reduced expression of eaf1 and eaf2 (supplementary material Fig. S6). Similarly, embryos with ectopic wnt8a also displayed a decrease in eaf1 and eaf2 expression (supplementary material Fig. S6Ac,g). However, the expression of eaf1 and eaf2 remained unchanged in embryos injected with frzb mRNA (supplementary material Fig. S6Ad,h), implying a complicated regulatory relationship between eaf1/2 and the Wnt/βcatenin signaling pathway.

Eaf1 and Eaf2 act as transcriptional repressors in patterning anterior neuroectoderm and mesoderm

Previous studies showed that Eaf1 and Eaf2 contain a transactivation domain in the C-terminus and that Eaf2 binds to specific nucleotides (Simone et al., 2001; Xiao et al., 2003; Xiao et al., 2006). These observations suggest that Eaf1 and Eaf2 might act as transcription factors. To explore this possibility, we made constructs by fusing full-length eaf1 or eaf2 with the transcriptional activator VP16 or with the transcriptional repressor engrailed (EnR) (Gómez-Skarmeta et al., 2001; Muhr et al., 2001). We also made constructs by fusing the N-terminal sequences (encoding amino acids 1-112) of eaf1 or eaf2 (eaf1-N or eaf2-N) or their C-terminal sequences (encoding amino acids 113-256 for eaf1-C and 113-259 for eaf2-C) with VP16 or EnR for mapping their functional domains (Fig. 3Aa).

Embryos injected with EnR mRNA or VP16 mRNA displayed the same phenotype as wild-type embryos (supplementary material Fig. S7A; data not shown), indicating that VP16 or EnR alone was not functional in vivo, probably owing to the lack of specific DNAor protein-binding domains. Most embryos injected with eaf1/2-EnR or eaf1/2-N-EnR mRNA (50 pg/embryo) displayed obviously dorsalized phenotypes (Fig. 3Ac-f; data not shown), similar to embryos injected with full-length eaf1 or eaf2 mRNA (Fig. 1C). By contrast, most embryos injected with eaf1/2-VP16 or eaf1/2-N-VP16 mRNA (50 pg/embryo) displayed phenotypes with obvious anterior neuroectoderm truncation (Fig. 3Ag-j; data not shown),

similar to Eaf morphants (Fig. 1A). However, embryos injected with either eaf1/2-C-EnR mRNA or eaf1/2-C-VP16 mRNA (50 pg/embryo) did not exhibit any abnormal phenotypes (supplementary material Fig. S7B; data not shown), similar to embryos injected with either EnR or VP16 mRNA alone (supplementary material Fig. S7A), further ruling out any nonspecific effect of EnR or VP16.

In addition, embryos injected with eaf1/2-EnR or eaf1/2-N-EnR mRNA exhibited expanded six3b and opl expression (Fig. 3Ba-j) and reduced egr2b expression (Fig. 3Bk-o), similar to embryos injected with full-length eaf1/2 mRNA (Fig. 1D). Embryos injected with eaf1/2-EnR or eaf1/2-N-EnR mRNA displayed enhanced expression of the dorsal-ventral patterning markers gsc (supplementary material Fig. S7Ca-e) and chd (supplementary material Fig. S7Cf-j), similar to embryos injected with full-length eaf1 and eaf2 mRNA (supplementary material Fig. S4Ad-f). Taken together, these phenotypes suggest that eaf1 and eaf2 might modulate embryogenesis by acting as transcriptional repressors rather than activators in vivo.

Interestingly, the phenotypes exhibited by embryos injected with eaf1/2-N or with full-length eaf1/2 fused with the EnR transcriptional repressor were similar to those of embryos injected with full-length eaf1/2 mRNA. This promoted us to further examine the function of the N- and C-termini of Eaf1 and Eaf2.

Both the N- and C-termini must be intact for Eaf1 and Eaf2 suppressive activity

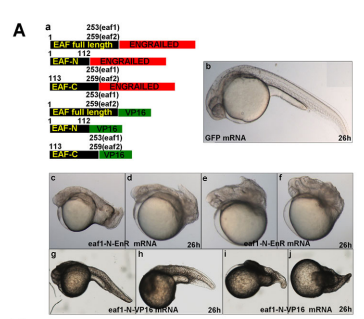
We next investigated the function of the N- and C-termini of Eafl and Eaf2 by ectopic expression in embryos. Embryos injected with eaf1-N or eaf2-N mRNA (50 pg/embryo) displayed remarkable anterior neural truncation (nearly 90%; Fig. 4A), in contrast to embryos injected with full-length eaf1 or eaf2 mRNA (Fig. 1C). Embryos injected with eaf1-C or eaf2-C mRNA (50 pg/embryo) displayed no obvious defects (data not shown). Furthermore, six3b expression was reduced dramatically and that of opl was almost completely abolished in embryos injected with eafl-N or eaf2-N mRNA (Fig. 4B; data not shown), whereas their expression was unchanged in embryos with ectopic expression of eaf1-C or eaf2-C (Fig. 4C; data not shown). After co-injecting embryos with fulllength eaf1 or eaf2 mRNA and eaf1-N or eaf2-N mRNA, the reduced expression of six3b was rescued (Fig. 4Da,b). In addition, co-injection of eaf1-N-EnR with eaf1-N mRNA also restored six3b expression (Fig. 4Db), which further reinforced the suppressor role of Eaf1.

Next, we determined whether the N-terminus of Eaf1/2 functions as a dominant-negative form to affect neural and mesodermal patterning by upregulating Wnt/β-catenin signaling. Co-injection of dn-Tcf mRNA counteracted the reduced six3b expression that resulted from ectopic expression of eaf1-N or eaf2-N (Fig. 4D). In addition, co-injection of β -catenin1-MO, but not β -catenin2-MO, effectively rescued six3b expression in embryos injected with eaf1-N or *eaf2*-N mRNA alone (Fig. 4Dc).

Taken together, these observations suggest that the C-terminus of Eaf1/2 might function as repressor or as a repressor recruiter and that both the N-terminus and the C-terminus must remain intact for Eaf1 and Eaf2 to function as suppressors in Wnt/β-catenin signaling.

Eaf1 and Eaf2 attenuate the transcriptional activity of β-catenin

To determine the mechanisms underling eaf1 and eaf2 antagonism of Wnt/β-catenin signaling, we used TopFlash luciferase reporter assays to monitor the transcriptional activity of β-catenin in



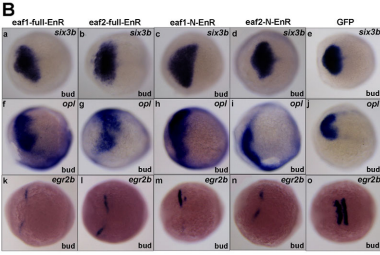


Fig. 3. Zebrafish *eaf1* and *eaf2* induce anterior neuroectoderm by acting as transcriptional repressors.

(A) (a) mRNAs used for injection encode full-length or the Nor C-terminus of Eaf1 or Eaf2 fused with the transcriptional activator VP16 or the engrailed transcriptional repressor domain (EnR). Numbers refer to amino acid residues in Eaf1 and Eaf2. (b) Embryos injected with GFP mRNA at 26 hpf exhibit normal morphology. (c-f) Embryos injected with eaf1-N-EnR mRNA display similar morphology to embryos injected with full-length eaf1 mRNA. (g-j) Embryos injected with eaf1-N-VP16 mRNA display phenotypes with obvious anterior neuroectoderm truncation, similar to Eaf morphants. (B) eaf1 and eaf2 act as anterior neuroectoderm inducers by repressing transcription. six3b (a-e) and opl (f-j) displayed expanded expression in embryos injected with eaf1/2-EnR or eaf1/2-N-EnR mRNA. (k-o) egr2b expression was reduced in embryos injected with eaf1/2-EnR or eaf1/2-N-EnR mRNA. Bao, dorsal views, anterior to the left. All mRNA injections were at 50 pg/embryo.

zebrafish embryos and cell lines (Aoki et al., 1999; Playford et al., 2000). First, we injected Eaf1-MO1 alone or together with the 8xTopFlash reporter and an internal control vector, pTK-renilla, and then performed luciferase assays. At 5 hours post-fertilization (hpf), embryos injected with Eaf MOs exhibited obviously enhanced reporter activity as compared with control embryos injected with a standard MO (Fig. 5A). Western blots revealed that injection of Eaf MOs did not change the protein levels of either total β -catenin or active β-catenin (supplementary material Fig. S8A). In addition, to determine whether ectopic expression of eaf1/2 mRNA in embryos could directly suppress the transactivity of β-catenin in vivo, we injected embryos at the one-cell stage with expression plasmids of Myc-tagged zebrafish Eaf1 (Myc-zeaf1), Eaf2 (Myc-zeaf2) or empty vector (control) together with the 8xTopFlash reporter as well as pTK-renilla. At a series of time points after injection (6, 8 and 10 hours), Myc-zeafl or Myc-zeaf2 dramatically suppressed 8xTopFlash activity as compared with the control vector (Fig. 5B), in contrast to what was observed in embryos with Eaf1/2 knockdown.

To further determine the role of eaf1 and eaf2 in Wnt/ β -catenin signaling, we also evaluated the effect of ectopic Eaf expression on β -catenin transcriptional activity in mammalian cell lines. In HEK 293 cells (Fig. 5Ca,b), zebrafish eaf1 and eaf2 dramatically

suppressed 8xTopFlash reporter activity induced by zebrafish β -catenin 1 or β -catenin 2. In addition, we found that eaf1 and eaf2 suppressed the endogenous β -catenin activity induced by conditioned medium containing Wnt3a (Fig. 5Cc). Again, zebrafish eaf1 or eaf2 did not affect the protein level of either β -catenin 1 or β -catenin 2 (supplementary material Fig. S8B).

Given that *c-Myc* represents a bona fide target of Wnt/β-catenin signaling (He et al., 1998), and that *c-Myc* is a well-defined proto-oncogene that is often upregulated in many types of cancers, we next evaluated whether *eaf1* or *eaf2* could also suppress *c-Myc* expression. Indeed, zebrafish *eaf1* or *eaf2* (Fig. 5Da,b) significantly suppressed the activity of a human *c-MYC* promoter in a dose-dependent manner. Western blots confirmed the expression of ectopic zebrafish Eaf1 and Eaf2 proteins (supplementary material Fig. S8C). Using semi-quantitative RT-PCR, we detected decreased expression of endogenous *c-MYC* mRNA 20 hours after transfection with zebrafish *eaf1* or *eaf2* (Fig. 5Dc), suggesting that they might suppress *c-Myc* expression through modulating Wnt/β-catenin signaling.

Eaf1 and Eaf2 interact with β-catenin 1 and β-catenin 2 to form a protein complex

Using co-immunoprecipitation assays, we found that zebrafish Eaf1 and Eaf2 strongly interact with zebrafish β -catenin 1 and β -catenin

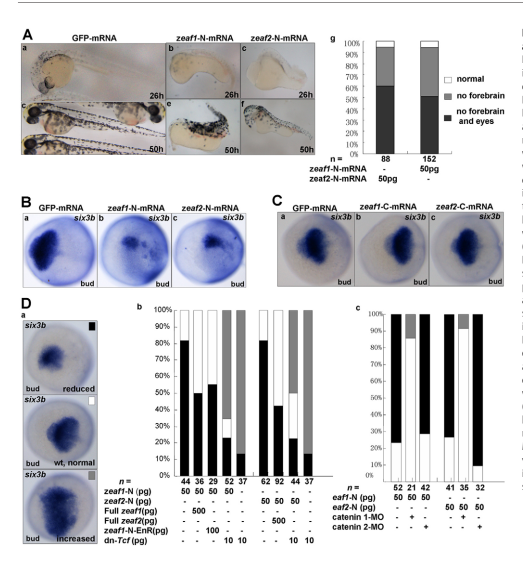


Fig. 4. The N-terminus of Eaf1 or Eaf2 acts as a dominant negative. (A) (a-f) Representative morphology of embryos injected with GFP mRNA or mRNA encoding the N-terminus of Eaf1 or Eaf2 (exons 1-3) at 26 or 50 hpf. (g) Embryos were scored morphologically at 50 hpf: white, the percentage of normal embryos; gray, embryos without forebrain; black, embryos without forebrain and eyes. (B,C) six3b expression was reduced in embryos injected with mRNA encoding the Nterminus (exons 1-3) of Eaf1 or Eaf2 (B) but was normal in embryos injected with mRNA encoding the C-terminus (exons 4-6) of Eaf1 or Eaf2 (C). (D) The N-terminus (exons 1-3) of Eaf1/2 suppresses forebrain formation by promoting enhanced Wnt/β-catenin activity. (a) Representative embryos showing normal, reduced and increased expression of six3b. (b) Embryos were scored for six3b expression after injection with mRNAs at the dosages indicated. (c) Abnormal expression of six3b in embryos injected with mRNA encoding the N-terminus (exons 1-3) of Eaf1 or Eaf2 was rescued by co-injection with β-catenin1-MO (8 ng/embryo) but not with β-catenin2-MO (8 ng/embryo). Ba-c,Ca-c,Da, dorsal views, anterior to the left. mRNA injections were at 50 pg/embryo unless stated otherwise (Db).

2 (Fig. 6B). Domain mapping indicated that Eaf1 and Eaf2 interact with the β -catenin center region that contains 12 Armadillo repeats and the C-terminus (Fig. 6D,E), but not the N-terminus (Fig. 6C). This might explain why eaf1 and eaf2 could also suppress the transactivity of the C-terminus as well as full-length β-catenin 1 and β-catenin 2. Furthermore, domain mapping for Eaf1 and Eaf2 revealed that the N-terminal region (amino acids 1-112; Fig. 6F-H), but not the C-terminal region (amino acids 113-253 for Eaf1 and 113-259 for Eaf2; Fig. 6I), bound to β -catenin 1/2. In addition, zebrafish Eaf1 and Eaf2 also interacted with c-Jun (Fig. 6J) (Gan et al., 2008).

To confirm the interaction between Eaf1/2, β-catenin 1/2 and c-Jun, we performed colocalization assays. Eaf1 and Eaf2 colocalized with β -catenin 1/2 and c-Jun in Cos-7 cells after transfection with GFP- or RFP-tagged proteins (supplementary material Fig. S9).

To determine whether *eaf1* or *eaf2* also suppresses β-catenin transactivity by altering the cytoplasmic-nuclear shuttling of βcatenin, we measured β-catenin levels in the cytoplasm and nucleus by western blot analysis after ectopic expression or knockdown of eaf1/2. Zebrafish β-catenin was found mainly in the nucleus, and altering eaf1/2 expression did not affect the distribution of the protein (supplementary material Fig. S10). Thus, eaf1 and eaf2 do not affect the translocation of β -catenin between the cytoplasm and nucleus.

The suppressive effect of Eaf1 and Eaf2 on Wnt/ β-catenin signaling is evolutionarily conserved across species

In our previous study, we found that zebrafish eaf1 and eaf2 share a high degree of identity with their human homologs EAF1 and EAF2. In addition, human EAF1/2 mRNA could efficiently rescue the phenotype associated with zebrafish eaf1/2 morphants (Liu et al., 2009). This prompted us to explore whether the suppressive function of eaf1 and eaf2 on Wnt/β-catenin signaling is conserved in humans.

Using zebrafish as our *in vivo* model, we overexpressed human EAF1/2 mRNA in embryos by mRNA injection and found that the human and zebrafish Eafs behaved in a very similar manner. At 72 hpf, the human EAF1/2-injected embryos exhibited an expanded forebrain and shortened bodies with either a short tail or no tail (75-85% of embryos; Fig. 7A,B), fully phenocopying the embryos injected with zebrafish eaf1/2 mRNA. We then used marker genes for further phenotyping. As expected, injection of human EAF1 or EAF2 mRNA resulted in expanded expression of six3b (Fig. 7C) and increased expression of both dkk1 and frzb (supplementary material Fig. S11A,B).

We next performed transcription assays using the TopFlash reporter. Human EAF1 and EAF2 significantly suppressed the reporter activity induced by ectopic expression of β -catenin in HEK

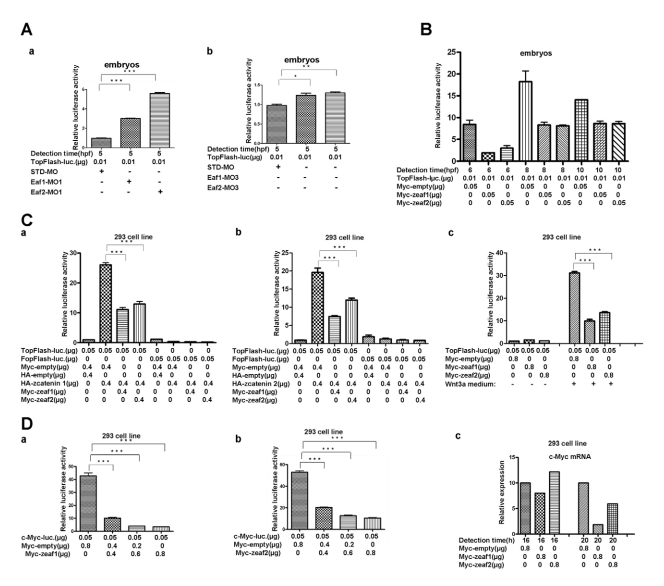


Fig. 5. *eaf1* and *eaf2* inhibit Wnt reporters. (**A**) Endogenous β-catenin transcriptional activity in zebrafish embryos was enhanced by knockdown of Eaf1 or Eaf2. One-cell stage embryos were injected with the plasmids indicated, together with 8xTopFlash as a reporter and *TK-renilla* as an internal control; luciferase activity was measured at 5 hpf. (**B**) Endogenous β-catenin transcriptional activity in zebrafish embryos was suppressed by ectopic expression of *eaf1* or *eaf2*. Luciferase activity was measured at 6, 8 or 10 hpf. (**C**) In the HEK 293 mammalian cell line, 8xTopFlash activity enhanced by Wnt/β-catenin was suppressed by overexpression of *eaf1* or *eaf2*. (a,b) 8xTopFlash activity enhanced by β-catenin 1 (zcatenin1) (a) or β-catenin 2 (zcatenin2) (b) was suppressed by both *eaf1* (zeaf1) and *eaf2* (zeaf2). (c) 8xTopFlash activity enhanced by addition of *Wnt3a*-conditioned medium was suppressed by ectopic expression of *eaf1* or *eaf2*. (**D**) The expression of the Wnt/β-catenin target gene human *c-MYC* was suppressed by *eaf1* or *eaf2*. (a,b) Dose-dependent downregulation of *c-MYC* promoter activity by *eaf1* (a) and *eaf2* (b). (c) Semi-quantitative RT-PCR analysis of endogenous human *c-MYC* expression after ectopic expression of *eaf1* or *eaf2*. Data from the luciferase reporter assays are reported as mean±s.d. of three independent experiments performed in triplicate; ****P<0.001, **P<0.05 (Student's *t*-test).

293 cells (Fig. 7D). Similarly, human *EAF1* and *EAF2* suppressed the *c-MYC* promoter activity (Fig. 7E).

Human EAF1 and EAF2 bound to human β -catenin as well as to other complex components, including Tcf, c-Jun and Axin (Fig. 7F,G). Similar to zebrafish *eaf1/2*, human *EAF1* and *EAF2* did not affect the protein stability of ectopic β -catenin or endogenous active β -catenin (supplementary material Fig. S11C,D). The cytoplasmic-nuclear translocation of β -catenin was also unaffected by overexpression of human EAF1 or EAF2 in HEK 293 cells (data not shown), and nor were the protein levels of the other β -catenin transcriptional complex proteins Tcf, c-Jun and Axin (supplementary material Fig. S11E-G).

Together, these observations suggest that the suppressive role of Eaf1 and Eaf2 in Wnt/ β -catenin signaling is evolutionarily conserved across species.

DISCUSSION

Zebrafish *eaf1* and *eaf2* are novel antagonists of Wnt/β-catenin signaling

In this report, loss-of-function of *eaf1* and *eaf2* in zebrafish embryos resulted in truncation of the anterior neuroectoderm, a similar morphology to embryos with enhanced Wnt signaling (Kelly et al., 1995; Kim et al., 2000; Kim et al., 2002). Gain-of-function of *eaf1* and *eaf2* resulted in enhanced expression of dorsal and anterior brain

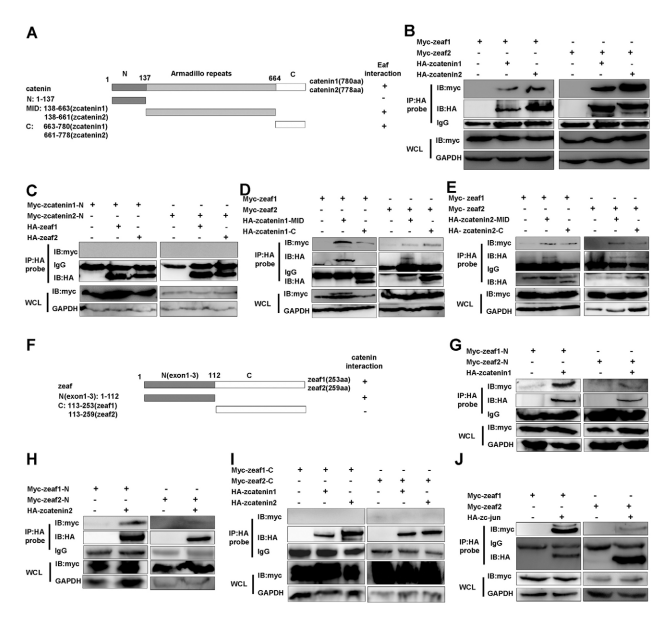


Fig. 6. Eaf1 and Eaf2 function as novel nuclear Wnt signaling components. (A) The domains in zebrafish β -catenin 1 and β -catenin 2 that interact with Eaf1 and Eaf2: +, interaction; -, no interaction. MID, the β -catenin center region that contains 12 Armadillo repeats. (B) Co-immunoprecipitation of Eaf1 or Eaf2 with full-length β-catenin 1 or β-catenin 2. (**C**) Eaf1 or Eaf2 did not interact with the N-terminus of β-catenin 1 or β-catenin 2. (**C**) Eaf1 or Eaf2 interacted with both the Armadillo repeats and the C-terminal transcriptional domain of β -catenin 1 (D) and β -catenin 2 (E). (F) The domains in Eaf1 and Eaf2 that interacted with β -catenin 1/2: N, exons 1-3 of Eaf1 or Eaf2; C, exons 4-6 of Eaf1 or Eaf2; +, interaction; -, no interaction. (**G,H**) β -catenin 1 (G) and β-catenin 2 (H) interacted with the N-terminus of Eaf1 or Eaf2. (J) β-catenin 1/2 did not interact with the C-terminus of Eaf1 or Eaf2. (J) Coimmunoprecipitation of Eaf1 or Eaf2 with full-length zebrafish c-Jun. WCL, whole cell lysate.

markers. This expression pattern mirrors that of wnt8 morphants, wnt8 mutants, or embryos expressing ectopic Wnt inhibitors (Lekven et al., 2001; Kim et al., 2002; Momoi et al., 2003). In addition, we observed enhanced expression of the Wnt/β-catenin maternal targets sqt and gsc at the blastula stage, the zygotic targets cdx4 and tbx6 at the later gastrula stage, and the direct target axin2 in midbrain at bud stage. All these observations indicate that eaf1 and eaf2 are required for forebrain and mesoderm patterning by negatively regulating Wnt/β-catenin signaling.

eaf1 and eaf2 counteracted the effects of enhanced Wnt/β-catenin signaling in apc mutants, as well as in embryos with ectopic expression of wnt8 or BIO treatment. However, β-catenin1-MO and dn-Tcf mRNA rescued the defects in gsc and six3b expression

exhibited by Eaf morphants. This suggests that eaf1 and eaf2 might antagonize Wnt/β-catenin signaling downstream of wnt8, but upstream of, or parallel to, β -catenin and tcf.

Interestingly, β-catenin2-MO effectively rescued dorsal gene expression in Eaf morphants, but caused severe forebrain defects in the same embryos as well as in those injected with mRNA encoding the N-terminus of Eaf1/2. This is in contrast to the effect of β catenin1-MO. This outcome is consistent with a previous report that β -catenin 1 and β -catenin 2 play essential but opposing roles in the formation of axis and neuroectoderm, and that β -catenin 2 is essential for dorsal mesoderm and forebrain formation (Bellipanni et al., 2006).

In a previous study we had shown that Eafs contribute to the regulation of convergence and extension movements through non-

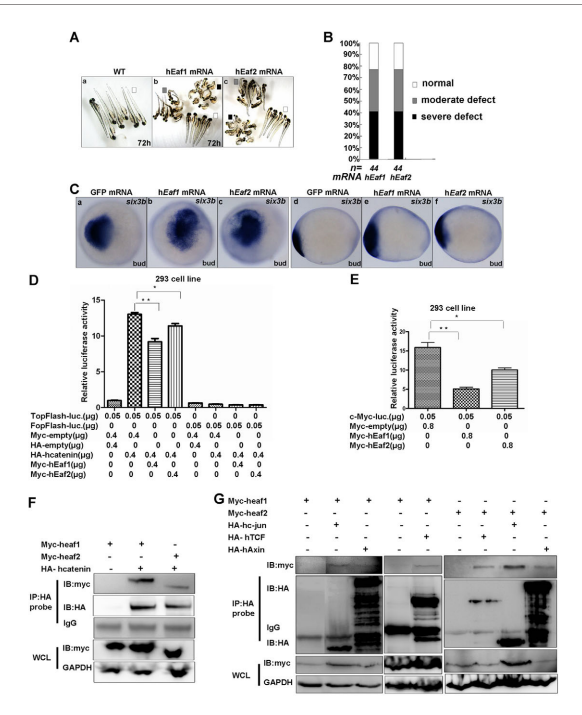


Fig. 7. Eaf1/2 activity is evolutionarily conserved from zebrafish to human. (**A**) Phenotypes of zebrafish embryos injected with human *EAF1* (hEaf1) or *EAF2* (hEaf2) mRNA at 72 hpf. (**B**) Morphology of zebrafish embryos injected with human *EAF1* or *EAF2* mRNA at 3 dpf (200-500 pg/embryo) was scored as normal (white), moderately defective (gray) or severely defective (black); embryos injected with *GFP* mRNA (200-500 pg/embryo) were used as a control. (**C**) *six3b* displayed enhanced expression in zebrafish embryos in which human *EAF1* or *EAF2* was ectopically expressed. (**D**) 8xTopFlash activity enhanced by human β-catenin was suppressed in the HEK 293 cell line by human EAF1 or EAF2. (**E**) Downregulation of *c-MYC* promoter activity by human full-length EAF1 or EAF2. Mean±s.d.; **P<0.01, *P<0.05 (Student's *t*-test). (**F**) Co-immunoprecipitation of human EAF1 or EAF2 with full-length human β-catenin. (**G**) Human EAF1 and EAF2 interact with components of the human β-catenin transcriptional complex, including TCF4, c-JUN and AXIN, *in vivo*. Ca-c, dorsal views, anterior to the left; Cd-f, lateral views, anterior to the left. WCL, whole cell lysate.

canonical signaling (Liu et al., 2009). Many factors, such as dkkl and naked, regulate both canonical and non-canonical Wnt signaling (Caneparo et al., 2007; Van Raay et al., 2007). In embryos with ectopic expression of factors that promote Wnt/ β -catenin signaling, such as wnt8a and active β -catenin, Eaf expression was reduced, but remained unchanged in embryos with ectopic expression of the

Wnt/ β -catenin inhibitor *frzb*. Together with the evidence that Eafs negatively regulate Wnt/ β -catenin signaling, this supports the possible existence of a regulatory feedback loop between *eaf1/2* and canonical Wnt signaling.

In this study, *eaf1* and *eaf2* did not appear to function redundantly, even though they exhibit partially redundant function in regulating

convergence and extension movements (Liu et al., 2009). Regarding the similar phenotypes shown in embryos injected with each MO alone (Eaf1-MO or Eaf2-MO), the function of *eaf1* and *eaf2* might be dosage dependent.

Eaf1 and Eaf2 are novel components of the β-catenin transcriptional complex

Multiple factors serve as either antagonists or activators in Wnt/ β -catenin signaling. These factors then execute their function by modulating the protein degradation, stabilization or distribution of β -catenin or of other components of the β -catenin transcriptional complex (Morin et al., 1997; Behrens et al., 1998; Gan et al., 2008). Here, we provide evidence that Eaf1 and Eaf2 not only bind to β -catenin, but also to other components of the β -catenin transcriptional complex, such as Tcf, Axin and c-Jun, which are evolutionarily conserved between zebrafish and human. However, overexpression or knockdown of Eafs did not alter the total or nuclear levels of β -catenin protein or of the other components of the β -catenin transcriptional complex, which suggests that mechanisms other than the regulation of protein stability and localization account for the suppressive role of Eaf1 and Eaf2.

β-catenin transcriptional activity is reported to emanate from its most N-terminal Arm repeat region, as well as from its C-terminal region, with the latter being the most potent transactivation domain (van de Wetering et al., 1997; Hecht et al., 1999). Eaf1/2 interacted with both the Arm repeat domain and C-terminus of β-catenin, and β-catenin interacted with the N-terminus of Eaf1/2. *In vivo*, embryos with ectopic expression of Eaf-EnR and Eaf-N-EnR show the same phenotype as embryos injected with mRNA encoding full-length eaf1/2. In all, these observations suggest that Eaf1/2 either act as a novel transcriptional repressor or recruit a repressor for the βcatenin transcriptional complex. Of note, Eafs bind to the same regions of β -catenin that commonly interact with other β -catenin antagonists, such as ICAT, Chibby and APC (Tago et al., 2000; Takemaru et al., 2003; Mosimann et al., 2009). This further supports the notion of a suppressive function for Eaf1 and Eaf2 in Wnt/βcatenin signaling.

Because they harbor transactivation domains in their C-terminus, Eaf1 and Eaf2 are implicated as transcriptional activators (Simone et al., 2001; Simone et al., 2003; Kong et al., 2005; Xiao et al., 2008). Here, we provided evidence that Eaf1 and Eaf2 actually function as repressors rather than activators to inhibit the transactivity of β -catenin. These results suggest that Eaf1 and Eaf2 might act to suppress transactivation of their binding partners, but directly activate the expression of their own downstream genes as transcription factors. This phenomenon has been reported for other transcription factors, such as p53. As a classic transcription factor, p53 can inhibit HIF1 α transactivity by direct interaction (An et al., 1998).

Eaf1 and Eaf2 may function as tumor suppressors through antagonizing Wnt/ β -catenin signaling

Compelling evidence points to a role for Wnt/ β -catenin signaling in cancer (Kim et al., 2002; Moon et al., 2004; Clevers, 2006). Cancer-associated mutations result in the constitutive activation of Wnt/ β -catenin signaling (Clevers, 2006). Indeed, multiple cancers, such as colon cancer, hair follicle tumors, prostate cancer and leukemia, show a high frequency of aberrant stabilization and constitutive activation of β -catenin (Morin et al., 1997; Chan et al., 1999; Jamieson et al., 2004; Lo Celso et al., 2004). In addition, transgenic mice with constitutive activation of β -catenin develop prostatic intraepithelial neoplasia (mPIN) (Yu et al., 2009; Yu et al., 2011). Eaf2 has been showed to have a tumor suppressive function

in multiple cancers (Xiao et al., 2003; Xiao et al., 2008). In this study, through both loss- and gain-of-function assays in the zebrafish model, we revealed that eaf1 and eaf2 antagonize Wnt/ β -catenin signaling. Furthermore, we found that Eaf1 and Eaf2 dramatically suppress β -catenin transcriptional activity by interacting with β -catenin and other components of the β -catenin transcriptional complex, which is evolutionarily conserved between zebrafish and human. Although further studies in human tumor samples or mammalian models are required, we propose that the human Eaf gene family might act as tumor suppressors through inhibiting Wnt/ β -catenin signaling.

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Competing interests statement

The authors declare no competing financial interests.

Supplementary material

Supplementary material available online at http://dev.biologists.org/lookup/suppl/doi:10.1242/dev.086157/-/DC1

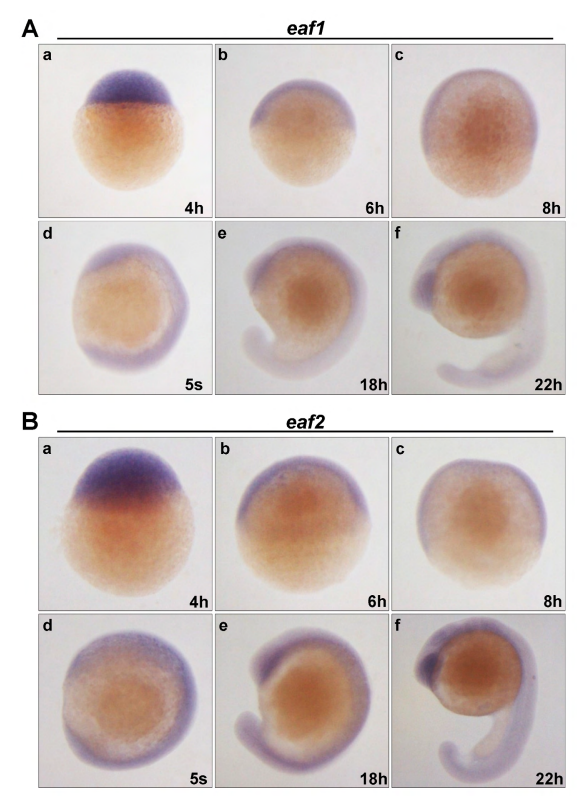
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Fire S1. The expression patterns of zebrafish eaf1 and eaf2 during embryogenesis. (A) eaf1 expression. (B) eaf2 expression. Lateral view, dorsal to the right.

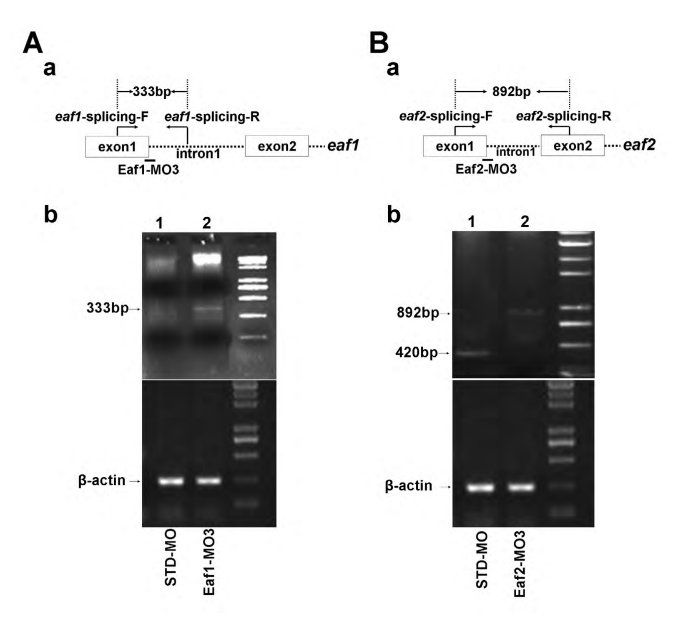


Fig. S2. Validation of Eaf1-MO3 and Eaf2-MO3. (A) (a) Schematic of *eaf1* exon-intron structure, Eaf1-MO3 targeting position and RT-PCR primer locations. (b) In STD-MO injection embryos, an expected 333 bp band could not be detected by RT-PCR, but in embryos with Eaf1-MO3 injection, a 333 bp band could be detected by RT-PCR. (B) (a) Schematic of *eaf2* exon-intron structure, Eaf2-MO3 targeting position and RT-PCR primer locations. (b) A 420 bp band was amplified from embryos injected with STD-MO by RT-PCR, but a 892 bp band was amplified from embryos injected with Eaf2-MO3, which contained intron 1 of *eaf2*. Embryos were collected at the bud stage for RNA extraction. Lane 1 is RNA from embryos injected with STD-MO, and lane 2 is RNA from embryos injected with Eaf1-MO3 (Ab) or Eaf1-MO3 (Bb).

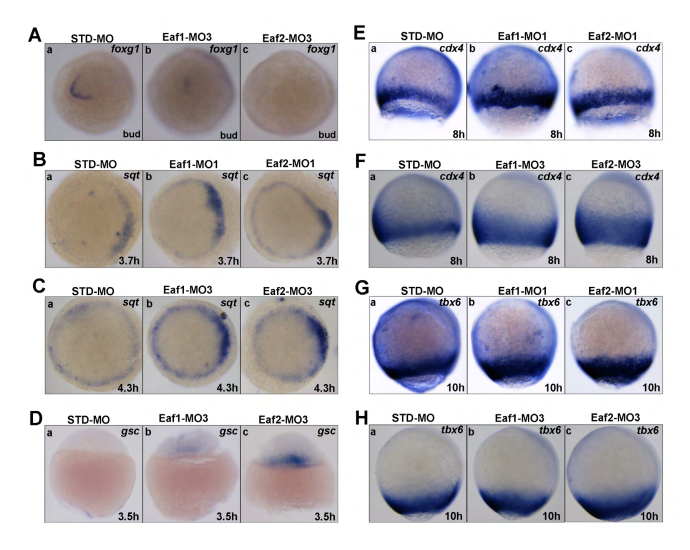


Fig. S3. Knockdown of zebrafish Eaf1 and Eaf2 causes defects in anterior neuroectoderm patterning and mesoderm patterning. (A) The expression of anterior neuroectoderm marker foxg1. (B-D) The expression of dorsal mesoderm marker genes sqt and gsc. (E-H) The expression of posterior ectoderm/mesoderm marker cdx4 and ventral mesoderm marker tbx6. (A) Dorsal view, anterior to the left; (D) dorsal view; (B,C) animal view, dorsal to the right; (E-H) lateral view, dorsal to the right.

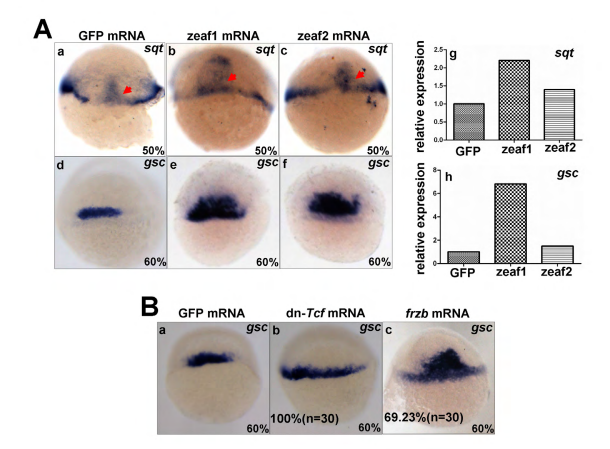


Fig. S4. Ectopic expression of zebrafish Eaf1 and Eaf2 increases the expression of *sqt* **and** *gsc.* (**A**) At the gastrula stage, dorsal marker genes *sqt* (a-c) and *gsc* (d-f) displayed expanded expression, particularly in the ventral domain. qRT-PCR was used to measure expression of *sqt* (g) and *gsc* (h) in these embryos. (**B**) In embryos injected with dn-*Tcf* mRNA (b) or *frzb* mRNA (c), the expression of *gsc* expanded ventral-laterally at the gastrula stage, compared with embryos injected with *GFP* mRNA (a). Dorsal views.

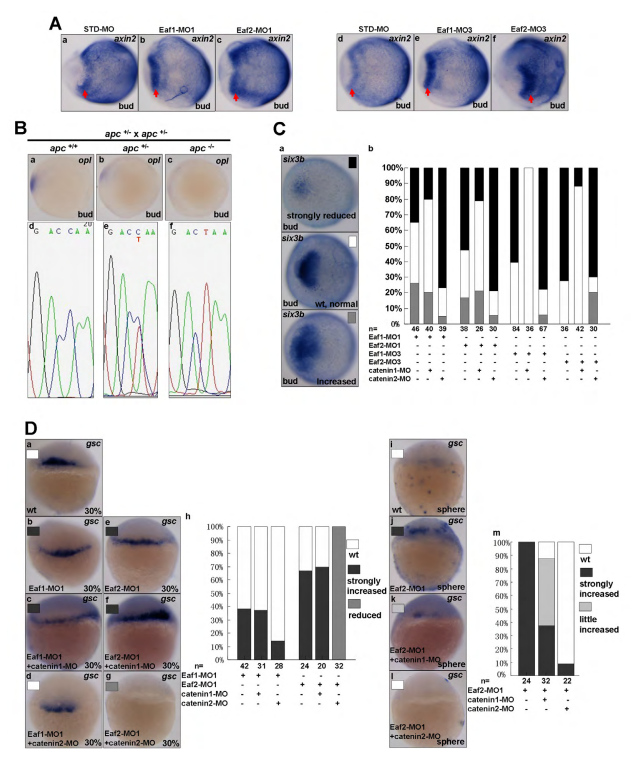


Fig. S5. *eaf1* and *eaf2* antagonize Wnt/β-catenin signaling. (A) The expression of *axin2*, a bona fide direct target of Wnt/β-catenin signaling at bud stage, was enhanced after knockdown of *eaf1* (b,e) or *eaf2* (c,f) (red arrows). (B) The offspring of $apc^{+/-} \times apc^{+/-}$ were genotyped by sequencing the mutant allele DNA fragment amplified from genomic DNA extracted from WISH-stained embryos. The embryos with normal *opl* expression contain the wild-type allele (C/C) (a,d), whereas the embryos with reduced *opl* expression contain the heterozygous allele (C/T) (b,e) and the embryos with strongly reduced *opl* expression contain the homozygous mutated allele (T/T) (c,f). (C) β-catenin1-MO, but not β-catenin2-MO, partially rescued the anterior brain defects in Eaf morphants. (D) At the 30% gastrula stage, β-catenin2-MO partially reduced the increased *gsc* expression associated with Eaf morphants (a-h); both β-catenin1-MO and β-catenin2-MO partially reduced the increased *gsc* expression of Eaf2-MO1 morphants at the sphere stage (i-m), and β-catenin2-MO was more effective (m). (A-C) Dorsal view, anterior to the left; (Da-g,i-l) dorsal view.

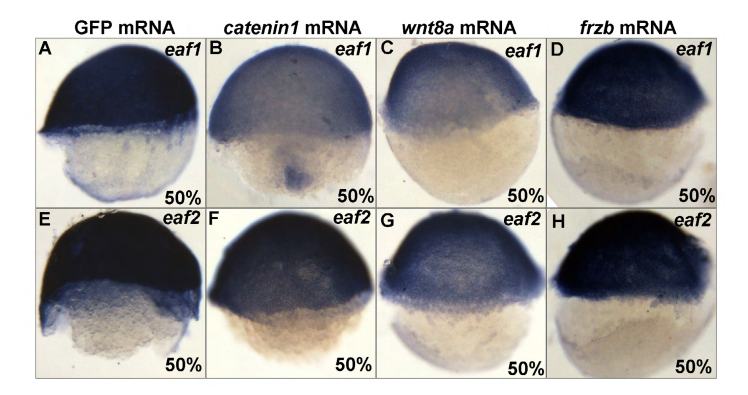


Fig. S6. The regulation of zebrafish *eaf1* and *eaf2* by Wnt/β-catenin signaling. The expression of *eaf1* and *eaf2* is downregulated in embryos with ectopic expression of active β-catenin I mRNA (**B**,**F**) or wnt8a mRNA (**C**,**G**) as compared with embryos injected with *GFP* mRNA (**A**,**E**), but was maintained in embryos injected with *frzb* mRNA (**D**,**H**). Lateral views, dorsal to the right.

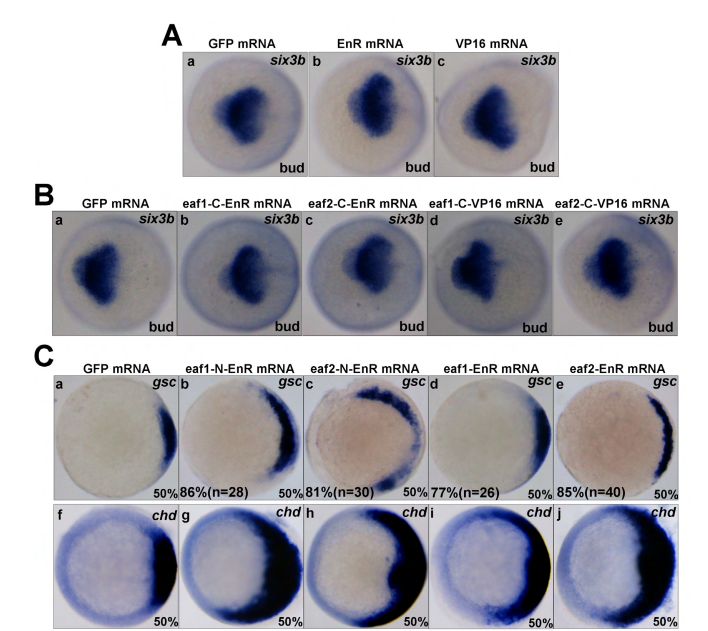


Fig. S7. VP16 and EnR are not functional *in vivo* and Eaf1/2 act as transcriptional repressors. (A,B) *Six3b* staining indicated normal patterning of anterior neural ectoderm in embryos injected with mRNA of the engrailed domain (EnR) or the VP16 domain (A), and in embryos injected with EnR or VP16 fused with the C-terminus of Eaf1 or Eaf2 (B). (C) In the embryos injected with *eaf1/2-EnR* or *eaf1/2-N-EnR* mRNA, the expression of the dorsal marker genes *gsc* and *chd* displayed similarly expanded patterns. (A,B) Dorsal views, anterior to the left; (C) animal views, dorsal to the right. All mRNA injections were 50-100 pg/embryo.

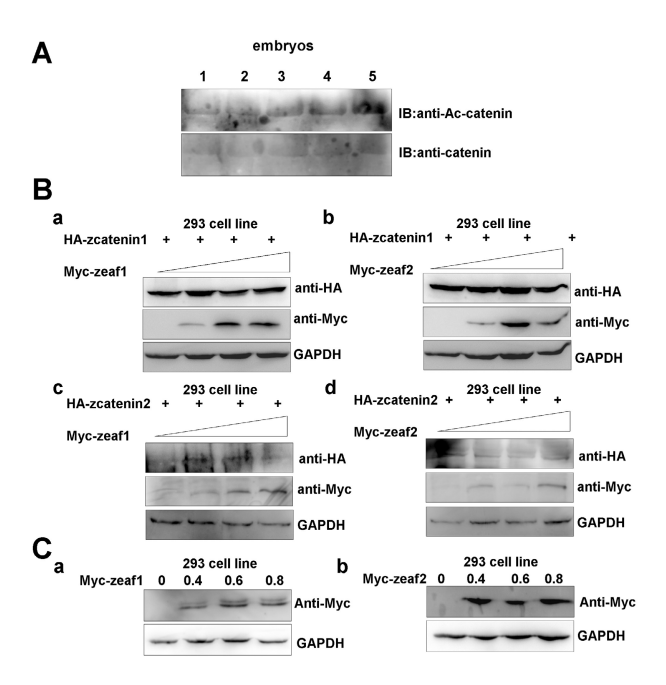


Fig. S8. Zebrafish Eaf1 or Eaf2 does not affect the protein level of either β-catenin 1 or β-catenin 2. (A) Western blot analysis for total β-catenin and active β-catenin protein (Ac-catenin) in embryos injected with STD-MO (lane 1), Eaf1-MO1 (lane 2), Eaf2-MO1 (lane 3), Eaf1-MO3 (lane 4) or Eaf2-MO3 (lane 5) (8 ng/per embryo). (B) The co-transfection of increased amounts of *eaf1* or *eaf2* together with β-*catenin 1* and β-*catenin 2* did not change the protein level of β-catenin 1 and β-catenin 2. (C) Western-blot analysis confirmed ectopic expression of zebrafish Eaf1 and Eaf2.

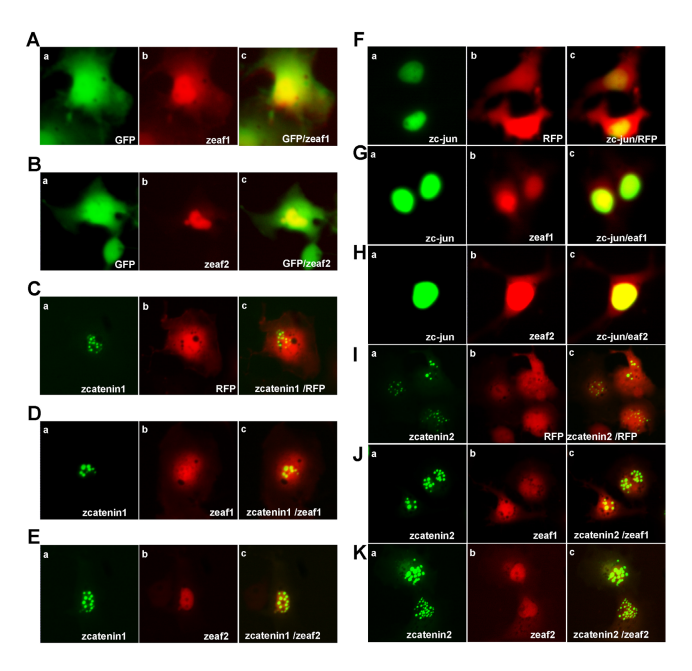


Fig. S9. Zebrafish Eaf1 or Eaf2 colocalizes with zebrafish β-catenin 1, β-catenin 2 and c-Jun. Cos-7 cells were transfected with: (**A**) empty GFP together with *eaf1*-RFP; (**B**) empty GFP together with *eaf2*-RFP; (**C**) empty RFP together with β-catenin1-GFP; (**D**) β-catenin1-GFP together with *eaf1*-RFP; (**E**) β-catenin1-GFP together with *eaf2*-RFP; (**F**) *c-jun*-GFP together with empty RFP; (**G**) *c-jun*-GFP together with *eaf1*-RFP; (**H**) *c-jun*-GFP together with *eaf2*-RFP; (**I**) β-catenin2-GFP together with *eaf1*-RFP; (**K**) β-catenin2-GFP together with *eaf2*-RFP. The cells were photographed under a fluorescence microscope 24 hours after transfection.

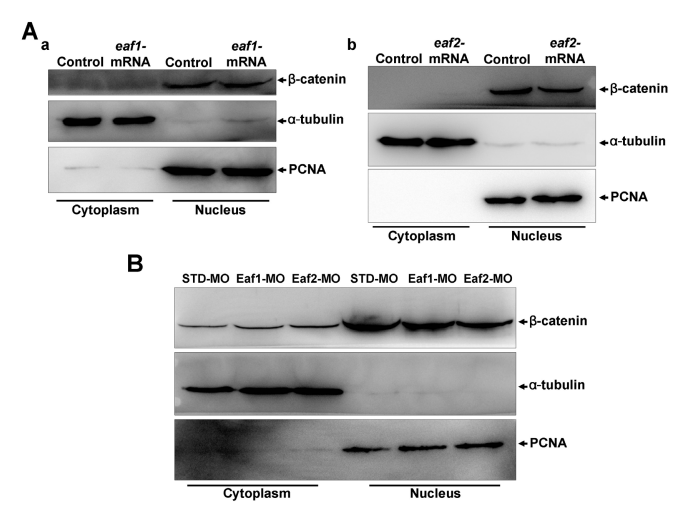


Fig. S10. Ectopic expression or knockdown of zebrafish eaf1 or eaf2 in embryos at the gastrula stage does not change cytoplasmic-nuclear translocation of β -catenin. (A) Ectopic expression of eaf1 (a) or eaf2 (b) did not change cytoplasmic-nuclear translocation of β -catenin. (B) Knockdown of eaf1 or eaf2 did not change cytoplasmic-nuclear translocation of β -catenin.

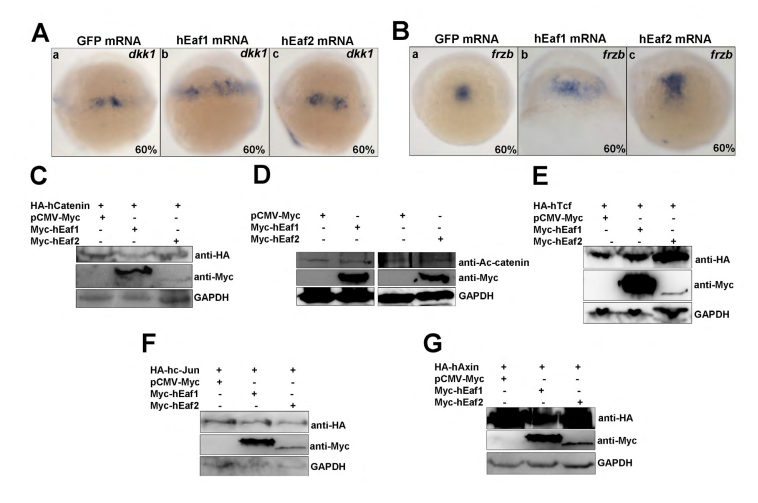


Fig. S11. Human Eaf function is evolutionary conserved in regulating Wnt/β-catenin signaling. (A,B) In zebrafish embryos injected with human EAF1 or EAF2 mRNA, dkkl (A) and frzb (B) displayed enhanced expression. (C) Overexpression of human EAF1 or EAF2 did not affect co-transfected human β-catenin stability. (**D**) Overexpression of human EAF1 or EAF2 did not change the endogenous active β-catenin protein level. (**E-G**) Ectopic expression of human TCF4 (E), human C-JUN (F) and human AXIN (G) after co-transfection with human EAF1 or EAF2 into the HEK 293T cell line.

Table S1. Primers used for qRT-PCR, probe amplification, splice MO validation, cDNA and promoter cloning

Primer	Sequence (5'-3')
sqtFq	GCTGGTGGAAGTGAAC
sqtRq	CCTTCACACCGATAAGCGTTG
gscFq	GCAAGAGACGACACCGAAC
gscRq	TGAACCAAACCTCTACCTTCTC
18sFq	GAGAAACGGCTACCACATCC
18sRq	CACCAGACTTGCCCTCCAA
c-mycFq (human)	GGCGAACACACACGTCTTGGA
c-mycRq (human)	CTTACGCACAAGAGTTCCGTAGC
18sFq (human)	TCAACTTTCGATGGTAGTCGCCGT
18sRq (human)	TCCTTGGATGTGGTAGCCGTTTCT
cdx4-probe-F	CGTCCATGAGGAACATACAGC
cdx4-probe-R	CAAGAGCCTCCAGCATTTCG
opl-probe-F	GGCGAAGTTACAGACAGA
opl-probe-R	GACATGACCGTATTGCTC
six3b-probe-F	TTTGGTCGTTGCCCGTAG
six3b-probe-R	CGTGATGCTGAAGCCTGT
eaf1-splicing-F	TGGTGAATTCAATGTGACGCGCAG
eaf1-splicing-R	GATTTGTTGGTGCGAATGTGGT
eaf2-splicing-F	AGAAGTGAGGCGTCTATTTCTGCC
eaf2-splicing-R	CTTGACCGCAATGTTGTTGCTGAG
zf-catenin2-F	ATATCTCTAGAAGGATTGACGCAACGATGGCTAG
zf-catenin2-myc-R	ATATCGCGGCCGCGTCCTTCGCTCAGCAGCTCTCTA-3
zf-catenin1-F	ATATCGAATTCGGTCTAGATGGCTACCCAGTCTGACTTGA

zf-catenin1-myc-R	ATATCGCGGCCGCTTACAGATCGGTGTCAAACCAGG
zf-c-jun-F	ATATCGAATTCGGTCTAGACCTTCTATGTCTACCAAGATGG
zf-c-jun-myc-R	ATATCGCGGCCGCGTCCTTCGGCTCTCCTCAGAAG
apc-sequencing-F	CTACCCAACTTTACCTATATCAG
apc-sequencing-R	GACTCTCAAAACTGTCAAGGG

Primers are zebrafish, except where marked for human.

Table S2. Morpholino sequences

Morpholino	Sequence (5'-3')
Eaf1-ATG-MO (Eaf1-MO1)	GCGGCGGTTCGAGCTGCCGTTCAT
Eaf2-ATG-MO (Eaf2-MO1)	ATGCTGTTCCATTCATTCTAATCCA
Eaf1-splice-MO (Eaf1-MO3)	GTCTCTTGATGGACTCACATCTG
Eaf2-splice-MO (Eaf2-MO3)	AAAAGATGCAACTTACATCGTACCG
β-catenin1-MO	CTGGGTAGCCATGATTTTCTCACAG
β-catenin2-MO	CCTTTAGCCTGAGCGACTTCCAAAC