## Report

## The *C. elegans* Zonula Occludens Ortholog Cooperates with the Cadherin Complex to Recruit Actin during Morphogenesis

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#### Summary

The dramatic cell-shape changes necessary to form a multicellular organism require cell-cell junctions to be both pliable and strong. The zonula occludens (ZO) subfamily of membrane-associated guanylate kinases (MAGUKs) are scaffolding molecules thought to regulate cell-cell adhesion [1–3], but there is little known about their roles in vivo. To elucidate the functional role of ZO proteins in a living embryo, we have characterized the sole C. elegans ZO family member, ZOO-1. ZOO-1 localizes with the cadherin-catenin complex during development, and its junctional recruitment requires the transmembrane proteins HMR-1/E-cadherin and VAB-9/claudin, but surprisingly, not HMP-1/a-catenin or HMP-2/β-catenin. zoo-1 knockdown results in lethality during elongation, resulting in the rupture of epidermal cell-cell junctions under stress and failure of epidermal sheet sealing at the ventral midline. Consistent with a role in recruiting actin to the junction in parallel to the cadherin-catenin complex, zoo-1 loss of function reduces the dynamic recruitment of actin to junctions and enhances the severity of actin filament defects in hypomorphic alleles of hmp-1 and hmp-2. These results show that ZOO-1 cooperates with the cadherin-catenin complex to dynamically regulate strong junctional anchorage to the actin cytoskeleton during morphogenesis.

## **Results and Discussion**

## ZOO-1, the Sole Zonula Occludens Ortholog in *C. elegans,* Localizes to Junctions during Morphogenesis

The *C. elegans* genome contains a single predicted ortholog of the zonula occludens protein family, ORF Y105E8A.26, which we have named *zoo-1*, for zonula occludens ortholog (Figure S1 available online). We assayed ZOO-1 expression via immunostaining (Figure 1; Figure S2); a *zoo-1::gfp* construct shows identical localization (Movie S1). During morphogenesis, ZOO-1 becomes enriched at the borders of epidermal cells (Figure 1A; Figures S2D–S2F); elongating embryos exhibit the strongest junctional accumulation (Figures S2G and S2H). ZOO-1 is also expressed in myoblasts and persists in mature muscle cells (Figure S2G). In contrast, *zoo-1::gfp* driven by an epithelial promoter shows no muscle-associated signal (data not shown); thus muscle-associated ZOO-1 signal is due to expression specifically in muscle.

In cultured epithelial cells, ZO-1 initially associates with the adherens junction (AJ) and segregates apically to the tight junction as cells mature [4–7]. The apical junction in epidermal cells of *C. elegans* has two subdomains with distinct multiprotein complexes, the cadherin-catenin and DLG-AJM complexes [8], which can be partially resolved via light microscopy in embryos [9]. Quantitative colocalization analysis shows a high degree of overlap between ZOO-1, HMP-1/ $\alpha$ -catenin, and JAC-1/p120 catenin but not between ZOO-1 and the DLG-1/AJM-1 complex (Figure S3).

### ZOO-1 Recruitment to Junctions Is Dependent on HMR-1/Cadherin and VAB-9/BCMP1 but Independent of HMP-1/ $\alpha$ -Catenin and HMP-2/ $\beta$ -Catenin

We next examined molecular requirements for ZOO-1 recruitment. Unlike AJM-1, which depends on DLG-1/Discs large for localization, ZOO-1 localizes properly in *dlg-1(RNAi)* embryos (Figures 1D-1F). Previous work in tissue culture has suggested that localization of vertebrate ZO-1 to the AJ may depend on  $\alpha$ -catenin [10, 11]. We tested this in vivo by immunostaining hmp-1(zu278) null embryos for ZOO-1. However, ZOO-1 junctional localization appears largely unaffected in hmp-1 zygotic null (data not shown) embryos, as it does in hmp-1(RNAi) (Figures 1G-1I) or hmp-2/β-catenin (RNAi) embryos (data not shown), in which both maternal and zygotic mRNA are removed [12, 13]. In contrast, hmr-1/E-cadherin (RNAi) completely disrupts epidermal ZOO-1 localization (Figures 1J-1L), although localization in muscle is unaffected. VAB-9/BCMP1 also localizes to the cadherin-catenin complex in epidermal cells in C. elegans [9]; ZOO-1 expression in vab-9(ju6) mutants is very similar to that in hmr-1(RNAi) embryos (Figures 1M-10). These results suggest that both HMR-1 and VAB-9 are essential for recruiting ZOO-1 to the apical junction, but that they act upstream of HMP-2 and HMP-1.

Vertebrate ZO proteins directly interact with multiple claudin family members [14, 15]. However, we could detect no effects on junctional recruitment of ZOO-1 in *clc-1/2* single or double RNAi embryos nor epithelial permeability defects in *zoo-1* loss-of-function embryos in a standard assay [16] (data not shown). These data suggest that ZOO-1 is not an essential component of the paracellular permeability pathway in *C. elegans*.

#### zoo-1 Knockdown Results in Morphogenesis Defects

To examine consequences of loss of *zoo-1* function, we analyzed two *zoo-1* mutants, but both result in incomplete loss of *zoo-1* gene function (Supplemental Results). In order to achieve more complete *zoo-1* loss of function, we performed RNAi in an RNAi hypersensitive background (*rrf-3(pk1426*); [17]). *zoo-1(RNAi);rrf-3* embryos have no detectable ZOO-1 expression as assessed via immunostaining (Figure S4), and knockdown can be achieved via multiple different RNAs that target *zoo-1* (data not shown). *rrf-3(pk1426*) homozygotes exhibit low levels of embryonic lethality (11.4% ± 6.3%, mean ± SD, n = 272) prior to the initiation of epidermal

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morphogenesis, because of gastrulation failure (data not shown). Although the penetrance of gastrulation defects in double mutants is similar to rrf-3 single mutants, zoo-1(RNAi) in an rrf-3(pk1426) background increased overall lethality to 33.2% ± 5.9% (n = 247) and yielded multiple morphogenetic defects (Figure 2). zoo-1(RNAi);rrf-3 embryos (Figures 2G-2I) properly complete ventral enclosure and initiate elongation; however, the rate of elongation is markedly slower than in wild-type (Figures 2A-2C; Movie S2) or rrf-3 (Figures 2D-2F; Movie S3) animals, and abnormal bulges develop along the body (Figure 2I). Body wall muscle is functional in arrested zoo-1;rrf-3 embryos, which continue to twitch, and muscle morphology appears normal via phalloidin staining (Figure S4H), suggesting that these defects are epidermal in nature. 6% of zoo-1(RNAi);rrf-3 embryos exhibit epidermal rupture during elongation (Figures 2J-2L; Movie S4). The distribution and dynamics of HMP-1::GFP and JAC-1::GFP are normal in living zoo-1(RNAi);rrf-3 embryos, and we could not detect enhancement of morphogenetic defects after ZOO-1 depletion in vab-9(ju6) null mutants (data not shown). The simplest interpretation of these results is that ZOO-1 acts downstream of VAB-9 to stabilize junctional intearity.

That zoo-1(RNAi);rrf-3 embryos rupture suggests reduced resistance of apical junctions to actomyosin-mediated contractility. We therefore generated hypocontractile zooFigure 1. ZOO-1 Junctional Recruitment Is Dependent on HMR-1/E-Cadherin and VAB-9/ Claudin, but Not on HMP-2/ $\beta$ -Catenin or HMP-1/ $\alpha$ -Catenin

(A-L) Confocal images of elongating embryos stained for ZOO-1 (green in [A], [D], [G], [J], [M]), AJM-1 as a junctional marker (red in [B], [E], [H], [K], [N]), and the merged image (C, F, I, L, O). Wild-type (A-C), *dlg-1(RNAi)* (D-F), and *hmp-1(RNAi)* (G-I) embryos display proper junctional localization of ZOO-1, despite disruption of AJM-1 localization in the case of dlg-1(RNAi) (E). (J-L) *hmr-1(RNAi)* embryo exhibits abrogated junctional ZOO-1 staining, though staining persists in sarcomeres (J, arrow).

(M–O) *vab-9(ju6*) embryo lacks junctional ZOO-1 staining, though ZOO-1 localization in sarcomeres in unaffected ([M], arrow). Scale bar represents 10  $\mu$ m.

1(RNAi);rrf-3 embryos by using simultaneous weak RNAi against let-502/Rho kinase, and we enhanced contractility by performing zoo-1(RNAi) in mel-11(it26)/myosin phosphatase mutants, in which myosin presumably remains phosphorylated and hence abnormally active [18]. let-502(RNAi) resulted in a reduction of rupture of zoo-1(RNAi);rrf-3 embryos from 12% to 4% (n = 66 and 116 embryos examined, respectively; significantly different, p < 0.04, Fisher's exact test), whereas zoo-1(RNAi) knockdown in mel-1(it26) homozygotes resulted in the appearance of early ruptures prior to the 1.5-fold stage (n = 53and 22 embryos examined for zoo-

1(*RNAi*);*mel-11(it26*) and *mel-11(it26*), respectively; Figures 2M–2O; Figure S5; significantly different from *mel-11* alone, p < 0.008). Based on these results, we conclude that ZOO-1 is especially important to provide mechanical stability to epidermal junctions.

# *zoo-1* (RNAi) Reduces Junctional Actin Recruitment, Leading to Perturbed Actin Filaments

We next visualized actin dynamics with an F-actin reporter expressed specifically in the epidermis, the actin-binding domain of VAB-10 fused to GFP [19]. We observed a significant decrease in actin localized near cell-cell junctions. Actin in this region aligns into a robust cable parallel to cell-cell boundaries in rrf-3 embryos (Figures 3A and 3C), whereas in zoo-1(RNAi);rrf-3 embryos, junctional actin is less robust (Figures 3B and 3D). Quantitative analysis (see Figure S6 for description) confirms these observations: the ratio of junctional to cytoplasmic actin in wild-type is 2.16 ± 0.26 (mean ± SD, n = 22 cells in 4 embryos measured) versus 1.4 ± 0.2 in zoo-1(RNAi);rrf-3 embryos (n = 23 cells in 5 embryos; significantly different based on a two-tailed Student's t test, p < 0.01). During elongation, actomyosin contractile forces act along circumferential actin bundles (CFBs), which attach at their ends to cell-cell junctions and are thought to distribute the forces driving elongation. In untreated embryos, CFBs are evenly spaced (Figure 3E). Strikingly, in zoo-1 (RNAi) embryos, some CFBs cluster abnormally



(Figure 3F), suggesting that ZOO-1 contributes to their anchorage.

Because some *zoo-1(RNAi);rrf-3* embryos rupture during elongation, we imaged F-actin during ventral enclosure, when midline junctional connections are established. In contrast to wild-type embryos, which accumulate robust junctional actin at the ventral midline (Figure 3A; Movie S5), in *zoo-1(RNAi);rrf-3* embryos that display midline bulges near the end of enclosure, we consistently found loss of accumulation of midline junctional actin (Figure 3B; n = 6/6 embryos with midline defects examined), or failure to establish a midline connection entirely between one or more cells (Movies S6 and S7; n = 4/6 embryos with midline junctional failure). Taken together, the abnormalities in actin organization we observe in *zoo-1* knockdown embryos provide a mechanical explanation for observed defects at the end of ventral enclosure and during elongation.

# *zoo-1* (RNAi) Enhances the Lethality of *hmp-1/α*-Catenin and *hmp-2/*β-Catenin Hypomorphs

Connecting the actin cytoskeleton to cell-cell junctions is a role traditionally assigned to the cadherin-catenin complex [20]. Because ZOO-1 recruitment is independent of both  $\alpha$ - and  $\beta$ -catenin, we hypothesized that ZOO-1 recruits actin to the junction in a parallel pathway. To test this hypothesis, we examined the combined effects of *zoo-1* (RNAi) and weak loss of function for  $\beta$ -catenin and  $\alpha$ -catenin, by using *hmp-2* (*qm39*) ([21]; M. Costa, personal communication) and *hmp-1(fe4)* [22],

Figure 2. Loss of *zoo-1* Function Causes Embryonic Lethality

Nomarski images of representative embryos undergoing elongation are shown. t = 0 correlates with 90 min after ventral enclosure.

(A–C) Wild-type embryo.

(D-F) rrf-3(pk1426) embryo.

(G-I) *zoo-1(RNAi);rrf-3(pk1426)* embryo exhibiting failed elongation with pronounced bodyshape defects.

(J–L) *zoo-1(RNAi);rrf-3(pk1426*) embryo that has ruptured from the posterior region. Note the delayed elongation of the *zoo-1(RNAi);rrf-3(pk1426*) embryos relative to wild-type.

(M–O) zoo-1(RNAi);mel-11(it26) embryo. Note the ventral rupture ([N], arrow).

Scale bar represents 10 µm.

respectively. hmp-2(qm39) displays 6% ± 1.2% (n = 832) embryonic and early larval lethality at 20°C (Table 1). In hmp-2(qm39);zoo-1(RNAi) embryos, lethality is significantly enhanced to 60% ± 4.6% (n = 508) and mutants exhibit delayed development (Table 1). Unlike wild-type embryos (Figure 4A), progeny of hmp-1(fe4) hermaphrodites show pronounced elongation defects (Figure 4B; Movie S8) and exhibit  $77.5\% \pm 7.9\%$  (n = 844) embryonic and early larval lethality (Table 1; [22]). Phalloidin staining of hmp-1(fe4) embryos confirms that the spatial arrangement of CFBs is occasionally perturbed ([27]; Figure 4E). zoo-1(RNAi) in hmp-1(fe4) mutants enhances overall lethality to 99.6% ± 1.8% (n = 818) and causes nearly all embryos to exhibit the

Humpback phenotype (Figure 4D; Movie S9). Approximately half of *zoo-(RNAi);hmp-1(fe4)* mutants ultimately rupture at various positions along the body axis (Figure 4C; Movie S10).

*zoo-1* loss of function also exacerbates the cytoskeletal defects observed in *hmp-1(fe4)* embryos: CFBs often cluster, resulting in inappropriately thick bundles (Figure 4F) similar to *hmp-1* null mutants and the most severe *hmp-1(fe4)* embryos [12, 22], suggesting that ZOO-1 and the cadherin complex act in parallel to stabilize actin at epidermal junctions. Loss of UNC-34/Ena also synergizes with *hmp-1(fe4)*, but unlike ZOO-1, UNC-34 is correctly localized in *hmr-1* mutant backgrounds [23]. We found no evidence for synergistic lethality between *zoo-1* and the null allele, *unc-34(gm104)* (data not shown).

In conclusion, we have provided in vivo analysis of ZOO-1/ ZO-1 in *C. elegans*, and we show that ZOO-1 acts at junctions along with core AJ proteins during epithelial morphogenesis. Recent studies in *Drosophila* have implicated ZO-1/Pyd at AJs, based on defects in cell rearrangement during tracheal morphogenesis in *pyd* mutants [24]. However, these same studies have also implicated *pyd* in nuclear functions. Because it lacks the nuclear localization sequence found in other ZO-1 orthologs, ZOO-1 provides a "natural experiment" that can identify exclusively non-nuclear roles for ZO-1 proteins.

In ZO-1/ZO-2/ZO-3 knockdown cells in culture, AJ maturation is delayed [25]. In contrast, we do not find a delay in recruitment of core AJ components in *zoo-1* knockdown embryos in vivo. It is possible that the extremely rapid kinetics



Figure 3. Loss of *zoo-1* Function Disrupts Actin Accumulation at Cell-Cell Junctions

(A and B) Ventral views of a wild-type (A) and *zoo-1(RNAi);rrf-3* (B) embryo at the end of ventral enclosure expressing a *gfp*-tagged fragment of *vab-10* that binds F-actin in epidermal cells [19]. In the wild-type embryo, two pairs of anterior cells have accumulated dense actin at the midline ([A], arrows), whereas only small actin puncta ([B], left arrow) or detached actin filaments ([B], right arrow) remain at the same position in the *zoo-1(RNAi);rrf-3* embryo ([B], arrows).

(C) Robust actin cables are visible at cell-cell borders in epidermal cells in a comma stage embryo (arrows).

(D) Actin is less evenly distributed at junctions in epidermal cells of comma stage *zoo-1* knockdown embryos (arrows).

(E and F) Embryos at the two-fold stage of elongation stained for F-actin. (E) Wild-type embryo.

(F) *zoo-1(RNAi)* embryo. Note the abnormal clustering of circumferential actin filament bundles in the *zoo-1(RNAi)* embryo (arrow).

Scale bars represent 5  $\mu$ m.

of junction formation in model invertebrates (minutes, as opposed to many hours in vertebrates) accounts for this difference, as has been previously suggested [22]. However, dynamic imaging of actin in living embryos after *zoo-1* knockdown revealed dramatic effects on actin recruitment at maturing junctions, which may be analogous to defects observed in the transition from "spot-like" to "belt-like" AJs in cultured cells after ZO protein depletion [25, 26]. Although ZO-1 directly binds actin [11, 27], the lack of sequence conservation in this region of ZOO-1 does not immediately suggest that ZOO-1 does so. Unfortunately, the dominant lethality of *zoo-1* transgenes has thus far precluded unambiguous structure-function analysis to address this issue.

C. elegans embryos undergoing epidermal morphogenesis do not require HMP-1/ $\alpha$ -catenin or HMP-2/ $\beta$ -catenin for junctional recruitment of ZOO-1. In contrast, ZOO-1 recruitment does depend on HMR-1/E-cadherin and VAB-9/BCMP1. Because HMR-1 is required for proper junctional localization of VAB-9 [9], the simplest explanation for these localization results is HMR-1 $\rightarrow$ VAB-9 $\rightarrow$  $\rightarrow$ ZOO-1. Because there are very few cytoplasmic residues in VAB-9 that could engage in direct binding to ZOO-1, we think it unlikely that the interaction between the two proteins is direct. Instead, another protein presumably recruits ZOO-1 to AJs. Future studies that characterize the binding affinities of ZOO-1 at epidermal junctions

Table 1. zoo-1 Lethality in rrf-3, hmp-1(fe4), and hmp-2(qm39)	
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Genotype	% Lethality	SD (nª)
 rrf-3(pk1426)	11.4	±6.3 (272)
zoo-1(RNAi);rrf-3(pk1426)	33.2	±5.9 (247)
hmp-1(fe4)	77.5	±7.9 (844)
zoo-1(RNAi);hmp-1(fe4)	99.6	±1.8 (818)
zoo-1(cxTi8317);hmp- 1(fe4)	99.8	±0.6 (1114)
hmp-2(qm39)	6	±1.2 (832)
zoo-1(RNAi);hmp- 2(qm39)	60	±4.6 (508)

<sup>a</sup>Number of embryos counted. Numbers are the sum of at least three separate experiments for each genotype.

should clarify the role of this highly conserved protein during morphogenesis.

#### Supplemental Data

Supplemental Data include Supplemental Results, Supplemental Experimental Procedures, six figures, and ten movies and can be found with this article online at http://www.current-biology.com/cgi/content/full/18/17/ 1333/DC1/.

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Figure 4. zoo-1(RNAi) Enhances the Elongation Defects of hmp-1(fe4) Mutants

(A–C) Nomarski images at 90 min time intervals of representative embryos undergoing elongation. (A) Wild-type embryo.

(B) *hmp-1(fe4)* embryo with visible body-shape defects.

(C) *zoo-1(RNAi);hmp-1(fe4)* embryo that has ruptured from the ventral surface (arrow).

(D) Distribution of embryonic lethal phenotypes of *hmp-1(fe4)* and *zoo-1(RNAi);hmp-1(fe4)* animals.

(E and F) Representative confocal images of F-actin staining in a *hmp-1(fe4)* (E) and *zoo-1(RNAi);hmp-1(fe4)* (F) embryo. The organization of circumferential actin filaments is consistently and markedly disrupted in *zoo-1(RNAi);hmp-1(fe4)* embryos.

Scale bar represents 10 µm.

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