# Two *Caenorhabditis elegans* calponin-related proteins have overlapping functions to maintain cytoskeletal integrity and are essential for reproduction

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#### Abstract

Multicellular organisms have multiple genes encoding calponins and calponin-related proteins, and some of these are known to regulate actin cytoskeletal dynamics and contractility. However, functional similarities and differences among these proteins are largely unknown. In the nematode Caenorhabditis elegans, UNC-87 is a calponin-related protein with seven calponin-like (CLIK) motifs and is required for maintenance of contractile apparatuses in muscle cells. Here, we report that CLIK-1, another calponin-related protein that also contains seven CLIK motifs, has an overlapping function with UNC-87 to maintain actin cytoskeletal integrity in vivo and has both common and different actin-regulatory activities in vitro. CLIK-1 is predominantly expressed in the body wall muscle and somatic gonad, where UNC-87 is also expressed. unc-87 mutation causes cytoskeletal defects in the body wall muscle and somatic gonad, whereas clik-1 depletion alone causes no detectable phenotypes. However, simultaneous depletion of *clik-1* and unc-87 caused sterility due to ovulation failure by severely affecting the contractile actin networks in the myoepithelial sheath of the somatic gonad. In vitro, UNC-87 bundles actin filaments. However, CLIK-1 binds to actin filaments without bundling them and is antagonistic to UNC-87 in filament bundling. UNC-87 and CLIK-1 share common functions to inhibit cofilin binding and allow tropomyosin binding to actin filaments, suggesting that both proteins stabilize actin filaments. Thus, partially redundant functions of UNC-87 and CLIK-1 in ovulation is likely mediated by their common actin-regulatory activities, but their distinct activities in actin bundling suggest that they also have different biological functions.

#### Introduction

Calponins and calponin-related proteins regulate actin filament bundling, actin filament stability, and actomyosin contractility in a wide variety of cell types (1-3). Many of these

proteins contain a single calponin-homology (CH) domain and various numbers of calponinlike (CLIK) motifs (also known as CLIK repeats). Although CH domains are recognized as actin-binding domains of several proteins, such as fimbrin,  $\alpha$ -actinin, and spectrin (4,5), the CH domain of calponin is not an actin-binding site (6-9). Instead, CLIK motifs in the Cterminus of calponin act as actin-binding sites (6,9,10). CLIK motifs are 23-amino-acid sequences and present in variable numbers in calponins and calponin-related proteins (9,11). Vertebrate SM22/transgelin and yeast calponinrelated proteins have only one CLIK motif (12-16). Vertebrate calponins have three CLIK motifs (17), whereas calponins in flatworms and mollusks have five CLIK motifs (18-20). UNC-87 in the nematode *Caenorhabditis elegans* has seven CLIK motifs (21). Importantly, the number of CLIK motifs correlates with the strength of actin-filament binding *in vitro* (22) and inhibitory effects on actin filament dynamics in cultured cells (10,23). These observations suggest that CLIK motifs are modular actin-binding sequences with incremental affinity with actin filaments depending on the number of motifs. However, whether all CLIK motifs have the same or different properties remains unknown.

Nematodes have uniquely evolved calponin-related proteins with multiple CLIK motifs and without a CH domain. C. elegans UNC-87, which contains seven CLIK motifs (21), is predominantly expressed in muscular tissues and associated with actin filaments (21,24,25). Two isoforms, UNC-87A and UNC-87B, are expressed by two alternative promoters for two separate first exons, and they share all CLIK motifs and similar biochemical properties (21,24). Mutations in the unc-87 gene cause disorganization of sarcomeric actin filaments in the body wall muscle due to defects in the maintenance of sarcomeres (21,26,27). UNC-87 is conserved in other nematode species and play a critical role in worm motility (28,29). In vitro, UNC-87 bundles actin filaments (22), inhibits

actin-filament severing by actin-depolymerizing factor (ADF)/cofilin (25), and inhibits actomyosin ATPase (24). In addition, the C. *elegans* genome has three other genes encoding similar calponin-related proteins with various numbers of CLIK motifs: *clik-1* (seven motifs), clik-2 (six motifs), and clik-3 (five motifs). A proteomic study reported that CLIK-1, CLIK-2, and CLIK-3 are enriched in the pharyngeal muscle (30). Null mutations of *clik-2* and *clik-3* cause pumping defects in the pharynx, whereas null mutation of *clik-1* causes no detectable phenotypes (31). Therefore, function of *clik-1* remains unknown. In this study, we aimed to determine the function of *clik-1*. Our results indicate that UNC-87 and CLIK-1 are coexpressed in muscle tissues and have overlapping and critical functions in the reproductive system by maintaining cytoskeletal integrity in the somatic gonad. Biochemical characterization showed that UNC-87 and CLIK-1 have common and distinct actinregulatory activities in vitro.

## Results

# CLIK-1 and UNC-87 are partially redundant and required for C. elegans reproduction

C. elegans has four genes (unc-87, clik-1, clik-2, and clik-3) encoding calponin-related proteins with no CH domains. Among them, primary structures of UNC-87B (hereafter referred to as UNC-87) and CLIK-1 (also known as T25F10.6, GenBank accession number: CCD74240) are very similar (42 % identity) containing seven CLIK motifs (C1 - C7) (Fig. 1A). Comparison of CLIK motifs from representative calponins and calponin-related proteins shows that CLIK motifs are highly conserved across species (Fig. 1B). However, sequences of C5, C6, and C7 of CLIK-1 are diverged in their N-terminal halves (Fig. 1B), suggesting that biochemical properties of CLIK-1 are somewhat different from those of UNC-87.

To determine expression pattern and subcellular localization of CLIK-1, green fluorescent protein (GFP) was fused to the Cterminus of CLIK-1 at the endogenous clik-1 locus using CRISPR/Cas9-mediated genome editing. CLIK-1-GFP was strongly expressed in the body wall muscle (Fig. 2A, C, E) and myoepithelial sheath of the somatic gonad (Fig. 2B, D, F), where CLIK-1-GFP localized in a sarcomeric pattern in the body wall muscle (Fig. 2A) and in a filamentous pattern in the myoepithelial sheath (Fig. 2B). Our preliminary study indicated that CLIK-1 co-localized with actin filaments (our unpublished observastions). However, we did not detect expression of CLIK-1-GFP in the pharynx, although we did detect expression of CLIK-1-GFP in the neurons surrounding the pharynx (our unpublished observations). Thus, our results disagree with the proteomics study reporting enrichment of CLIK-1 in the pharynx (30). Body wall muscle and myoepithelial sheath of the gonad are tissues where UNC-87 is also expressed and associated with actin filaments (21,24,25), suggesting that CLIK-1 and UNC-87 regulate actin filaments in the same cells. Additional characterization of tissue distribution and subcellular localization of CLIK-1-GFP will be reported elsewhere.

In a recent study, Wang et al. reported that *clik-1* gene knockout did not cause any obvious phenotypes (31), and we confirmed their observations using a separately isolated clik-1-null strain [clik-1(ok2355)] that contains a 1-kb deletion in the *clik-1* gene (our unpublished observation). However, we found that knockdown of *clik-1* by RNA interference (RNAi) in an unc-87 null mutant [unc-87(e1459)] (21,24) caused sterility with severe cytoskeletal defects in the reproductive system (Fig. 3 and 4). When worms were treated with control RNAi, wild-type worms actively moved and produced many progeny (Fig. 3A and I), whereas unc-87(e1459) worms were slowmoving and produced much less progeny (Fig. 3C and I). RNA interference of *clik-1* did not affect movement and fecundity in wild-type

background (Fig. 3B and I), but enhanced impaired worm motility to nearly paralysis (Fig. 3D and I) and caused sterility in *unc*-87(e1459) (Fig. 3D, see the absence of eggs and small worms on the culture plate). In the body wall muscle, actin was organized into a striated sarcomeric pattern in wild-type with control RNAi (Fig. 3E) or *clik-1(RNAi)* (Fig. 3F). Sarcomeric actin in unc-87(e1459) with control RNAi was disorganized with accumulation in aggregates (Fig. 3G), and the actin disorganization was not apparently enhanced by clik-1(RNAi) in unc-87(e1459) (Fig. 3H), suggesting that enhancement of paralysis by clik-1(RNAi) is not due to structural defects in the sarcomeres. Currently, we do not know why motility of unc-87(e1459); clik-1(RNAi) worms are severely impaired.

Simultaneous depletion of unc-87 and clik-1 disrupted cytoskeletal integrity in the somatic gonads, which explains why worm reproduction was strongly impaired (Fig. 4). In *C. elegans* hermaphrodites, oocytes are surrounded by the myoepithelial sheath in the gonad that provides contractile forces during ovulation of mature oocytes (32,33). The myoepithelial sheath is a smooth-muscle-like somatic tissue that contains non-striated actomyosin networks (34-36). The MYO-3 myosin heavy chain was used as a marker for the myoepithelial sheath (34,36,37). In wild-type with control RNAi, the myoepithelial sheath contained networks of actin and MYO-3 myosin and covered multiple oocytes (as shown by clusters of condensed chromosomes) (Fig. 4A-D). RNAi of *clik-1* in wild-type did not cause alteration in the actomyosin networks in the myoepithelial sheath (Fig. 4E-H). In unc-87(e1459) with control RNAi, actin filaments were somewhat disorganized, and the myoepithelial sheath was not as extended as wild-type with control RNAi (Fig. 4I-L and Q). Strikingly, in unc-87(e1459) with clik-1(RNAi), actin filaments were only sparsely distributed, and the myoepithelial sheath was shortened and nearly collapsed to one side of the gonad (Fig.

4M-P and Q). The oocytes in these worms contained large accumulations of DNA (Fig. 4O, arrow) as a result of endomitotic DNA replication during ovulation failure (38). These phenotypes indicate that UNC-87 and CLIK-1 have partially redundant function to maintain actin cytoskeletal integrity in the myoepithelial sheath and to execute proper ovulation that is an essential process in worm reproduction.

# CLIK-1 competes with UNC-87 for actin filament binding and inhibits filament bundling

To characterize biochemical properties of CLIK-1 and compare them with those of UNC-87, we prepared recombinant CLIK-1 protein with no extra tag sequence (Fig. 5A). When CLIK-1 was incubated with filamentous (F-) actin, CLIK-1 co-sedimented with F-actin after ultracentrifugation (Fig. 5B) indicating that CLIK-1 bound to F-actin. Quantitative analysis of the F-actin co-sedimentation assays at various CLIK-1 concentrations demonstrated that CLIK-1 bound to F-actin with a dissociation constant (Kd) of  $0.51 \pm 0.094 \ \mu M \ (n = 3)$ , which is higher than the estimated Kd ( $\sim 0.15 \mu$ M) for UNC-87 binding to F-actin (25). Binding was saturated at ~0.25 (mol CLIK-1/ mol actin) indicating that CLIK-1 binds to actin filaments with a stoichiometry of 1:4 (Fig. 5C). Under low-speed centrifugation (18,000 x g, 10 min), CLIK-1 did not increase actin in the pellets (Fig. 5B and D), whereas UNC-87 increased actin in the pellets by bundling filaments (22) (Fig. 5D). Therefore, CLIK-1 binds to actin filaments but, unlike UNC-87, lacks filament bundling activity.

Because CLIK-1 and UNC-87 localize to actin filaments in the same cells, we examined their functional relationship when the two proteins are both present. Not surprisingly from their sequence similarity, CLIK-1 and UNC-87 competed for F-actin binding (Fig. 6A-F). In the F-actin co-sedimentation assays at high speed (200,000 x g, 20 min) in the presence of a constant UNC-87 concentration, increasing concentrations of CLIK-1 reduced UNC-87 in

the pellets (Fig. 6A and B). In the opposite experiments with a constant CLIK-1 concentration, increasing concentrations of UNC-87 reduced CLIK-1 in the pellets (Fig. 6C and D). UNC-87 was more resistant to dissociation than CLIK-1 and more effective to dissociate the counterpart than CLIK-1, which is consistent with the higher affinity of UNC-87 with actin than that of CLIK-1. Interestingly, the competition between UNC-87 and CLIK-1 resulted in antagonistic modulation of actin filament bundling. UNC-87 bundled actin filaments and increased actin in the pellets under low-speed centrifugation (Fig. 6E and F). However, increasing concentrations of CLIK-1 decreased actin in the pellets even in the presence of UNC-87 (Fig. 6E and F). Their antagonistic effects on actin bundling were further demonstrated by direct observation of fluorescently labeled actin filaments. Actin filament bundles were formed in the presence of UNC-87 (Fig. 6, compare G and H), but not CLIK-1 (Fig. 6I). When both UNC-87  $(1 \mu M)$ and CLIK-1 (2 µM) were present, very few actin bundles were formed (Fig. 6J and K). The concentration of CLIK-1 needed to be higher than that of UNC-87 to cause significant reduction of actin bundles (Fig. 6K). Thus, the balance of UNC-87 and CLIK-1 can determine the extent of actin filament bundling in vitro.

# Tropomyosin binds to actin filaments simultaneously with CLIK-1 or UNC-87 and partially inhibits filament bundling

Although the biochemical experiments suggest that CLIK-1 is an inhibitor of actin filament bundling, we did not detect excessive actin filament bundling in the body wall muscle or myoepithelial sheath of *clik-1(RNAi)* worms or *clik-1* knockout worms (our unpublished observation). Therefore, we reasoned that other actin binding protein(s) can inhibit actin filament bundling by UNC-87. Since tropomyosin is a major F-actin binding protein, we examined how tropomyosin interacts with

actin filaments in the presence of UNC-87 or CLIK-1. The *lev-11* gene encodes at least six tropomyosin isoforms, and LEV-11A is a major high-molecular-weight isoform in muscle cells (39-42). LEV-11A bound to actin filaments in the presence of either UNC-87 or CLIK-1 (Fig. 7A-C). In addition, in the presence of a saturating concentration of LEV-11A, actin bundling, as determined by low-speed actin sedimentation, was inhibited when the UNC-87 concentrations were low (Fig. 7D-F). At high UNC-87 concentrations, actin filament bundling was no longer inhibited by LEV-11A (Fig. 7D-F). Therefore, tropomyosin, in addition to CLIK-1, can also inhibit UNC-87-induced actin filament bundling under certain conditions.

# CLIK-1 and UNC-87 prevent ADF/cofilin from F-actin binding

UNC-87 stabilizes actin filaments by preventing biding of actin depolymerizing factor (ADF)/cofilin to the filaments (25). Similarly, CLIK-1 inhibited ADF/cofilin binding to actin filaments. UNC-60B, a muscle-specific ADF/cofilin isoform in C. elegans, severs actin filaments, and UNC-60B-bound actin filaments can be sedimented by ultracentrifugation (43-46). UNC-60B, which co-sedimented with Factin, decreased when increasing concentrations of UNC-87 (Fig. 8A and C) or CLIK-1 (Fig. 8B and C) were included. UNC-87 was more efficient to prevent actin binding of UNC-60B than CLIK-1 (Fig. 8C), which is consistent with the difference in their affinity with actin. Thus, UNC-87 and CLIK-1 share a common actinstabilizing function by antagonizing ADF/cofilin.

# Discussion

Here, we demonstrate that two calponinrelated actin binding proteins, UNC-87 and CLIK-1, are critical for cytoskeletal integrity in the *C. elegans* reproductive system. Depletion of both UNC-87 and CLIK-1 cause severe

disorganization of actin filaments in the myoepithelial sheath of the somatic gonad, which results in sterility. Single depletion of either UNC-87 or CLIK-1 caused much milder or no phenotypes, indicating that UNC-87 and CLIK-1 have partially redundant functions in *vivo*. A previous single-cell proteomic study reported that CLIK-1 is enriched in the pharyngeal muscle cells (30). However, we do not detect strong expression of CLIK-1 in the pharynx. Instead, CLIK-1 expression is strong in other contractile tissues, including body wall muscle and myoepithelial sheath and overlaps with UNC-87 expression. Interestingly, biochemical analysis reveals both common and distinct actin-regulatory properties of UNC-87 and CLIK-1. Both proteins bind to actin filaments and prevent binding of ADF/cofilin to actin filaments. However, UNC-87 bundles actin filaments, but CLIK-1 does not. Their competitive binding to actin filaments modulates the extent of filament bundling when the two proteins co-exist. Thus, our current study uncovers novel functional aspects of calponinrelated proteins in the regulation of the actin cytoskeleton.

The myoepithelial sheath of the somatic gonad (ovary) is highly sensitive to depletion of UNC-87 and CLIK-1. The myoepithelial sheath is a monolayer of smooth-muscle-like cells that contain non-striated contractile actin networks (34-36). Its contraction is tightly coupled with oocyte maturation and pushes only one mature oocyte out of the ovary for subsequent fertilization (32,33). Contraction of the myoepithelial sheath is regulated by both troponin-tropomyosin complex (47-49) and phosphorylation of myosin light chain (37). UNC-87 has an inhibitory effect on actomyosin contractility, and may facilitate relaxation of the myoepithelial sheath (24). Severe disorganization of the actin filaments in the myoepithelial sheath in double depletion of UNC-87 and CLIK-1 suggest that the two calponin-related proteins have redundant roles to stabilize actin filaments. CLIK-1 shares a

common function with UNC-87 to protect actin filaments from ADF/cofilin (Fig. 8). ADF/cofilin and actin-interacting protein 1 are required for assembly of actin networks in the myoepithelial sheath (50,51). However, excessive activities of these proteins are expected to destabilize actin filaments. Either one of UNC-87 or CLIK-1 is sufficient to stabilize actin filaments, but loss of both proteins can cause severe actin disorganization such that the myoepithelial sheath is no longer functional. In addition, ovulatory contraction of the myoepithelial sheath involves extensive morphological changes and mechanical stress (32,33), and these calponin-related proteins may have a role in providing mechanical stability to the actin filaments as demonstrated biophysically for mammalian calponin (52,53).

To our knowledge, CLIK-1 is the only calponin-related protein lacking actin filament bundling activity. Because only a limited number of calponin-related proteins have been characterized biochemically, there may be other calponin-related proteins with variable functions. Vertebrate calponin bundles actin filaments in vitro (54,55) and play important roles in cytoskeletal structures containing actin bundles (10,56). Vertebrate SM22/transgelin (57,58) and yeast calponin-related proteins (12-14,59) have only one CLIK motif and still bundles actin filaments in vitro. A calponinrelated protein with no actin bundling activity, such as CLIK-1 or an equivalent protein, may compete with conventional calponin and/or SM22/transgelin to modulate the extent of actin bundle formation. Although such an antagonistic relationship was not clearly detected between UNC-87 and CLIK-1 in the body wall muscle and myoepithelial sheath in vivo, it may be of functional significance in other cell types, or other actin-binding proteins, such as tropomyosin, may participate in the modulation of actin bundle formation.

The differences in the actin-regulatory activities of UNC-87 and CLIK-1 suggest that not all CLIK motifs behave in the same manner.

Previous studies suggested that CLIK motifs are functionally similar modular motifs because the number of CLIK motifs correlates with the effectiveness in stabilizing actin filaments in cultured cells (10,23). The difference between UNC-87 and CLIK-1 in actin bundling activity indicate that some of the CLIK motifs in UNC-87 specifically mediate actin bundling. Since UNC-87 is monomeric (22), a single UNC-87 molecule needs to bind to multiple actin filaments to bundle them, suggesting that some of the CLIK motifs preferentially bind to a second filament. By contrast, CLIK-1 does not bundle actin filaments. Therefore, all the CLIK motifs in CLIK-1 bind to the same filament. There may be a difference in cooperativity among actin binding sites. In UNC-87, a first actin-binding site may prevent a second actinbinding site from binding to the same filament, while, in CLIK-1, a first actin-binding site may promote binding of a second actin-binding site to the same filament. Alternatively, the structure of UNC-87 may be suited to cross-bridge multiple filaments, whereas that of CLIK-1 may be optimal to bind to a single filament. Further dissection of biochemical properties of CLIK motifs should inform us about how different CLIK motifs contribute to functional differentiation of calponin-related proteins.

## **Experimental Procedures**

#### C. elegans strains and culture

Worms were cultured following standard methods (60) . The following strains were obtained from the *Caenorhabditis* Genetics Center and used in this study: N2 wild type, CB1459 *unc-87(e1459)*, and RB1820 *clik-1(ok2355)*. *clik-1(ok2355)* was outcrossed three times and used in this study (outcrossed strain: ON225).

#### CRISPR/Cas9 genome editing

The GFP sequence was inserted in the genome in-frame at the 3'-end of the *clik-1* coding region by CRISPR/Cas9-mediated

genome editing using vectors and protocols described by Dickinson et al. (61). A single guide (sg) RNA target sequence (GATGTAATTACCACATACGA) was cloned into pDD162 (Addgene Plasmid # 47549) for expression of both sgRNA and Cas9 nuclease. Homology arms of ~500 bp each were fused with the GFP::self-excising cassette region from pDD282 (Addgene Plasmid #66823) by fusion PCR using Q5 High-Fidelity DNA Polymerase (New England Biolabs) and used as a homologous repair template. Hygromycinresistant worms with a roller phenotype were isolated as knock-in worms. They were treated at 34 °C for 4 hours to induce excision of the self-excising cassette. In the next generation, non-roller GFP-positive worms were isolated, and three independent *clik-1::GFP* strains, ON352 clik-1(kt1 [clik-1::gfp]), ON353 clik-1(kt2 [clik-1::gfp]), and ON354 clik-1(kt3 [clik-1::gfp]), were established.

# **RNA** interference experiments

RNA interference experiments were performed by feeding with *Escherichia coli* HT115(DE3) expressing double-stranded RNA as described previously (62). The RNAi experiments were started by treating L1 larvae, and phenotypes were observed when they grew to adult worms after three days. Control RNAi experiments were performed using the unmodified L4440 plasmid vector (kindly provided by Andrew Fire, Stanford University) (63). An RNAi clone for *clik-1* (V-4N16) was obtained from Source BioScience. Images of worms on agar plates were captured using a Nikon Coolpix 995 digital camera mounted on a Zeiss Stemi 2000 stereo microscope.

#### Fluorescence microscopy

Live worms were anesthetized in M9 buffer containing 0.1% tricaine and 0.01% tetramisole for 30 min, mounted on 2% agarose pads, and observed by epifluorescence and differential interference contrast optics using a Nikon Eclipse TE2000 inverted microscope (Nikon Instruments, Tokyo, Japan).

Staining of whole worms with tetramethylrhodamine–phalloidin (Sigma-Aldrich) was performed as described previously (64).

Gonads were dissected from adult hermaphrodites on polylysine-coated glass slides as described previously (36). They were fixed by methanol at  $-20^{\circ}$ C for 5 min and washed by phosphate-buffered saline (PBS) and primary antibodies in PBS containing 1% BSA. After being washed with PBS, they were treated with fluorophore-labeled secondary antibodies, followed by washing with PBS.

Primary antibodies used were rabbit anti-actin polyclonal (Cytoskeleton, Denver, CO) and mouse anti–MYO-3 monoclonal (5-6; ref (65)) antibodies. Secondary antibodies used were Alexa 488–labeled goat anti-mouse IgG from Life Technologies and Cy3-labeled goat anti-mouse IgG from Jackson ImmunoResearch (West Grove, PA).

Fixed samples were mounted with ProLong Gold (Life Technologies) and observed by epifluorescence using a Nikon Eclipse TE2000 inverted microscope with a CFI Plan Fluor ELWD 40× (dry; NA 0.60) or Plan Apo  $60 \times$  (oil; NA 1.40) objective. Images were captured by a SPOT RT monochrome chargecoupled device camera (Diagnostic Instruments, Sterling Heights, MI) and processed by IPLab imaging software (BD Biosciences) and Photoshop CS3 (Adobe, San Jose, CA) or by ORCA Flash 4.0 LT monochrome scientific complementary metal-oxide-semiconductor camera (Hamamatsu Photonics, Shizuoka, Japan) and processed by Nikon NIS-Elements and Adobe Photoshop CS3. Quantification of the myoepithelial sheath lengths was performed using ImageJ.

# **Protein preparation**

Actin was prepared from rabbit muscle acetone powder (Pel-Freeze Biologicals) as described by Pardee and Spudich (66). Recombinant LEV-11A (tropomyosin) (41), UNC-87B (22), and UNC-60B (ADF/cofilin) (45) were expressed in *Escherichia coli* and purified as described previously. Unstained molecular weight markers (Nacalai USA, catalog number 29458-24) were used in SDS-PAGE.

The open reading frame of *clik-1* cDNA was amplified with reverse-transcriptase-PCR and cloned at the NdeI - BamHI sites of pET-3a (Novagen) with no extra tag sequence. E. coli BL21(DE3) pLysS was transformed with the expression vector, cultured in M9ZB medium containing 50 µg/ml ampicillin and 34 µg/ml chloramphenicol at 37 °C until A600 reached 0.6  $cm^{-1}$ . Then, protein expression was induced by adding 0.1 mM isopropyl β-dthiogalactopyranoside for 3 hr at 37 °C. The cells were harvested by centrifugation at 5000  $\times$ g for 10 min and disrupted by sonication in a buffer containing 0.1 M KCl, 1 mM EDTA, 20 mM Tris-HCl (pH 7.5), 0.2 mM dithiothreitol (DTT) and 1 mM phenylmethanesulfonyl fluoride. The homogenates were centrifuged at  $25,000 \times g$  for 30 min, and the supernatants were fractionated by 35 % saturated ammonium sulfate (194 g/l). Insoluble proteins were separated by centrifugation at 20,000 x g for 20 min, dissolved in a small volume of buffer A (0.1 M NaCl, 0.2 mM DTT, and 20 mM Tris-HCl, pH 7.5), and dialyzed overnight against buffer A. The dialyzed proteins were cleared by centrifugation at 20,000 x g for 20 min, applied to a Mono-Q 4.6/100PE column (GE Healthcare), and eluted with a linear gradient of NaCl from 0.1 to 0.5 M. Fractions containing CLIK-1 were applied to HiLoad 16/60 Superdex 75 (GE Healthcare) and eluted with a buffer containing 0.1 M KCl, 0.2 mM DTT, 20 mM HEPES-KOH, pH 7.5. Pure CLIK-1 fractions were pooled and concentrated by a MacroSep Centrifugal Device (molecular weight cutoff at 10K) (Pall Corporation), and stored at -80 °C. Protein concentrations were determined by a BCA Protein Assay kit (Thermo Scientific).

## F-actin sedimentation assays

Filamentous actin (5 or  $10 \mu$ M) was incubated with proteins of interest in F-buffer (0.1 M KCl, 2 mM MgCl<sub>2</sub>, 20 mM HEPES-KOH, pH 7.5, 1 mM DTT) at room temperature. To examine F-actin binding of the proteins of interest by high-speed co-sedimentation, the samples were ultracentrifuged at 42,000 rpm (200,000 xg) for 20 min using a Beckman 42.2Ti rotor. To examine F-actin bundling by low-speed sedimentation, the samples were centrifuged at 15,000 rpm (18,000 x g) for 10 min using a Hettich MIKRO 200 centrifuge. Supernatants and pellets were separated, adjusted to the same volumes, and examined by SDS-PAGE (12% acrylamide gel). For quantitative analysis of high-speed F-actin cosedimentation, control experiments were performed without actin, and non-specific sedimentation of CLIK-1 was determined and subtracted from sedimentation of CLIK-1 in the presence of actin. The gels were stained with Coomassie Brilliant Blue R-250 (National Diagnostics) and scanned by an Epson Perfection V700 scanner at 300 dpi. Band intensity was quantified with ImageJ.

## Microscopic observation of actin bundles

Direct observation of actin bundles by fluorescent microscopy was performed as described previously (24,67) using DyLight 549labeled actin (68). Briefly, DyLight 549-labeled

actin filaments (2 µM actin, 20 % labeled) were incubated with proteins of interest for 5 min at room temperature in F-buffer, and 2 µl of the samples were applied to nitrocellulose-coated coverslips (22 x 40 mm) and mounted with noncoated coverslips (18 x 18 mm). They were immediately observed with epifluorescence using a Nikon Eclipse TE2000 inverted microscope with a Plan Apo  $60 \times$  (oil; NA 1.40) objective. Images were captured by a SPOT RT monochrome charge-coupled device camera (Diagnostic Instruments, Sterling Heights, MI) and processed by IPLab imaging software (BD Biosciences) and Photoshop CS3 (Adobe, San Jose, CA). For quantification of actin bundles, regions of interest of 12.7 µm x 12.7 µm (100 x 100 pixels) were randomly selected. Then, using ImageJ, a threshold was set to eliminate single filaments, and number of pixels above the threshold regardless the intensity was counted.

# Statistical analysis

Data in Fig. 3I, 4Q, and 6K were analyzed by one-way analysis of variance with the Holm-Sidak method of pairwise multiple comparison procedures using SigmaPlot v13.0 (Systat Software, San Jose, CA).

# Data availability

All data are contained in the manuscript. Raw data are available from S. O. upon request.

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This paper is dedicated to the memory of Kanako Ono who passed away during the preparation of this manuscript.

# Author contribution

S. O. conceived the project and wrote the paper. S. O. and K. O. performed experiments, analyzed data, and prepared figures.

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# Figure legends

**Figure 1. Domain organizations of calponin and calponin-related proteins.** (A) Schematic representation of domain organizations of vertebrate calponin (CNN1, calponin 1; CNN2, calponin 2; CNN3, calponin 3), vertebrate SM22/transgelin (TGN1, SM22α/transgelin 1; TGN2, transgelin 2; TGN3, transgelin 3), *C. elegans* UNC-87, and *C. elegans* CLIK-1. CH: calponin-homology domain; CLIK:

calponin-like motif. (B) Sequence alignment of CLIK motifs from calponin and calponin-related proteins. Human calponin 1 (CNN1) (NCBI Reference Sequence: NM\_001299); human SM22α/transgelin (TGN1) (NCBI Reference Sequence: NM\_001001522); *Drosophila melanogaster* MP20 (NCBI Reference Sequence: NM\_057295); *C. elegans* UNC-87 (NCBI Reference Sequence: NM\_001025922); *C. elegans* CLIK-1 (GenBank accession number: CCD74240).

**Figure 2. Expression and localization of CLIK-1 in** *C. elegans.* GFP was fused to the C-terminus of CLIK-1 using CRISPR/Cas9-mediated genome editing. Localization of CLIK-1-GFP (A, B) and differential interference contrast images (C, D) were examined in live adult hermaphrodites. Merged images are shown in E and F. Expression of CLIK-1-GFP in the body wall muscle (A, C, E) and the myoepithelial sheath of the somatic gonad (B, D, F) are highlighted.

**Figure 3. Simultaneous depletion of** *unc-87* and *clik-1* causes impaired worm motility and sterility. Wild-type or *unc-87(e1459)* worms were treated with control RNAi or *clik-1(RNAi)*. (A – D) Live worms on the culture plates are shown. Large worms are the RNAi-treated worms. Small worms and eggs are F1 progeny from the treated worms. Bar, 1.0 mm. (E-H) Organization of actin filaments in the body wall muscle as observed by staining with tetramethylrhodamine-phalloidin. Bar, 10 µm. (I) Worm motility was quantified as number of beating per 30 sec. n = 10. Boxes represent the range of the 25th and 75th percentiles, with the medians marked by solid horizontal lines, and whiskers indicate the 10th and 90th percentiles. NS, not significant. \*\*\*, p < 0.001.

Figure 4. Simultaneous depletion of *unc-87* and *clik-1* causes severe disorganization of actin filaments in the somatic gonad. Wild-type or *unc-87(e1459)* worms were treated with control RNAi or *clik-1(RNAi)*. (A-P) Dissected gonads were stained for F-actin (A, E, I, M), MYO-3 (myosin) (B, F, J, N), and DNA (C, G, K, O). The proximal side of the gonads are oriented to the right. Merged imaged (F-actin in red, MYO-3 in green, and DNA in blue) are shown in the right column (D, H, L, P). Bar, 50 µm. (Q) Lengths of the myoepithelial sheath as marked by the MYO-3 staining were measured. Sample sizes (n) were 29 (wild-type; control RNAi), 28 (wild-type; *clik-1(RNAi)*), 26 (*unc-87(e1459);* control RNAi), and 27 (*unc-87(e1459); clik-1(RNAi)*). Boxes represent the range of the 25th and 75th percentiles, with the medians marked by solid horizontal lines, and whiskers indicate the 10th and 90th percentiles. NS, not significant. \*\*\*, p < 0.001.

Figure 5. CLIK-1 binds to actin filaments without bundling them. (A) Purity of recombinant UNC-87  $(0.5 \ \mu g)$ , recombinant CLIK-1  $(0.5 \ \mu g)$ , and rabbit muscle actin  $(0.2 \ \mu g)$ , was examined by SDS-PAGE (12 % acrylamide gel). Molecular weight markers in kDa are shown on the left. (B) F-actin sedimentation assays at high speed (200,000 x g, 20 min) (top) and low speed (18,000 x g, 10 min) (bottom). F-actin (10  $\mu$ M) was incubated with 0 – 2  $\mu$ M CLIK-1 for 1 hr and centrifuged at the indicated conditions. Supernatants (s) and pellets (p) were separated and examined by SDS-PAGE. Positions of CLIK-1 (C) and actin (A) are indicated on the right. (C) Quantitative analysis of the high-speed F-actin sedimentation assays. The high-speed F-actin sedimentation assays were performed using 5  $\mu$ M F-actin and 0 – 10  $\mu$ M CLIK-1. Molar ratios of CLIK-1 to actin in the pellets (actin-dependent sedimentation of CLIK-1 was determined by subtracting non-specific sedimentation of CLIK-1) [bound CLIK-1 (µM)/actin (µM)] are plotted as a function of free CLIK-1. A dissociation constant (Kd) was determined from three independent experiments (means  $\pm$  standard deviation). (D) Quantitative analysis of the low-speed F-actin sedimentation assays. The low-speed F-actin sedimentation assays were performed with 10 µM F-actin and  $0 - 10 \mu M$  UNC-87 or CLIK-1. Percentages of actin in the pellets (bundled actin filaments) were plotted as a function of concentration of UNC-87 or CLIK-1. Data are means ± standard deviation from three independent experiments.

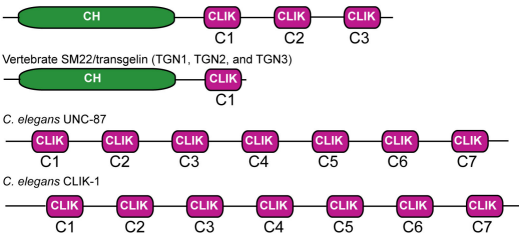
Figure 6. Competitive binding of UNC-87 and CLIK-1 to actin filaments. (A, B) F-actin (10 µM) was incubated with 1  $\mu$ M UNC-87 and 0 – 10  $\mu$ M CLIK-1 and examined by high-speed co-sedimentation assays. Supernatants and pellets were analyzed by SDS-PAGE (A). Relative amounts of UNC-87 in the pellets (100 % in the absence of CLIK-1) were quantified and plotted as a function of CLIK-1 concentration (B). Data are means  $\pm$  standard deviation from three independent experiments. (C, D) Factin (10  $\mu$ M) was incubated with 2  $\mu$ M CLIK-1 and 0 – 5  $\mu$ M UNC-87 and examined by high-speed cosedimentation assays. Supernatants and pellets were analyzed by SDS-PAGE (C). Relative amounts of CLIK-1 in the pellets (100 % in the absence of UNC-87) were quantified and plotted as a function of UNC-87 concentration (D). Data are means  $\pm$  standard deviation from three independent experiments. (E, F) F-actin (10  $\mu$ M) was incubated with 2.5  $\mu$ M UNC-87 and 0 – 10  $\mu$ M CLIK-1 and examined by lowspeed sedimentation assays. Supernatants (s) and pellets (p) were analyzed by SDS-PAGE (E). Percentages of actin in the pellets (bundled actin filaments) were plotted as a function of CLIK-1 concentration. Data are means  $\pm$  standard deviation from four independent experiments. (G – K) Direct observation of actin bundling by fluorescence microscopy. DyLight 549-labeled actin (2  $\mu$ M) was incubated with buffer only (G) or buffer with 1 µM UNC-87 (H), 2 µM CLIK-1 (I), or 1 µM UNC-87 and 2 μM CLIK-1 (J) and observed with fluorescence microscopy. Bar, 10 μm. Actin bundle formation was quantified (K) from randomly selected regions of interest (100 x 100 pixels) (n = 25). Boxes represent the range of the 25th and 75th percentiles, with the medians marked by solid horizontal lines, and whiskers indicate the 10th and 90th percentiles. NS, not significant. \*, 0.01 ; \*\*, <math>0.001 ; \*\*\*, p< 0.001.

Figure 7. Tropomyosin binds to F-actin in the presence of UNC-87 or CLIK-1 and partially inhibits UNC-87-mediated actin filament bundling. (A - C) F-actin (10 µM) was incubated with 1 µM LEV-11A (tropomyosin) and 0 – 10 µM UNC-87 (A) or CLIK-1 (B) and examined by high-speed cosedimentation assays. Supernatants and pellets were analyzed by SDS-PAGE. Relative amounts of LEV-11A in the pellets (100 % in the absence of UNC-87 or CLIK-1) were quantified and plotted as a function of UNC-87 or CLIK-1 concentration (C). Data are means ± standard deviation from three independent experiments. (D – F) F-actin (10 µM) was incubated with 0 - 10 µM UNC-87 in the absence (D) or presence of 2 µM LEV-11A (E) and examined by low-speed sedimentation assays. Supernatants (s) and pellets (p) were analyzed by SDS-PAGE. Percentages of actin in the pellets (bundled actin filaments) were plotted as a function of UNC-87 concentration. Data are means ± standard deviation from three independent side pendent experiments.

**Figure 8. UNC-87 and CLIK-1 inhibit F-actin binding of ADF/cofilin.** F-actin (10  $\mu$ M) was incubated with 10  $\mu$ M UNC-60B (ADF/cofilin) and 0 – 5  $\mu$ M UNC-87 (A) or CLIK-1 (B) and examined by high-speed co-sedimentation assays. Supernatants and pellets were analyzed by SDS-PAGE. Relative amounts of UNC-60B in the pellets (100 % in the absence of UNC-87 or CLIK-1) were quantified and plotted as a function of UNC-87 or CLIK-1 concentration (C). Data are means ± standard deviation from three independent experiments.

# Α

Vertebrate calponin (CNN1, CNN2, and CNN3)

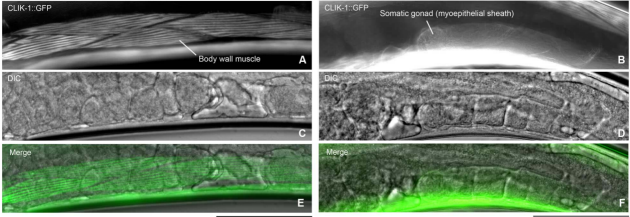


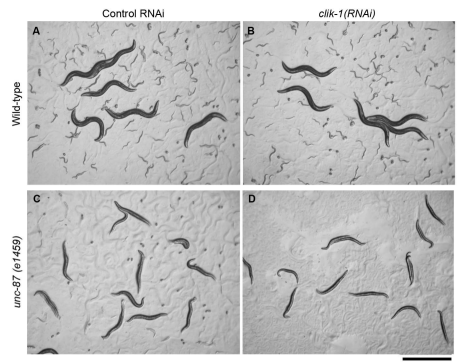
В

HUMAN CNN1	C1	IGLQMGTNKFASQQGMTAYGTRR
HUMAN CNN1	C2	ISLQMGTNKGASQAGMTAPGTKR
HUMAN CNN1	C3	VSLQMGSNKGASQRGMTVYGLPR
HUMAN TGN1	C1	IGLQMGSNRGASQAGMTGYGRPR
DM MP20	C1	VG <mark>LQAG</mark> S <mark>NKGATQAG</mark> Q-NL <mark>G</mark> AGR
CE UNC-87	C1	IPS <mark>QAGWNK</mark> GD <mark>SQ</mark> KL <mark>MTNFG</mark> TPR
CE UNC-87	C2	VRLQSGTNKYCSQRGMTGFGSGR
CE UNC-87	C3	VRLQAGTNKYDSQKGMTGFGTGR
CE UNC-87	C4	IPLQSGTNKFASQKGMTGFGTAR
CE UNC-87	C5	IPSQMGSNQYASQKGMTGFGQPR
CE UNC-87	C6	VRLQSGTNRFASQAGMIGFGTCR
CE UNC-87	C7	IPS <mark>QAGWNK</mark> GD <mark>SQ</mark> KK <mark>MTSFG</mark> APR
CE CLIK-1	C1	IPIQAGSNKYASQKGMTGFGVPR
CE CLIK-1	C2	TWLQSGTNKFASQKGQSGFGAVR
CE CLIK-1	C3	VPLQSGTNKLASQAGMTGIGMPR
CE CLIK-1	C4	IHLQMGTNKFANQAGMTGFGMPR
CE CLIK-1	С5	SRQTSGWKEG <mark>ASQ</mark> AGMTGFGAFR
CE CLIK-1	C6	IPY <mark>QMGVN</mark> FLE <mark>SQAGKTGFG</mark> QPR
CE CLIK-1	C7	FWTGQKEADH <mark>ANQTGMTAFG</mark> TP <mark>R</mark>

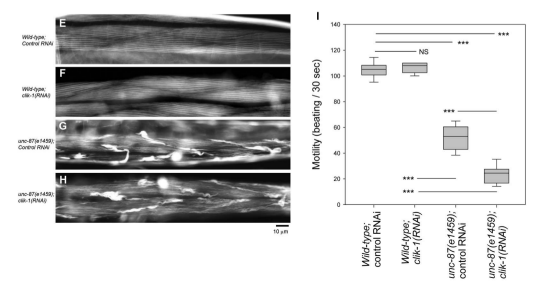
Consensus

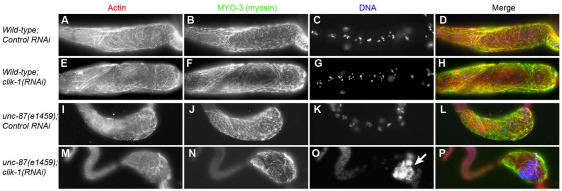
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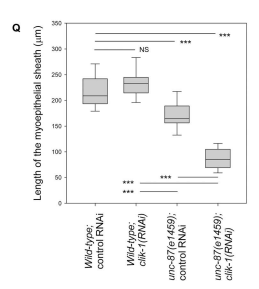


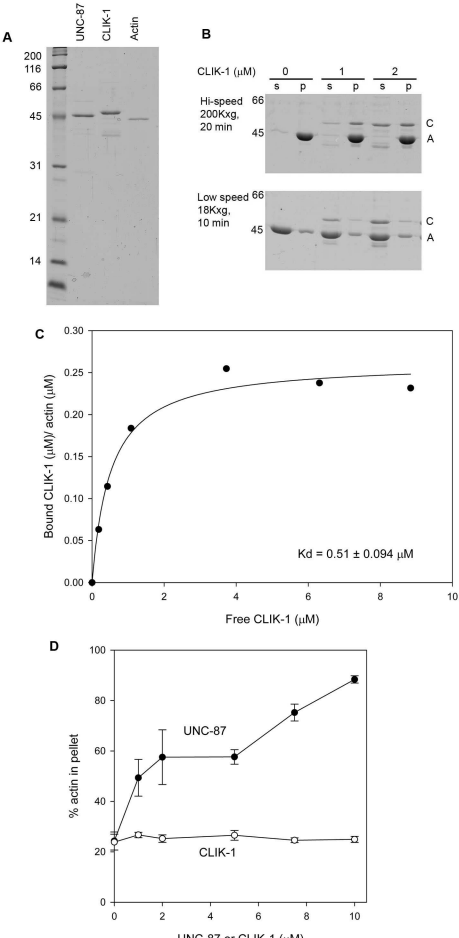






50 µm





UNC-87 or CLIK-1 (µM)

