
Small GTPases*

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*Edited by Iva Greenwald. Last revised February 23, 2016. Published in its final form August 16, 2018. This chapter should be cited as: Reiner D. J., Lundquist E. A. Small GTPases. (August 16, 2018), WormBook, ed. The *C. elegans* Research Community, WormBook, doi/10.1895/wormbook.1.67.2, <http://www.wormbook.org>.

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Abstract

Members of the protein superfamily of small guanosine triphosphatases, also known as small GTPases, small G-proteins, or the Ras superfamily, are involved in nearly every aspect of cell biology. Small GTPases are tightly regulated molecular switches that make binary on/off decisions through controlled loading of GTP (activation) and hydrolysis of GTP to GDP (inactivation). Small GTPases typically function as nodal points that integrate broad upstream regulatory inputs and disseminate broad effector outputs. The superfamily comprises five families that are conserved across eukaryotes: Ras, Rho, Rab, Arf, and Ran. Each family, besides Ran, has radiated functionally since our last common ancestor with fungi, and certain subfamilies persist throughout metazoa. The double genome duplication leading to vertebrates resulted in two to four genes for many subfamilies, plus some novel mammalian additions. Here we discuss general principles of small GTPase biology, survey the *C. elegans* complement of small GTPases and how they compare to their mammalian counterparts, and note atypical nematode members that do not fall into discrete subfamilies. We do not discuss the multitude of other proteins with catalytic guanosine triphosphatase domains that fall outside the small GTPase/Ras superfamily.

1. Overview

The Ras (Rat sarcoma) superfamily of small GTPases is an ancient group of molecular switches whose functions have radiated phylogenetically to encompass broad areas of cell biology. The founding members of the superfamily are the Ras proto-oncogenes, discovered in the early 1980s, which are conserved from yeast to *C. elegans*, *Drosophila*, and mammals. Ras itself, however, composes a tiny part of the overall signaling capabilities of the small GTPase superfamily. Members of the superfamily are key regulators of most processes in the cell, including differentiation, proliferation, vesicle and organelle dynamics and transport, nuclear dynamics, and regulation of the cytoskeleton. What they share in common is the GDP/GTP switch or, in some cases, just the structure of the GTPase domain, even if catalytically inactive.

In addition to being the founding member of the superfamily, Ras is also the reference protein of the Ras family of small GTPases—one of five families in the Ras superfamily: the others being Rho (Ras homology), Arf (ADP-ribosylation factor), Rab (Ras-like in brain), and Ran (Ras-like nuclear). (Figure 1). Ras family members generally participate in signal transduction, though, like many superfamily members, they perform functions outside of their stereotypical roles. Rho family members typically regulate cytoskeletal dynamics. The Rab family members generally control fusion and trafficking of vesicles and larger bodies. Functions of Arf family members partially overlap with those of Rabs, and are generally involved in vesicle biogenesis and recycling, intracellular trafficking, and cytoskeletal regulation. The single Ran controls nuclear translocation and structure.

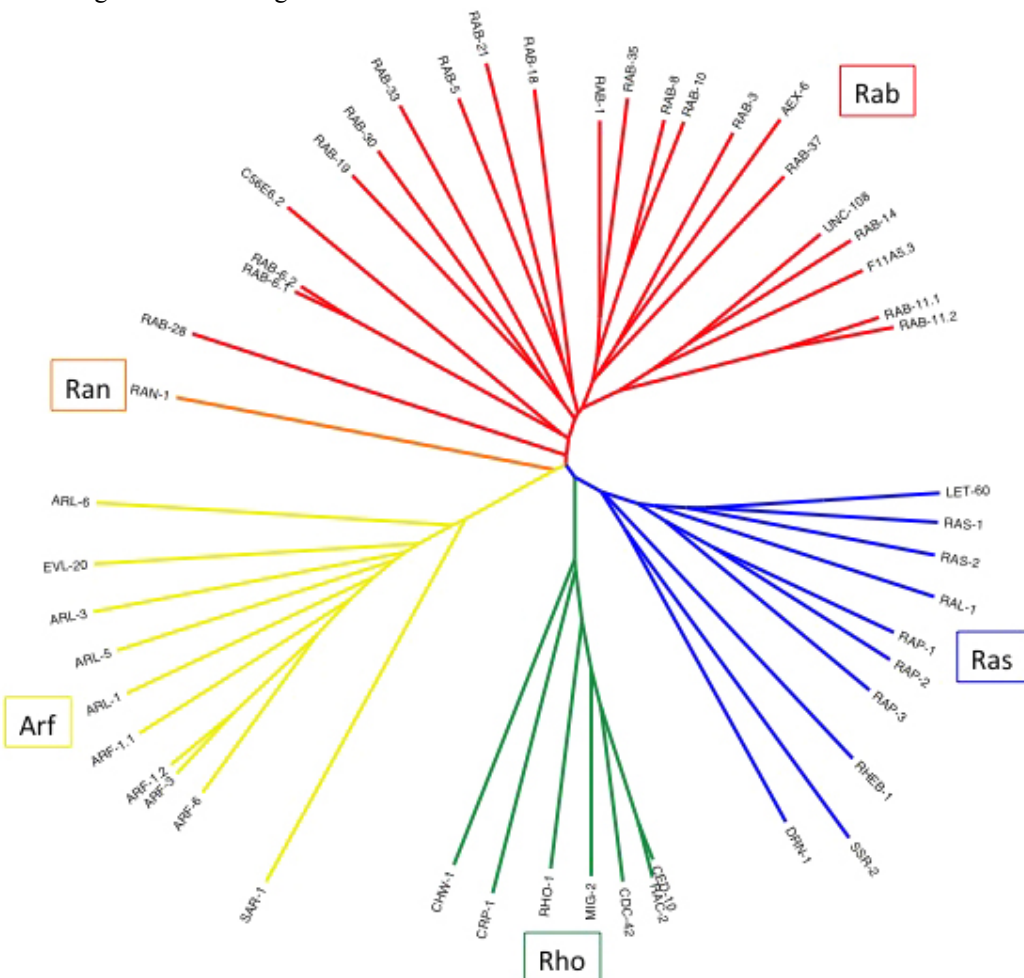


Figure 1. A dendrogram of the *C. elegans* small GTPases. The Ras superfamily consists of the Ras, Rho, Rab, and Arf families, with Ran comprising its family of one. The core GTPase domain is the unifying feature of this group, while N- and C-terminal extensions are much less conserved and could distort alignments. Thus only the core GTPase domain was used for the alignment and generation of this rooted tree (via Clustal X, with trees generated in FigTree).

Small GTPases cycle between two basic states: inactive GDP-bound (Ras•GDP), and active, GTP-bound (Ras•GTP) (Figure 2). Two major classes of regulators control the GTP binding state of small GTPases. Guanine nucleotide exchange factors (GEFs) eject GDP from Ras•GDP, and the much higher levels of free GTP than GDP in the cell favor GTP loading to produce Ras•GTP. The catalytic activity of intrinsic GTP hydrolysis is typically poor. But GTPase activating proteins (GAPs) stimulate GTP hydrolysis to convert the protein from Ras•GTP to Ras•GDP.

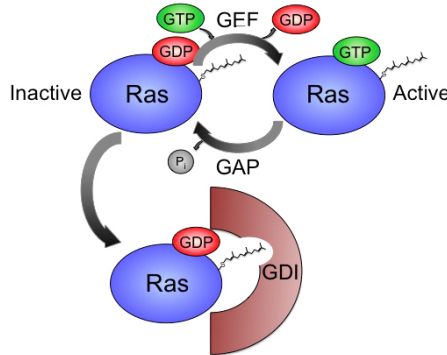


Figure 2. The small GTPase cycle. GEFs promote GTP loading to activate small GTPases, while GAPs stimulate intrinsic GTP hydrolysis to inactivate small GTPases. Guanine nucleotide dissociation inhibitors (GDIs) sequester GDP-bound, prenylated GTPases, and can sequester or chaperone them among plasma membrane and subcellular compartments.

GEFs and GAPs are multi-domain proteins that integrate diverse arrays of cellular inputs to confer precise spatiotemporal control of small GTPase activation (Bos et al., 2007). GEF catalytic domains vary widely; canonical types can be identified by homology, typically within GTPase families (e.g., *S. cerevisiae* CDC25 domains for the Ras family, DH-PH and DOCK domains for the Rho family, and DENN domains for the Rab family). Additionally, non-canonical GEFs are still being identified based on function rather than homology (Bos et al., 2007). GEF and GAP domains are typically unconserved between GTPase families. Distinctive GAP domains are specific to each GTPase family: RasGAP, RhoGAP, and TBC domains are diagnostic for the Ras, Rho, and Rab families, respectively (Barr and Lambright, 2010). GEFs and GAPs can be found with a wide diversity of accompanying domains, and it is thought that different GEFs and GAPs orchestrate activity of the same small GTPase in different tissues or in different subcellular compartments of the same tissue type. For example, 69 different mammalian DH-PH RhoGEFs and 11 DOCK-ELMO RhoGEFs have been identified to regulate 22 Rho family members (Rossman et al., 2005). The diversity of inputs to GEFs and GAPs encompasses all of cell biology—from direct binding by signaling proteins like heterotrimeric G-proteins, to second messengers like cAMP, to structural proteins (Bos et al., 2007).

The common structural and biochemical feature of the small GTPase superfamily is the approximately 170 residue G domain, which governs nucleotide binding, GTP hydrolysis, and effector interactions. The GTPase domain is defined by five G-boxes with certain structurally conserved residues across the superfamily: G1 (GxxxxGKS/T), G2 (T), G3 (DxxGQ/H/T), G4 (T/NKxD), and G5 (C/SAK/L/T) (Wennerberg et al., 2005) (Figure 3). Effector binding is governed by the Switch I and Switch II regions of the GTPase, which undergo a conformation change upon GTP binding, particularly the core effector-binding region in Switch I, around the G2 box (Holbrook and Kim, 1989). Additional N- and C-terminal extensions further regulate function, mostly through membrane association in defined subcellular compartments, typically the plasma membrane but extending to most organelles and vesicles.

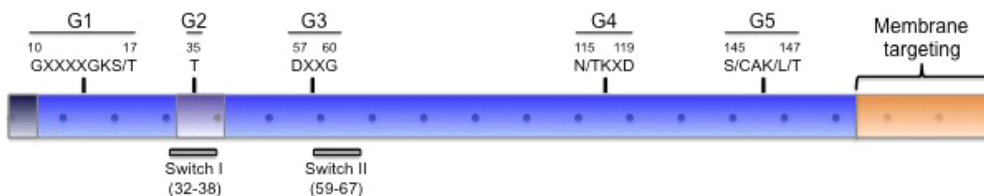


Figure 3. Functional domains of LET-60/Ras inferred from mammalian Ras. Blue indicates the G domain (also known as the GTPase domain; residues 5–166 in LET-60 and mammalian H/N/K-Ras), purple indicates the core effector-binding region that mediates direct interaction with effectors, and orange indicates the C-terminal hypervariable region and CAAX (HVR + CAAX) that is typically lipid modified and governs membrane localization. The five G motifs (also called G boxes) and consensus sequences among small GTPases are shown above the diagram with canonical *C. elegans* LET-60 and mammalian H/N/K-Ras numbering. Switch I and Switch II are regions whose conformation changes with the change from GDP loading (inactive) to GTP loading (active) states. Each dot represents 10 amino acids. This figure was adapted from (Wennerberg et al., 2005).

An ironic feature of small GTPases, which frequently confuses newcomers, is that the enzymatic function of small GTPases—GTP hydrolysis to generate GDP and free inorganic phosphate—inactivates the protein. Thus selectively altering the catalytic function of a small GTPase without disrupting protein structure locks the protein in the “on” state, while promoting enzymatic function hastens inactivation. Disruption of protein structure with strong loss-of-function or null mutations abrogates signaling.

The highly conserved structure of the Ras superfamily G domain has allowed the characterization and broad application of genetic tools developed through biochemical studies and mostly validated in mammalian cell-based assays (Wennerberg et al., 2005). Based on mutations found in oncogenic Ras (and following Ras numbering), substitutions at positions 12, 13, and 61 result in constitutive activity: canonical mutations are G12V, G13E, and Q61L, with G13E causing only moderate activation. S17N causes a dominant-negative GTPase; this reagent should be used with caution, since the S17N GTPase sequesters interacting GEFs, thus blocking their activation of shared substrates and potentially yielding misleading results. The F28L or “fast-cycling” mutation confers a GEF-insensitive but GAP-sensitive gain of function, which is particularly useful in generating intermediate-level activation, or for GTPases where GDP-GTP cycling is more important than the actual nucleotide-bound state (Reinstein et al., 1991). Variations in applicability occur across the superfamily, so utilization of these tools for each of its mammalian ortholog(s) should be assessed before applying them to a *C. elegans* protein for the first time.

Most but not all Ras superfamily small GTPases are targeted to specific subcellular compartments by membrane-targeting sequences and lipid modification. Ras, Rho, and Rab family members are generally C-terminally modified via farnesyl, geranylgeranyl, or palmitoyl lipids covalently linked to C-terminal cysteines, followed by cleavage of the residues after the modified cysteine (Wennerberg and Der, 2004) (Figure 4). A myristoyl group generally N-terminally modifies Arf family members, and Ran is not modified. For unknown reasons, certain subfamilies in each family are unmodified. Where present, the lipid modification inserts into the membrane bilayer, tethering the small GTPase to the membrane. The typically strong subcellular localization of each small GTPase is collectively dictated by specific small GTPase lipid modification, composition of local membranes, C-terminal electrostatic properties (including post-translational modifications like phosphorylation), and trafficking by chaperone proteins. For most modified small GTPases the lipid modification is essential for targeting and activity. For example, Cys to Ser mutation of the C-terminal CAAX sequence to SAAX results in a non-functional protein, a good control for ectopic expression studies (as is the wild-type GTPase, which due to tight regulatory control is usually inactive when ectopically expressed). In the CRISPR-Cas9 genome-editing era, the SAAX mutation is a good tool to eliminate small GTPase function without disrupting gene structure and attendant stoichiometry of regulatory factors. The utility of this approach, as well as engineered catalytic site mutations for other proteins, is becoming evident as we learn more about the impact of non-canonical regulatory networks, particularly those of non-coding RNAs (Tay et al., 2014).

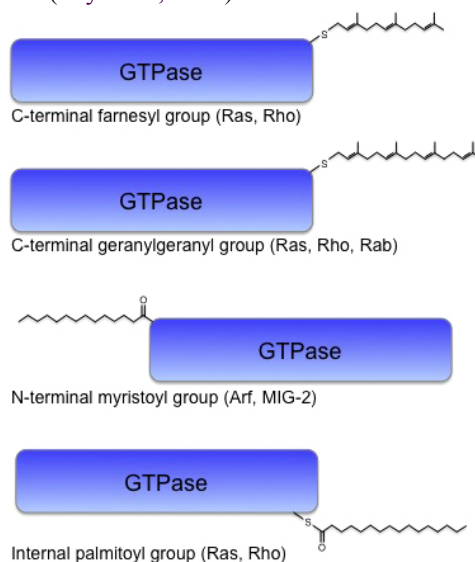


Figure 4. Lipid modifications of the Ras superfamily. Many Ras superfamily small GTPases across species are lipid modified. These examples are taken from mammalian benchmarks, but also include possible myristylation of MIG-2. This figure is an adaptation of (Wennerberg et al., 2005).

Yeast Ras is modified at the C-terminus by farnesyl isoprenoid lipid at the CAAX sequence and this observation was validated in mammalian cells (Casey et al., 1989; Jackson et al., 1990). This feature of Ras biology suggested that treatment with farnesyltransferase inhibitors (FTIs) could block oncogenic Ras signaling. This approach was defeated by two circumstances: first, there are many farnesylated proteins in the mammalian genome, and thus FTIs have substantial and unavoidable off-target effects; and second, when K-Ras4B farnesylation was blocked the protein was alternatively geranylgeranylated, thus bypassing blockade (Lerner et al., 1997; Sun et al., 1998). Otherwise, Ras and members of its superfamily are considered to be “undruggable,” though certain highly specific covalent modifications have been successful (Ostrem et al., 2013).

Of the Ras superfamily members in *C. elegans*, only **LET-60/Ras** has been tested for lipid transferase inhibitor function: addition of the classic farnesyltransferase inhibitor (FTI) was shown to suppress the *let-60* gain-of-function **multivulva** phenotype, consistent with inhibition of **LET-60** membrane targeting (Hara and Han, 1995). HMG-CoA reductase inhibitors lovastatin and fluvastatin can generally block protein prenylation. While not validated with **LET-60**, these more general small molecule inhibitors should inhibit membrane localization of small GTPases, and have been used in *C. elegans* to guide dosing (Morck et al., 2009).

The C-terminal extension of the small GTPase after the G domain is called the “hyper-variable region” because it is poorly conserved even among paralogs in the same subfamily, though there may be conservation of total charge, post-translational sites, and of course the CAAX motif at the very C-terminus, which together influence protein function and signaling outcomes. In addition to lipid modification of the CAAX sequence targeting the protein, the hyper-variable region also contributes to subcellular localization through electrostatic charges that better fit the lipid composition of certain subcellular compartments (Cox et al., 2015). This phenomenon was most dramatically illustrated for the oncogenic human K-Ras4B splice variant, which has a lysine-rich and hence positively charged hyper-variable region. Compared to the alternatively spliced K-Ras4A or other Ras genes N-Ras and H-Ras, which have less basic hyper-variable regions, K-Ras4B plasma membrane targeting is altered and oncogenic potential is much greater (Hancock, 2003). C-terminal electrostatic charge is important enough to K-Ras4B activity that it is thought to explain the enigmatic role of plasma membrane voltage, and attendant mis-regulation of ion channels, in targeting of K-Ras to lipid rafts containing elevated phospholipids (phosphatidylserine and phosphatidylinositol 4,5-bisphosphate) (Zhou et al., 2015). Additionally, each phosphorylation event of a hyper-variable region introduces two negative charges, and alters the electrostatic charge and hence targeting to membrane compartments. The phenomenon was demonstrated in mammalian cells by PKC phosphorylation of K-Ras (Bivona et al., 2006), Aurora A phosphorylation and PP2A dephosphorylation of RalA in the Ras family (Wu et al., 2005; Sablina et al., 2007), and PKA phosphorylation of RhoA in the Rho family (Ellerbroek et al., 2003). In each of these cases the phosphorylation events physically re-targeted the small GTPase and also altered its functional output by altering encounters with effectors. Thus C-terminal phosphorylation may be a common mechanism for dynamically switching small GTPase output.

Another mechanism of small GTPase regulation is that of the chaperone, typically termed a guanine nucleotide dissociation inhibitor (GDI) (Schmick et al., 2015). GDIs have affinity for the GDP-bound, inactive form of the lipid-modified GTPase and thus can shield the hydrophobic lipid-modified tail of the small GTPase from the hydrophilic cytosol during travel between membrane compartments. The mechanisms of GDI action have been investigated for Rab, Rho, and Ras family GDIs (Garcia-Mata et al., 2011; Chandra et al., 2012; Wandinger-Ness and Zerial, 2014). This subject has not been investigated in *C. elegans* beyond the biochemical properties of the single **RHI-1/RhoGDI** (Yap et al., 1999). **RHI-1** promotes increased lifespan of animals (Ha et al., 2006). **rhi-1(RNAi)** in an RNAi-hypersensitive background confers locomotion defects (Simmer et al., 2003) as expected for general deregulation of Rho family members.

Small GTPase output is controlled through conformational shift of the Switch I and Switch II regions upon GTP binding (Colicelli, 2004). Switch I includes the effector-binding domain, a nine-residue core binding sequence that is frequently absolutely conserved between metazoan orthologs. This is not the sole sequence that dictates effector partners, since other sequences across the surface of the protein can influence stability of the GTPase-effector interface. But the region mediates activity-dependent effector interactions across the superfamily, and reasonably selective effector-binding mutations can be found within the core effector-binding domains that do not otherwise disrupt GTPase function. In many cases the different effectors have been identified, but the mechanisms by which the small GTPase switches effectors, or how multiple effectors with different functions are orchestrated, are poorly understood. In effectors the domains that mediate GTPase binding are substantially less conserved across species than are the GTPases that bind them. Thus, from an evolutionary perspective the exceptionally high conservation of effector-binding domains may reflect the use of multiple effectors. One GTPase

and one effector could theoretically evolve together, resulting in substantial changes in sequence. But if a GTPase needs to interact with multiple binding partners, its effector binding sequence is much more constrained evolutionarily.

2. The Ras family

Since Ras oncoproteins are the most frequently mutated oncoproteins, with approximately 35% of all cancers harboring activating Ras mutations (Bos, 1989), Ras itself is an exceptionally well-studied protein. Additionally, Ras is the founding member of the Ras small GTPase superfamily, and all superfamily members have the fundamental Ras small GTPase architecture. Because of its status as a pioneer, many of the general mechanistic principles of small GTPase superfamily members were initially established in studies of Ras across eukaryotes from yeast to humans, including *C. elegans*. In this section, however, we focus just on immediate members of the Ras family. Ras family members share the trait that most of them function in some aspect of signal transduction. Core effector binding domains are probably the main determinants of binding partners, and are generally well conserved phylogenetically (Figure 5).

Rap1A	Q GIFVEK <u>YDPTIEDSYRKQVEVD-CQ</u>
Rap1B	Q GIFVEK <u>YDPTIEDSYRKQVEVD-AQ</u>
RAP-1	Q GIFVEK <u>YDPTIEDSYRKQVEVD-GQ</u>
Rap2A	TGTFIEK <u>YDPTIEDFYRKEIEVD-SS</u>
Rap2B	TGSFIEK <u>YDPTIEDFYRKEIEVD-SS</u>
RAP-2	SSTFIEK <u>YDPTIEDFYRKEIEVD-GQ</u>
RAP-3	Q GIFVHT <u>YDATIEDSYRKLSKVD-AE</u>
M-Ras	QKIFVPD <u>YDPTIEDSYLKHTeid-NQ</u>
RAS-2	QKQFVD <u>YYDPTIEDQYI</u> QHCEID-GN
R-Ras	QSYFVSD <u>YDPTIEDSYTKICSVd-GI</u>
R-Ras3	QSYFVT <u>DYDPTIEDSYTKQCVid-DR</u>
RAS-1	Q RYFVQD <u>YDPTIEDSYTKQCFVd-ED</u>
Rala	YDEFVED <u>YEPTKADSYRKVVLD-GE</u>
RalB	YDEFVED <u>YEPTKADSYRKVVLD-GE</u>
RAL-1	YDEFVEE <u>YEPTKADSYRKVVLD-GE</u>
N-Ras	QNHFVDE <u>YDPTIEDSYRKQVVID-GE</u>
H-Ras	QNHFVDE <u>YDPTIEDSYRKQVVID-GE</u>
K-Ras	QNHFVDE <u>YDPTIEDSYRKQVVID-GE</u>
LET-60	QNHFVEE <u>YDPTIEDSYRKQVVID-GE</u>
Rheb1	EGEFSEGY <u>DPTVENTYSKIVTLG-KD</u>
Rheb2	EGQFVDS <u>YDPTIENTFTKLITVN-GQ</u>
RHEB-1	Q NIFPER <u>YESTIEDQHSKHIAAF-HR</u>
Di-Ras1	KGTFRDT <u>YIPTIEDTYRQVISCD-KS</u>
Di-Ras2	KGTFR <u>ESYIPTVEDTYRQVISCD-KS</u>
Di-Ras3	SGNFR <u>HEYLPTIENTYCQLLGCS-HG</u>
DRN-1	K GTFNEN <u>YVPTIEDTYRQVISCNQ</u> KN

Figure 5. Effector-binding region of human vs. *C. elegans* Ras family members. The entire effector-binding region of human and *C. elegans* proteins was aligned. The core sequence, thought to govern direct protein-protein interactions, is underlined. Core residues not conserved in the subfamily are bolded and blue, and the DRN-1 insertion is bolded and red. Human orthologs are listed above *C. elegans* orthologs. *C. elegans* protein names have a dash and number. Some human proteins have alternate names (see Table 1, Section 9).

2.1. LET-60

let-60 encodes the single *C. elegans* ortholog of mammalian H-, N- and K-Ras. The lysine-rich (and hence very basic) C-terminal hyper-variable region of **LET-60** is most similar to K-Ras4B, the most oncogenic mammalian splice variant. The name derives from the lethality conferred by loss-of-function mutations.

The best-studied *C. elegans* function of **LET-60/Ras** is in transducing the EGF signal that patterns the vulval precursor cells (VPCs): many missense alleles were isolated based on defective patterning of the vulva (Ferguson et al., 1987). A signal originating in the anchor cell (AC) induces vulval fate specification (see WormBook chapter [Vulval Development](#)). The signal consists of EGF, which signals through the **LET-23** EGF receptor (EGFR) (Aroian et al., 1990). The resulting signaling cascade is conserved throughout metazoans (Egan and Weinberg, 1993). A key link between upstream growth factor receptors like EGFR and downstream Ras was discovery of the **SEM-5/Grb2** SH3-SH2-SH3 receptor tyrosine kinase (RTK) adaptor (Clark et al., 1992; Lowenstein et al., 1992; Pawson, 1992;

Rozakis-Adcock et al., 1992). SEM-5/Grb2 in turn recruits the Sos/SOS-1 RasGEF to the RTK signaling complex (Bonfini et al., 1992; Chardin et al., 1993; Chen et al., 1997; Chang et al., 2000). Both activating and temperature-sensitive alleles of *sos-1* have been characterized (Rocheleau et al., 2002; Modzelewska et al., 2007).

LET-60/Ras is necessary and sufficient for vulval induction (Beitel et al., 1990; Han et al., 1990; Han and Sternberg, 1990). Reduced LET-60 signaling causes an under-induced *vulvaless* (Vul) phenotype, while gain of LET-60 function causes 1° hyper-induction and *multivulva* (Muv) animals (Figure 6). Three putative RasGAPs have been identified, and *GAP-1* and *GAP-3* function redundantly to repress LET-60 activity during vulval development, while different combinations of GAPs regulate LET-60-dependent functions in other tissues (Hajnal et al., 1997; Stetak et al., 2008).

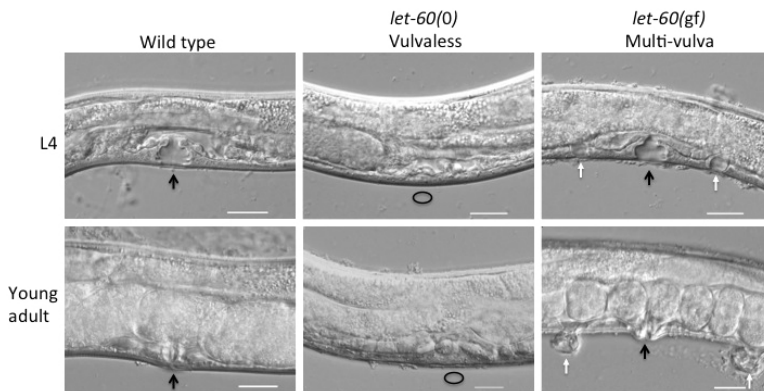


Figure 6. LET-60 functions as a binary switch controlling vulval induction. Vulval development is shown at the late L4 (top row) and adult (bottom row) stages. LET-60 activated by EGF and EGFR induces a single 1° cell flanked by two 2° cells, which divide through stereotyped lineages and undergo morphogenesis to form the L4 “Christmas tree” invagination (above) and mature vulva (below). Black arrows indicate normal vulvae, white arrows indicate ectopic pseudovulvae, and circles represent missing vulvae. The *let-60(dx16)* null deletion allele confers a 100% *vulvaless* phenotype (middle) (Gumienny et al., 1999). The *let-60(n1046gf)* G13E moderately activating allele (right) induces ectopic 1° cells, which go on to form ectopic pseudovulvae (scale bar = 20 μ m). Image credit: Neal Rasmussen from the Reiner lab.

Most identified functions of LET-60 are associated with the LIN-45/Raf Ser/Thr kinase (Han et al., 1993), which, at around the same time, was defined as the central Ras effector in other systems, such as *Drosophila* R7 photoreceptor specification (Ambrosio et al., 1989) and mouse fibroblast oncogenic transformation, activating a MEK-ERK MAP kinase signaling cascade to regulate transcription (Kyriakis et al., 1992; Moodie and Wolfman, 1994).

LET-60 activation of LIN-45/Raf also plays a central role in sex myoblast migration (Sundaram et al., 1996), specification of the excretory duct cell (Yochem et al., 1997), P12 neuroectoblast induction (Jiang and Sternberg, 1998), excretory/fluid homeostasis (Schutzman et al., 2001), and olfaction (Hirotsu et al., 2000). LET-60 also functions non-autonomously to regulate neuroblast delamination (Parry and Sundaram, 2014), and, with LIN-12/Notch, controls development of the excretory tube (Abdus-Saboor et al., 2011). LET-60 also triggers multiple ERK-dependent events in the germline (Eisenmann and Kim, 1997; Arur et al., 2009). For the cell fate patterning amongst groups of equipotent cells, LET-60 functions downstream of LET-23/EGFR; for fluid homeostasis LET-60 functions downstream of EGL-15/FGFR (see WormBook chapter Canonical RTK-Ras-ERK signaling and related alternative pathways).

The first Ras effector discovered was yeast adenylyl cyclase, and its Ras binding domain is a series of distinctive leucine-rich repeats (LRRs). Researchers initially thought that mammalian oncogenic Ras signaling would similarly be through adenylyl cyclase, but metazoan adenylyl cyclases do not have the LRR domain. Instead, the major Ras effectors in mammalian cells were found to be Raf kinase, and later PI3 Kinase, RalGEF, and others (Repasky et al., 2004). Nonetheless, LRR domains are still relevant for Ras signaling across phyla, and multiple Ras partners contain an LRR domain. A critical member of this group was discovered as a suppressor of activated Ras: SOC-2/SUR-8 (mammalian Shoc2) functions as a scaffold for Ras and Raf, and may be a major regulatory point for Ras signaling (Selkors et al., 1998; Sieburth et al., 1998). Additionally, missense mutation of Shoc2 causes defects in the RASopathy spectrum (Cordeeddu et al., 2009). RASopathies are a series of heritable disorders in which derepression of the Ras→Raf→MEK→ERK cascade causes heritable birth defects, mental retardation, and increased risk of cancer (Ratner and Miller, 2015).

Oncogenic activating mutations in Ras have mutations at positions 12, 13, and 61 as canonical activating mutations, with different alleles predominating in different tumor types (Pylayeva-Gupta et al., 2011). Mutations have also been developed to select for different effector usage. In the background of the mammalian Ras G12V activating mutation, the E37G mutation confers selectivity for RalGEF: the E37G compromises Ras binding to Raf and PI3K effectors, but leaves Ras→RalGEF interactions intact (White et al., 1995; Rodriguez-Viciano et al., 2004). The G12V,E37G mutant Ras has been well validated in subsequent cancer studies (Hamad et al., 2002), thus providing an excellent tool for dissecting Ras→RalGEF signaling (see Section 2.7). The *let-60(s1124)* (E37K) mutation was isolated as a strong loss of function allele, and mutates the same E37 residue as the RalGEF-selective allele, presumably blocking Raf binding. In mammals Ras T35S and Y40C are selective for Raf and PI3K activation, respectively (White et al., 1995); of these, T35S and E37G have been validated in *C. elegans* (Zand et al., 2011).

A series of loss-of-function, dominant-negative, and gain-of-function alleles defined the *let-60* locus (also known as *lin-34*, from the gain-of-function alleles) (Beitel et al., 1990; Han and Sternberg, 1991). Isolated based on vulval development phenotypes (or lethality from strong alleles due to failed induction of the excretory duct cell, see below), this collection of *let-60* alleles reflects the diversity of Ras alleles defined in other organisms.

The reference gain-of-function *let-60* allele is *n1046gf*, which causes a G13E substitution. Oncogenic position 13 alleles in Ras are found in cancers and are moderately activating. Four additional G13E activating alleles have been found for *let-60* (*n1700*, *n1849*, *sy103*, *sy130*), and all cause similar phenotypes. Two non-canonical gain-of-function alleles (i.e., not position 12, 13, or 61 mutations found in cancer) have also been isolated in screens: *ay75gf,ts* (G60R) confers a semi-dominant and temperature-sensitive defect in fluid homeostasis, consistent with constitutive activation of the EGL-15/FGFR cascade that regulates that process; and *ay75gf,ts* confers less penetrant fertility and vulval hyper-induction than the canonical G13E alleles like *n1046* (Schutzman et al., 2001). *let-60(ga89gf,ts)* (L19F) confers semi-dominant and temperature-sensitive defects in fertility, consistent with activation of the Raf-MEK-ERK cascade at multiple points in the germline, but only weakly induces vulval hyper-induction (Eisenmann and Kim, 1997; Arur et al., 2009). Notably, *let-60(n1046gf)* alone promotes neither fluid homeostasis defects nor sterility, though it can do so in sensitized backgrounds. Though all three of these defects are Raf-dependent, the Raf cascade is evidently activated at different levels in different tissues, or different sets of co-effectors are activated in different tissues. Also, the absence of identified completely activating lesions at positions 12 and 61 suggests that such mutants would be lethal or sterile.

Dominant-negative *let-60* alleles were also identified in mutant screens (Han and Sternberg, 1991). *let-60* alleles *sy101* and *sy99* (G10R), *sy94* (K16N), and *sy93* (D119N) all confer a dominant-negative vulvaless phenotype. None of them corresponded to canonical S17N dominant-negative mutations identified from mammals, perhaps, as hypothesized for strong gain-of-function mutations, due to potential toxicity of strong alleles.

2.2. RAS-1

ras-1 encodes the ortholog of mammalian R-Ras1 and R-Ras2/TC21. Very little has been done with the *C. elegans* **RAS-1**, with one exception. RASopathies are a series of mostly overlapping syndromes caused by mutations that weakly activate the Ras→Raf→MEK→ERK MAP kinase cascade (Rauen, 2013). In a RASopathy patient, putative activating mutations in R-Ras were found. Heat-shock induction of *C. elegans* **RAS-1** harboring the homologous mutation induced weak vulval morphogenesis defects compared to wild-type **RAS-1**, suggesting that **RAS-1** may contribute to LET-60/Ras-LIN-45/Raf signaling (Flex et al., 2014), with the caveat that ectopic over-expression of mutationally activated **RAS-1** could inappropriately engage effectors that typically associate with LET-60. R-Ras, TC21, and M-Ras (**RAS-2**, Section 2.3) share the trait of binding the RA domain of RalGEFs in vitro (Rodriguez-Viciano et al., 2004), raising the possibility that **RAS-1** or **RAS-2** could activate RalGEF (see Section 2.7). However, deletion alleles of neither *ras-1* nor *ras-2* conferred vulval phenotypes consistent with RalGEF interaction, but the double mutant has not been described (D.Reiner, unpublished data).

A key observation in mammalian R-Ras biology was the finding that the semaphorin receptors, plexins, function as ligand-dependent GAPs with specificity for R-Ras and M-Ras. R-Ras (and M-Ras) in turn activate integrins, so semaphorin-plexin signaling inhibits integrin activation, thus allowing growth cone collapse (Puschel, 2007). Later studies challenged this model, instead arguing that plexins function as Rap-specific GAPs (Wang et al., 2012; Wang et al., 2013). Plexins also impact Rho family signaling, so the story is liable to be complex. In *C. elegans*, plexins function in collaboration with Eph receptors to coordinate development of sensory rays in the male tail and embryonic ventral enclosure (Ikegami et al., 2004; Ikegami et al., 2012), but connections to Ras family members have not been observed.

2.3. RAS-2

ras-2 encodes the ortholog of mammalian M-Ras/R-Ras3. **RAS-2** is one of the *C. elegans* Ras-like proteins whose effector-binding domain is not absolutely conserved: a mammalian M-Ras effector-binding domain Ser that is conserved in human and *C. elegans* **LET-60**/Ras, Rap1, and **RAS-1** is a Gln in **RAS-2**, perhaps indicating that **RAS-2** effector partners are not conserved with mammalian partners. A major point of interest with mammalian M-Ras is that it binds the LRR domain of Shoc2, thus regulating the Ras-Raf scaffold and regulating a major oncogenic cascade (Rodriguez-Viciiana et al., 2006; Young et al., 2013). However, no functions have been associated in **RAS-2** in *C. elegans*.

2.4. RAP-1

rap-1 encodes the ortholog of mammalian Rap1A/Krev-1 and Rap1B. Rap1 started out with misdirection and continues to confound simple categorization. Rap1/**RAP-1**, as well as M-Ras and R-Ras/TC21/RAS1, have effector binding domains identical to those of **LET-60**/Ras. This observation, coupled with similar overall structure across the rest of the proteins, led early researchers to believe that extensive functional redundancy existed across the protein group. Subsequent experiments argue otherwise, with only Ras itself demonstrating widespread and central roles in biology across species. The effector domain identity between Ras and Rap1, then, remains enigmatic. Perhaps the functional differences are defined by divergent C-terminal membrane localization sequences, resulting in interaction with different partners in different subcellular compartments (Prior and Hancock, 2012).

Rap1 was initially identified as a Ras-like protein (Ras proximal), and over-expression assays led cancer researchers to believe that Rap1 functioned as a competitive inhibitor of Ras activation of Raf kinase (Frische and Zwartkruis, 2010). More precise experiments later disproved this notion, but the possible connection between Rap1 and Raf is still controversial in the mammalian literature, compounded by barriers to genetic analysis presented by two Rap1 genes, three Ras genes, and three Raf genes. In contrast, experiments with the single Rap1 gene in *Drosophila* support the model that Rap1 binds and activates Raf1 (Mishra et al., 2005; Mavromatakis and Tomlinson, 2012). The issue remains unaddressed in *C. elegans*.

In most systems Rap1 (and Rap2) are known for regulating biology of cell junctions, and many Rap1 (and Rap2) effectors impact junctional biology and/or morphogenesis (Kooistra et al., 2007). In *C. elegans*, **RAP-1** and **RAP-2** redundantly function in the hypodermis to control larval molting, such that few double mutant animals survive. **RAP-1** and **RAP-2** are jointly activated in this process by **PXF-1** ortholog of the mammalian RapGEF PDZGEF (Pellis-van Berkel et al., 2005). The **PXF-1** larval arrest mutant phenotype was partially rescued by heat-shock expression of the **RAP-1** G12V gain-of-function. **PXF-1** promoted GTP exchange specifically on **RAP-1** and **RAP-2**, but not **LET-60**/Ras or **RAL-1**. The *pxf-1* promoter::GFP fusion was expressed in hypodermis in pulses corresponding with larval molts, consistent with function at that time and place. Other structural defects were characterized. Perhaps surprisingly, compared to the role of Rap1 in other systems, *rap-1* mutant animals are viable. The *rap-1(pk2082)* nonsense allele was initially characterized as conferring growth and structural defects, but additional outcrossing and comparison to the *tm861* deletion allele (Frische et al., 2007; D. Reiner, unpublished results) indicates that the growth defects of *rap-1* mutant animals, if any, are mild.

The expected **RAP-1** contribution to junctional biology was found in a synthetic lethal RNAi screen, which identified the related Ras family small GTPase, **RAL-1**, as providing redundant functions in hypodermal junction formation and cadherin complex localization, resulting in rupturing of embryos (Frische et al., 2007) (see Section 2.7). Components of the **RAL-1** signal were analyzed, but **RAP-1** was not. **RAP-2** apparently played no role in this redundant event. The endogenous *rap-1* locus was tagged, using CRISPR technology, with fluorophore mNeonGreen at the 5' end, creating an N-terminally tagged protein that localized to most or all plasma membranes (Dickinson et al., 2015).

RNAi screens for defective male somatic gonad identified both RapGEF PDZGEF/**PXF-1** and **RAP-1** (Kalis et al., 2010), and *pxf-1* was identified in an RNAi screen for resistance to paralysis induced by the acetylcholine esterase inhibitor aldicarb (Sieburth et al., 2005). The *rgef-1* gene encodes the *C. elegans* ortholog of mammalian CalDAG GEF/RasGRP2, which promotes exchange on both Ras and Rap1 (Raaijmakers and Bos, 2009). The **RGEF-1** exchange activity is conserved in *C. elegans* but the reported RapGEF function in olfaction functions through **LET-60**/Ras and not **RAP-1** (Chen et al., 2011).

All RapGEFs are conserved in *C. elegans*, but many RapGEFs also activate Rap2, and some can activate Ras and other Ras family members. In addition to the aforementioned **PXF-1/PDZGEF** and **RGEF-1/CaldAGGEF**, *C. elegans* expresses the putative RapGEF **EPAC-1/cAMP-activated RapGEF**, but only phenotypes associated with **DRN-1** (ortholog of mammalian Di-Ras1/Rig, Di-Ras2, and Di-Ras3/Noey2/ARHI, see Section 2.9) have been identified (Tada et al., 2012). The RapGEF C3G/Y34B4A.4 is encoded in *C. elegans*, but no phenotype is associated with this protein. Worms also encode the combination RapGEF and phospholipase epsilon **PLC-1/PLC-ε**. PLC-ε has been identified as an effector for Ras in *C. elegans* and other systems (Bunney and Katan, 2006). **PLC-1** is required for fertility and ovulation (Shibatohge et al., 1998; Kariya et al., 2004) and embryonic morphogenesis (Vazquez-Manrique et al., 2008). It is unclear whether these activities are connected with small GTPase signaling. Oddly for Ras family GEFs, across metazoans PDZGEF, EPAC, and PLC-ε also contain Ras Association (RA) domains, which potentially serve as Rap1 effectors or points of cross talk with other related Ras family proteins that can also bind RA domains (Raaijmakers and Bos, 2009). Large sets of Rap effectors have been characterized, and many of them are shared with Rap2.

Rap1 and Rap2 have also been suggested to be inhibitory substrates of the semaphorin receptor, plexin. Activated plexins functions as GAPs, though whether for Raps, R-Ras, or both is unclear (Puschel, 2007; Wang et al., 2012; Wang et al., 2013). The Rap subfamily is also notable for having a Thr at position 61 (Ras numbering, compared to the Gln at position 61 in most Ras family members). This change is thought to confer GAP specificity (Daumke et al., 2004), and is shared only with the Di-Ras subfamily among Ras family members.

2.5. RAP-2

rap-2 encodes the ortholog of mammalian isoforms Rap2A, Rap2B, and Rap2C. Rap2 and Rap1 junctional functions in other systems are significantly overlapping, though individual activities have been defined (Raaijmakers and Bos, 2009). In *C. elegans* the only demonstrated biological outcome of **RAP-2** activity is its redundant regulation of molting in concert with **RAP-1**, downstream of the GEF **PXF-1** (see Section 2.4) (Pellis-van Berkel et al., 2005). Mammalian Rap2 signals to Ezrin via TNIK, and this general relationship was supported in *C. elegans* by localized enrichment of phospho-Ezrin in a **rap-2** null mutation background (Gloerich et al., 2012). Beyond this loose connection Rap2 effectors have not been characterized.

The *rap-2(gk11)* mutation has been shown, through redundancy with the **rap-1** null allele, to strongly reduce **rap-2** function, but single mutant animals are superficially wild type (Pellis-van Berkel et al., 2005). No other functions have been associated with RAP-2.

2.6. RAP-3

rap-3 encodes a divergent Rap-like protein. Overall GTPase domain structure, hypervariable region, and CAAX sequences are most similar to Rap proteins, particularly Rap1. However, key changes in the **RAP-3** core effector-binding region, particularly a substitution of Ala for a Pro conserved in all Ras family members, suggests that **RAP-3** does not functionally overlap its phylogenetically conserved brethren **RAP-1** and **RAP-2**. **RAP-3** is conserved in related nematodes but not humans or *Drosophila*, suggesting that **RAP-3** represents a nematode-specific addition that may engage distinct effectors. No functions have been ascribed to **RAP-3**.

2.7. RAL-1

ral-1 encodes the ortholog of mammalian RalA and RalB. By RNAseq the *C. elegans* **ral-1A** transcript encodes a Ral ortholog that is well conserved across the entire GTPase. The minority **ral-1B** transcript is predicted to encode a RAL-1B protein with a unique 41 residue N-terminal extension that is unprecedented in other species. It is unclear whether **ral-1B** transcript produces protein, and it has not been detected in related nematode species.

The function of RAL is to promote 2° vulval cell fate (see [Canonical RTK-Ras-ERK signaling and related alternative pathways](#)) and shorten lifespan (WormBook chapters [Dauer](#) and [Obesity and the regulation of fat metabolism](#)). The main Ral claim to fame in mammals is as a non-canonical effector of the Ras GTPase (see Section 2.1). This is a case of one Ras subfamily activating another Ras subfamily: activated Ras binds the RA (Ras Association) domain of RalGEF, thereby recruiting the GEF to the membrane and stimulating GTP exchange activity of Ral. Ral is generally in the vicinity of its activated GEF since, like Ras, Ral is membrane localized via its C-terminus. Indirect genetic results from *Drosophila* suggest that Rap1 (Section 2.4) also binds and activates RalGEF (Mirey et al., 2003), but these results are also consistent with parallel signaling. In all systems examined,

Rap1, like Ras, can bind the RalGEF RA domain in vitro or in yeast two hybrid assays, but evidence is lacking in vivo, including in *C. elegans* vulval patterning (Zand et al., 2011) (see below).

Unusual for a GEF and GTPase pair, RalGEF and Ral are monogamous: RalGEFs are thought to activate only Ral, and Ral is activated only by RalGEFs (Feig, 2003). Invertebrates have single RalGEF genes (encoded by *rgl-1* in *C. elegans*) compared to four in mammals. Deletion mutants of RalGEF are superficially wild type (T. Zand, K. Monahan and D. Reiner, unpublished results). Mammals and *Drosophila*, but not *C. elegans*, also encode Ras-independent RalGEFs, RalGPS1/2, that do not have an RA domain and are thought to be regulated by lipid signaling, but little is known about these proteins.

Despite exceptionally high sequence identity throughout the Ral GTPase domain, mammalian RalA and RalB, though both contributing to cancer, perform distinct functions in the cell. The RalA mouse knockout caused embryonic lethality while the RalB knockout is viable. The RalA/B double knockout is more severe than the RalA single knockout, consistent with diverging but also overlapping functions (Peschard et al., 2012). This difference between RalA and RalB functions is conferred by the hyper-variable C-terminal membrane targeting sequences (Feig, 2003; Camonis and White, 2005). *C. elegans* and *Drosophila* encode single Ral proteins.

A regulatory switch from Ras→Raf to Ras→RalGEF→Ral resolved a contradiction between morphogen gradient and sequential induction models of vulval cell fate patterning. Understanding this switch requires understanding of vulval cell fate patterning (see Vulval Development; Sherwood et al., 2005; and Canonical RTK-Ras-ERK signaling and related alternative pathways). A long-standing controversy in the field is the contradiction between two general models of vulval patterning by EGF. Classic developmental experiments and manipulation of strength of EGF and EGFR activity led to the morphogen gradient model (Figure 7). This model posits that an EGF gradient dictates the 3°-3°-2°-1°-2°-3° pattern of vulval fates, with 1° fate being determined by closest proximity to the AC, and hence EGF (Sternberg and Horvitz, 1986; Sternberg, 1988; Sternberg and Horvitz, 1989; Katz et al., 1995; Katz et al., 1996). Mutant screens identified under-induced (Vulvaless; Vul) and hyper-induced (Multivulva, Muv) mutants defining the core genes involved in patterning the vulva (Ferguson et al., 1987). Cloning of these genes led to identification of two signaling axes, EGFR→Ras→Raf→MEK→ERK MAP kinase, and Notch→CSL transcription factor, that are necessary for 1° and 2° signals, respectively (reviewed in WormBook chapters: LIN-12/Notch signaling in *C. elegans*; Notch signaling: genetics and structure; Vulval development; Canonical RTK-Ras-ERK signaling and related alternative pathways). The sequential induction model arose from findings that EGFR and Notch are necessary and sufficient in 1° and 2° cell fates, respectively (Seydoux and Greenwald, 1989; Koga and Ohshima, 1995; Simske and Kim, 1995). Initial EGF signal activates the Ras→Raf→MEK→ERK cascade to induce 1° cell fate. In presumptive 1° cells, this ERK MAP kinase cascade also induces Notch ligands, which induce two neighboring VPCs to become 2° (Chen and Greenwald, 2004). Importantly, 1° and 2° cell patterning signals are antagonistic. Notch activates expression of ERK phosphatase (LIP) in presumptive 2° cells to block inappropriate ERK activity (Berset et al., 2001), while LIN-12/Notch receptor is internalized and degraded in presumptive 1° cells to similarly block inappropriate Notch activation in presumptive 1° cells (Shaye and Greenwald, 2002; Shaye and Greenwald, 2005). Importantly, transcription of many modifier genes is re-programmed between initial EGF induction and the first cell division of VPCs (Berset et al., 2001; Yoo et al., 2004; Berset et al., 2005; Yoo and Greenwald, 2005; Zand et al., 2011). Consequently, vulval development can be divided into two stages: initial specification, which is reversible, and commitment/reinforcement, which is irreversible and results in final cell fate (Vulval development).

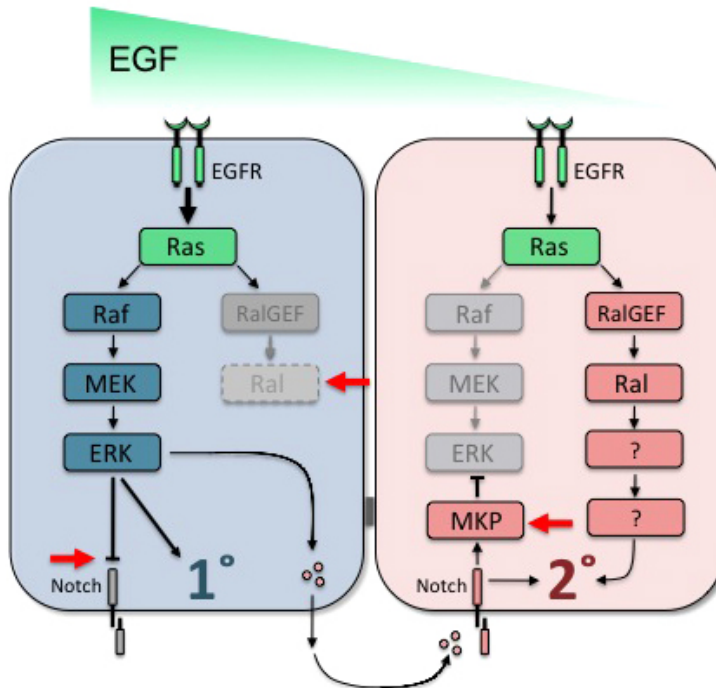


Figure 7. Vulval Ras switches effectors depending on cell position in the EGF gradient. Two models of vulval cell fate patterning, the morphogen gradient model and the sequential induction model, were formerly considered contradictory (Kenyon, 1995). The discovery that Ras→RalGEF→Ral mediates lower dose EGF signal to promote 2° fate reconciled the two models, and the combined model is shown here. Red arrows show key points at which signals are switched from supporting one cell type to the other. LIN-12/Notch is internalized and degraded in presumptive 1° cells to prevent contradictory Notch signaling. Notch-dependent transcriptional up-regulation of LIP-1 ERK phosphatase in presumptive 2° cells represses inappropriate MPK-1/ERK activation. Ral promoter activity is excluded from presumptive 1° but persists in presumptive 2° cells, thus preventing inappropriate activation of Ral.

That conflicting signals exist in VPCs prior to their final fate commitment allowed identification of additional signaling cascades (Zand et al., 2011; Nakdimon et al., 2012), particularly identification of Ras→RalGEF→Ral, which explains how high and low doses of EGF can induce different outcomes in Ras signaling. In presumptive 2° cells Ras also signals, but through RalGEF→Ral to promote 2° fate rather than Raf→MEK→ERK (Figure 7). Ras→RalGEF→Ral signal is dispensable for core patterning of the vulval precursor cells (Zand et al., 2011), but is an important positive regulator of Notch signal in VPCs. Ras→RalGEF→Ral thus mediates lower dose EGF signal in the morphogen gradient. And so the gradient is overlaid on the 1°-2° sequential induction by Ras→Raf→MEK→ERK and Notch. This work resolved the controversy in the field of how to reconcile conflicting observations of sequential induction vs. morphogen gradient patterning of the VPCs (Katz et al., 1995; Kenyon, 1995; Koga and Ohshima, 1995; Simske and Kim, 1995; Katz et al., 1996). This Ras switch in effectors is orchestrated at a cellular level by two complementary mechanisms. First, a *ral-1* transcriptional reporter is initially expressed in all of the vulval precursor cells, but after the onset of EGF signaling, expression is excluded from presumptive 1° cells while persisting in presumptive 2° cells. Thus unknown transcriptional circuitry insulates presumptive 1° cells from inappropriate Ras→RalGEF→Ral signaling. In addition, the Ras→Raf output of ERK phosphorylation is quenched in presumptive 2° cells by Notch-dependent transcription of the LIP-1/MKP/ERK phosphatase and other Notch-dependent transcriptional client genes (Berset et al., 2001; Yoo et al., 2004; Berset et al., 2005; Yoo and Greenwald, 2005).

This study was the first example of Ras dynamically switching effectors; it was previously assumed that different cell types were programmed to respond with different effectors. But Ras and its effectors Raf and RalGEF are all present in naïve VPCs before the onset of EGF signaling. Thus, a key unresolved question is whether and how Ras discriminates between partners that appear otherwise equivalent (Reiner, 2011). This mechanism may contribute to the 99.8% accuracy of VPC patterning (Braendle and Felix, 2008), particularly given the centrality of Ras→Raf in the process and the pitfalls of inappropriate signaling.

As with many ancient small GTPases, **RAL-1** and *Drosophila* Ral share exceptionally high identity with RalA and RalB throughout the GTPase domain, including 100% identity in the core effector-binding region, suggesting that **RAL-1** and RalA/B regulate a common suite of effectors. Because of their roles in RalA/B-dependent oncogenic

signaling, three canonical Ral binding partners have received most of the attention. RalBP1 (also known as RLIP76, RLIP1, and RIP1) contains a RhoGAP domain shown in vitro to be selective for Rac and Cdc42, has membrane transporter ATPase activity, and can scaffold proteins participating in endocytosis (Gentry et al., 2014). Phosphorylation of the C-terminus of mammalian RalA relocalizes it to the mitochondrial membrane, where it signals through RalBP1 to promote mitochondrial fission at mitosis (Kashatus et al., 2011). *C. elegans* RLBP-1 contains Ral-binding and RhoGAP domains, but it is unknown whether all of the mammalian RalBP1 functions are conserved and what functions RLBP-1 performs. Biological functions of *C. elegans* RLBP-1 are unknown.

Sec5 and Exo84 are well-validated binding partners of mammalian RalA/B and are essential and non-essential components of the exocyst complex, respectively. The exocyst is a phylogenetically conserved heterooligomeric complex that controls exocytosis and diverse aspects of cell behavior (Shirakawa and Horiuchi, 2015; Wu and Guo, 2015). The yeast exocyst uses Rab and Rho for this purpose, but metazoans seem to have replaced those functions with Ral (Novick and Guo, 2002). *ral-1(tm2760)*, a deletion in intron 3 that perturbs the splice donor sequence, confers sterility (Zand et al., 2011), as does the putative null deletion *tm5205*. Depletion of both maternal and zygotic *ral-1* disrupted cell polarity, tubulogenesis, and exocytosis, and these functions are closely associated with the PAR proteins (Armenti et al., 2014). Disruption of maternal and zygotic SEC-5, a central exocyst complex component from yeast to humans, confers the same phenotype, suggesting that RAL-1 participation as a member of the exocyst regulates fundamental cell biological events. Null mutants for RGL-1/RalGEF are superficially wild type (T. Zand, K. Monahan, and D. Reiner, unpublished results), consistent with the RAL-1 functions in the exocyst being activation-independent.

Exo84 and Sec5 serve as RalA/B docking sites for the exocyst in mammals. Exo84 and Sec5 also function as signaling scaffolds, coupling Ral to downstream activities. Sec5 mediates RalB-dependent activation of TBK1 in innate immunity and tumor survival (Chien et al., 2006). Mammalian Exo84 mediates RalB-dependent activation of autophagosome assembly (Bodemann et al., 2011) and mTORC1 activation (Martin et al., 2014) (see below). But many signaling intermediaries that execute these Ral-Sec5- and Ral-Exo84-dependent functions remain unknown.

In *C. elegans*, Exo84 and Sec5 have only been connected with Ral in regulation of cadherin-based junctions in embryonic hypodermal cells (Frische et al., 2007). In a screen for hypodermal phenotypes, *rgl-1* and *ral-1* were recovered as genes that confer a synthetic lethal RNAi phenotype in animals mutant for the RAP-1 small GTPase (Section 2.4). Also identified were genes encoding Ral binding partners SEC-5 and EXOC-8. No other characterized Ral-dependent process has used both exocyst partners, so perhaps Ral engages two genetically separable but required activities in this process. *rlbp-1* and genes encoding other exocyst components were not synthetic lethal with *rap-1* mutations. RAL-1 and RAP-1 signaling redundantly control cadherin complex localization to junctions of embryonic hypodermal cells, and hence double mutants disrupt epithelial integrity, migration, and morphogenesis. Upstream input has not been tested, particularly by LET-60/Ras, which might be expected. RAP-2 (Section 2.5) was not found to play a role in this process.

The existence of mammalian RalGAP biochemical activity was known for decades (Emkey et al., 1991), but the molecular identity of RalGAP was only identified relatively recently (Shirakawa et al., 2009; Chen et al., 2011). RalGAP is a heterodimer consisting of a catalytic α subunit and regulatory β subunit. The homology of the GAP domain and unusual regulatory architecture draws a parallel to the Tuberous Sclerosis Complex (TSC) heterodimeric RhebGAP (see Section 2.8). Determination that Akt phosphorylation inhibits RalGAP like it does TSC/RhebGAP further underscores the similarity between RalGAP and TSC/RhebGAP (Chen et al., 2011; Leto et al., 2013). However, nematode genomes do not express TSC orthologs. *C. elegans* *hgap-1* and *hgap-2* encode orthologs of RalGAP α and β subunits, respectively. Based on the reasoning that the ancestral nematode must have had compensating RhebGAP activity to allow loss of TSC/RhebGAP, RalGAP is a candidate for a shared Ral and Rheb GAP in *C. elegans*, hence the intentionally noncommittal name of Heterodimeric GTPase Activating Protein (HGAP). Putative null mutation of *hgap-1* or *hgap-2* reduced lifespan, as predicted for a TSC-like complex (Martin et al., 2014). This reduced lifespan phenotype was reversed by *ral-1(RNAi)*, consistent with the HGAP complex repressing RAL-1 activity as a RalGAP but leaving potential function as a RhebGAP unresolved. HGAP-2 was previously identified in an RNAi screen for decrease of DAF-2/InsR mutant longevity (Samuelson et al., 2007), but because RalGAP was as yet undiscovered, its role in signaling was not recognized. RAL-1 was identified as an aging gene in an RNAi screen for sensitivity to oxidative stress (Kim and Sun, 2007).

Based on the expectation that a common ancestor of *C. elegans* and mammals used both RalGAP and TSC/RhebGAP to regulate TOR, crosstalk was found between mammalian RalGAP and TSC/RhebGAP, and RalB was found to signal through Exo84 to directly activate TORC1. However, RalB localized TORC1 to the plasma membrane to mediate RalB-dependent invasion activity in cell culture, contrasting with canonical Rheb-TORC1

activation of anabolic metabolism at the lysosome. Thus Ral and Rheb may recruit TORC1 to different subcellular compartments to mediate distinct activities (Martin et al., 2014). RalB→Exo84 activates autophagy under starvation conditions (Bodemann et al., 2011). Autophagy is catabolic, is generally regulated in opposition to TORC1, and is directly inhibited by TORC1 kinase activity. Thus these two activities of RalB in cell culture may be contradictory. Alternatively, perhaps RalB switches between promoting catabolism when cells are starved and anabolism when cells are well fed. These signaling permutations suggest a complex regulatory dance in the interpretation of upstream nutritional status by Ral and Rheb.

In support of a mammalian study showing RalA-JNK-FOXO4 signaling in stress response, exogenous mammalian GST-JNK was phosphorylated in response to heat shock in a **RAL-1** dependent manner in worm extracts, and upon heat shock transgenic *C. elegans* FOXO/DAF-16::GFP translocated to the nucleus in a **RAL-1** dependent manner. Exogenous tagged mammalian RLF/RalGEF and RalA bound to the JIP1 JNK scaffold upon stimulation by reactive oxygen species, suggesting that Ral contributes to a central scaffold to promote JNK activation (van den Berg et al., 2013).

2.8. RHEB-1

rheb-1 encodes the single *C. elegans* ortholog of mammalian Rheb1 and Rheb2/RhebL1. While **RHEB-1** is clearly most similar to mammalian Rhebs, certain sequence features of the nematode protein are unconventional. First, mammalian and *Drosophila* Rhebs diverge from other Ras family members at Ras oncogenic mutation positions 12 and 13 (Ras numbering), having an “RC or “RS” instead of “GG”. *C. elegans* **RHEB-1** has a “PH” at positions 12, 13 (Figure 8). Second, though the mammalian and *Drosophila* Rheb core effector-binding domains are distinct from those in the rest of the Ras family, certain sequence elements are conserved throughout. Strikingly, the *C. elegans* **RHEB-1** core effector-binding domain diverges at multiple positions, including the third position Proline conserved in most Ras family members (Figure 8). Even yeast Rhebs, *S. cerevisiae* Rhb⁺ and *S. pombe* Rheb, have better conserved effector-binding domains than *C. elegans* **RHEB-1**. RHEB-1 may therefore be the exception that proves the rule in co-conservation of small GTPases and effector domains (see above): Rheb across species only has a single described effector, the Target of Rapamycin (TOR) Ser/Thr kinase, so perhaps there is greater tolerance of changes in the Rheb core effector binding region, which are presumably reflected in its downstream partner. Alignment of the *C. elegans*, *Drosophila*, and two human Rheb sequences throughout the GTPase domain indicates that, in *C. elegans*, Rheb has converged toward Ral (D. Reiner, unpublished observations) (Figure 8). As a control, *Drosophila* Rheb has not converged toward *Drosophila* Ral. Perhaps this convergence accompanied the evolutionary loss of TSC in *C. elegans*: if the HGAP-1/2 RalGAP performs both RalGAP and RhebGAP activities in *C. elegans*, it stands to reason that the described changes in **RHEB-1** to become more Ral-like facilitated co-regulation of these protein, which are separately regulated in other systems.

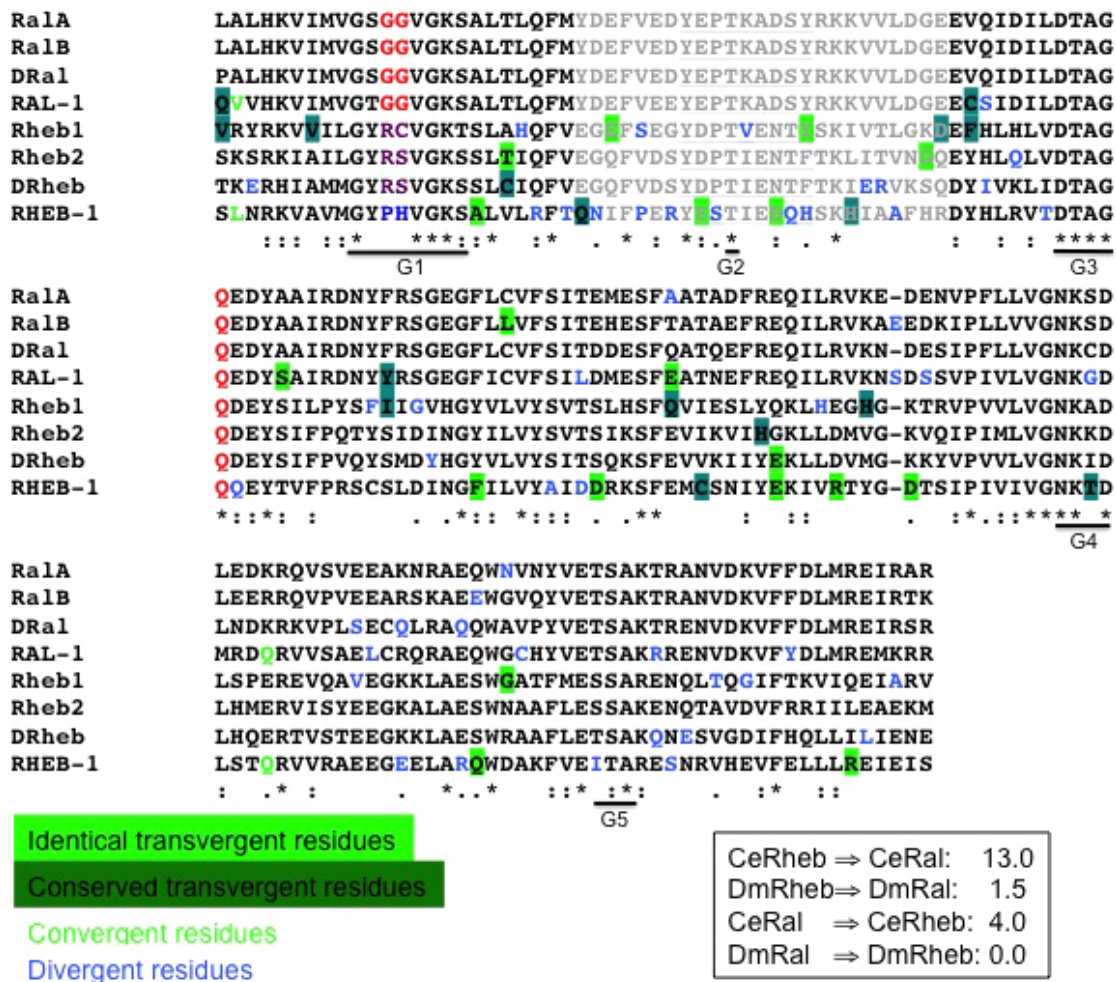


Figure 8. *C. elegans* Rheb has converged towards the Ral subfamily. The convergence was observed for *C. elegans* Rheb and Ral, but not *Drosophila*. Core GTPase sequences of human, fly, and worm Rheb subfamily and Ral subfamily proteins were aligned and scored for number of convergent vs. divergent residues. Residues that converge towards the consensus of the compared subfamily (here called “transvergent”) and are identical (light green highlighting) were valued at 1, with conservative transvergence (dark green highlighting) valued at 0.5. True convergence of a residue from both subfamilies to a new residue type (light green letter) was also valued at 0.5, while divergence (blue letter) was valued as zero. Events in the broad effector binding region (grayed, with core underlined) or G boxes were rated with higher impact (0.5 becoming 1.0). By this standard *C. elegans* RHEB-1 converged toward *C. elegans* RAL-1, with *Drosophila* and human Rheb and Ral as negative controls (Fisher's Exact Test, $P < 0.0001$). *C. elegans* RHEB-1, and to a lesser degree *C. elegans* RAL-1, also contains significantly more divergent residues, perhaps reflecting a protein under more intense selective pressure than is typical. This convergent evolution may reflect the loss of TSC/RhebGAP but retention of HGAP/RalGAP in the *C. elegans* clade, so the RHEB-1 protein is adapted to co-regulation by RalGAP. Classic Rheb changes at positions 12 and 13 (Ras numbering) are indicated by purple (blue in RHEB-1).

Rheb is known as a key activator of TOR. TOR is associated with two complexes. TORC1 incorporates the co-factor Raptor and is a critical control point for promoting anabolic metabolism at the expense of catabolic anabolism (autophagy) (Dibble and Cantley, 2015). TORC2 mediates a feedback loop to Sgk and Akt in both mammals and *C. elegans* (Jones et al., 2009; Soukas et al., 2009). Rheb, and its inhibitory TSC RhebGAP, has different effects on TORC1 and TORC2: Rheb is well established as directly promoting TORC1 activation, but Rheb may inhibit TORC2 activation (Yang et al., 2006; Groenewoud and Zwartkruis, 2013).

Consistent with the model of RHEB-1 activating TOR and anabolic metabolism, which generally decreases lifespan, *rheb-1(RNAi)* extends lifespan (Honjoh et al., 2009). However, the same study found that in an intermittent fasting paradigm, which extends lifespan, *rheb-1(RNAi)* decreases lifespan. That is, RHEB-1 is required for the intermittent fasting lifespan extension. These contradictory observations suggest that RHEB-1- and TOR-dependent regulation of lifespan are likely to be context dependent. RHEB-1 signaling may regulate both insulin signaling and FoxA, which are anti- and pro-longevity, respectively (Honjoh and Nishida, 2011). RHEB-1 and TOR have also been implicated as repressors of the mitochondrial unfolded protein stress response (Haynes et al., 2007), which also impacts lifespan (Bennett and Kaeberlein, 2014).

As noted in Section 2.7, *C. elegans* expresses all of the components of TOR system signaling except the tuberous sclerosis complex (TSC) RhebGAP (Long et al., 2002). TSC is present in *Drosophila* and all other metazoans besides nematodes; *S. pombe* has TSC while *S. cerevisiae* does not, but all of these organisms express Rheb, and Rheb loss is consistent with TOR loss biologically (van Dam et al., 2011). As noted in Section 2.7, the *C. elegans* HGAP complex could functionally replace the TSC/RhebGAP as well as function as a RalGAP. The decreased lifespan caused by disruption of **HGAP-1** or **HGAP-2** is consistent with the effect expected from derepressed **RHEB-1**.

The divergence of *C. elegans* **RHEB-1** and absence of TSC/RhebGAP raises the question of whether Rheb function is conserved with other species. Deletion of *C. elegans* TORC1 components TOR and Raptor resulted in mid-larval arrest with an atrophied gut and RNAi depletion extended lifespan (Hara et al., 2002; Long et al., 2002; Vellai et al., 2003; Jia et al., 2004). Mutants for *rheb-1* have not been described.

Mutating most Ras family members to a constitutively activate GTP-bound form is simple: mutate the G12, G13, or Q61 oncogenic Ras mutations (LET-60/Ras numbering). Not so with Rheb, because these positions are not conserved, and Rheb structure is atypical for the Ras family. In vitro mammalian Rheb1 is unusually highly GTP-bound, though GTP hydrolysis can be stimulated by addition of the TSC RhebGAP. Introducing the Q64L (Q61L in Ras numbering) mutation to Rheb1 confers a 2-3-fold increase in activity, much lower than activation conferred by homologous changes in other Ras family members (Inoki et al., 2003). The search for activating Rheb mutations in various systems has yielded only weak mutations (Mazhab-Jafari et al., 2014). With the high level of sequence divergence of *C. elegans* **RHEB-1**, it is uncertain whether the corresponding mutations would increase activity level. Ectopic expression of *C. elegans* RHEB-1(Q71L,Q72D) under control of its own promoter, an attempt to mutationally activate **RHEB-1**, confers weak TORC1 activation in an assay of rescue of fatty acid depletion (Zhu et al., 2013). Activation of RagA (see Section 7), another major input into TORC1, robustly rescued fatty acid depletion, suggesting that RHEB-1(Q71L,Q72D) is not constitutively active.

Dominant-negative mutations are useful for studying small GTPases function (see above) (Han and Sternberg, 1991). Canonical Ras dominant-negative mutation in mammalian Rheb1-S20N (S17N by Ras numbering) has no effect (Inoki et al., 2003), which, with the relatively high intrinsic Rheb1-GTP state could argue that Rheb1 does not require a GEF. *Drosophila* TCTP has properties expected from a RhebGEF (Hsu et al., 2007), with modest support from mammalian cells (Dong et al., 2009). In other contexts the mammalian TCTP does not perform RhebGEF activity or have genetic characteristics expected of RhebGEF in cell culture (Rehmann et al., 2008; Wang et al., 2008). Therefore it is still unknown whether there exists a RhebGEF in any system.

2.9. DRN-1

drn-1 encodes the *C. elegans* ortholog of mammalian Di-Ras1/Rig, Di-Ras2, and Di-Ras3/Noey2/ARHI. The Di-Ras subfamily shares typical sequence features of the Ras family, with some differences that may confer Rap-like GAP specificity. The Rap family has a Thr at position 61 instead of the Gln of Ras (Ras numbering; see Section 2.4) and the Di-Ras family, including **DRN-1**, has a Ser at 61. Not surprisingly, mammalian Rap1GAP stimulates Di-Ras1 GTP hydrolysis, while typical RasGAPs do not (Gasper et al., 2010). The *C. elegans* Rap1GAP ortholog **F53A10.2**, and perhaps other Rap-specific GAPs may therefore also function as a GAP for DRN-1.

Di-Ras subfamily effector-binding domains contain modest but presumably important differences from the rest of the Ras family (Figure 5), suggesting that different signaling partners are used. The mammalian Di-Ras family members are all tumor suppressor genes (typically brain, breast, and ovarian), which is a departure from typical oncogenic role of Ras in cancer.

C. elegans **DRN-1** is broadly expressed in neurons. The *ok400*, *tm2916*, and *tm2926* deletion alleles, all putative nulls due to dramatic disruption of conserved protein structure, confer resistance to the acetylcholine esterase inhibitor aldicarb, consistent with decreased cholinergic synaptic release in the mutants. This phenotype was transgenically rescued by putative gain-of-function, but not dominant-negative, *drn-1* constructs (Tada et al., 2012). Genetic interactions with $G\alpha_o$, $G\alpha_q$, and $G\alpha_s$ heterotrimeric G-protein α subunit mutants, all shown to regulate synaptic release, are consistent with **DRN-1** functioning downstream of **GSA-1/G α** , which stimulates cAMP production. Deletion of the cAMP-dependent RapGEF *epac-1* also confers aldicarb resistance (Tada et al., 2012), and thus by association may be a **DRN-1** GEF. In the absence of known Di-Ras subfamily effectors, it is unclear whether the neurobiological phenotypes of *drn-1* mutants relate to the tumor suppressor phenotypes caused by loss of subfamily members in mammals.

A singular feature of *drn-1* is the organization of the *drn-1* transcripts. The *drn-1* gene as annotated in Wormbase (WS247 and earlier versions) predicts two 5'UTR variants with different transcription initiation sites, one spliced and one not. However, using the first AUG start codon in hypothetical translation results in a 125 residue protein truncating the N-terminus of the GTPase domain and disrupting putative GTPase structure and effector binding. Mammalian Di-Ras subfamily members, like Ras itself, have an approximately 170 residue core GTPase domain, plus N- and C-termini extensions. There is no precedent for Ras family members with a truncated N-terminus; not only does that abolish guanine nucleotide cycling, it disrupts the protein structure used to interact with other proteins. Despite the absence of a conventional protein product, *drn-1* mutations confer a neuronal phenotype. Thus, *drn-1* is unlikely to be a pseudogene.

Notably, sequences 5' of the first putative AUG, if translated, would produce a hypothetical N-terminus of DRN-1 homologous to mammalian Di-Ras subfamily members: the entire GTPase domain would be conserved (Figure 9). Transcript *C54A12.4.2*, based on RNAseq and RT-PCR data (V. Muñiz-Medina and D. Reiner, unpublished results), contains a long putative 5'UTR. There are no AUG or termination codons in frame upstream of the annotated AUG. We hypothesize that non-canonical translation initiation codons are used to produce a protein with the entire conserved GTPase domain: the minimal size based on sequence conservation and Ras family N-termini is a 192 residue DRN-1 protein. This hypothetical protein, the “minimal DRN-1 protein,” has a functional GTPase domain, conservation of the entire GTPase sequence, and a function C-terminal membrane-targeting region. Therefore, we speculate that non-AUG translation initiation is used to express DRN-1.

Di-Ras1	-----	MPEQSNDYRVVVFGAGGVG
Di-Ras2	-----	MPEQSNDYRVAVFGAGGVG
Di-Ras3	---MGNASFGSKEQ KLL ---	KRLRLLPALLILRAFKPHRKIRDYRVVVVGTVAGVG
CeDRN-1	SAFQSSPTASNGGSPR LL LDTDFSIEATAATTAGSGSKVAEASTSDYRVAVFGAGGVG	
CbrDRN-1	SAFQSSPTASNGGSPR LL EADFSIEDTAATTAGSGSRVAEASTSDYRVAVFGAGGVG	
CbnDRN-1	FLQSPLPKKECSTPSH LL LETDSSEATATNTTAGSGSRVAEASTSDYRVAVFGAGGVG	
 Plausible CeDRN-1 start codon: CUGCUG (Leu-Leu)		
Plausible CbrDRN-1 start codon: CUGCUG (Leu-Leu)		
Plausible CbnDRN-1 start codon: CUACUG (Leu?-Leu)		

Figure 9. Alignment of human and Caenorhabditid Di-Ras subfamily predicted N-termini. GTPase domain sequences are in blue, established mammalian sequences in black, the inferred DRN-1 N-terminal extension in purple, and ambiguous further N-terminal extension in gray. Dileucine repeats are bolded. Di-Ras3 (ARHI, Noey2) has an N-terminal extension containing three dileucine motifs. Assuming a non-AUG initiation codon, transcript *C54A12.4.2*, predicted by RNAseq, aligns with upstream genomic sequence in *C. briggsae* (Cbr) and *C. brenneri* (Cbn), all of which could start with a dileucine repeat. Hypothetical translation of the overlapping transcript *C54A12.4.1* also contains a dileucine, but it precedes a stop codon and so it may be a computational artifact. Detailed analysis of the *drn-1* gene structure may resolve these questions.

Though rare, there is a precedent for non-AUG initiation across many organisms (Ivanov et al., 2011). Where known, non-AUG codons generally follow the template of “NUG,” with “CUG” being a common initiator. Leaky translation initiation mutants in *C. elegans* use start codons with similarity to “AUG,” with “CUG” being among the most efficient (Maduzia et al., 2010; Zhang and Maduzia, 2010). A “CUGCUG” sequence lies in the *C54A12.4.2* transcript in frame with the annotated AUG and with no intervening terminators, resulting in a hypothetical DRN-1 containing an N-terminal extension starting with LL dual leucine (Figure 9). The mammalian Di-Ras3/Noey2/ARHI protein has an N-terminal extension with three LL pairs. Perhaps there is a functional parallel between Di-Ras3/Noey2/ARHI and DRN-1 putative N-terminal extensions. The DRN-1 ortholog in other *Caenorhabditis* species suggests a long N-terminal extension with highly identical core Di-Ras sequences. The Wormbase entry for both *C. briggsae* and *C. brenneri* DRN-1, chosen because of available RNAseq data, predict the same gene organization: the first AUG sequence would result in a truncated protein, but alternate initiation codons would produce proteins conserved throughout their length with CeDRN-1.

Transcript *C54A12.4.1*, which partially overlaps *C54A12.4.2*, is not supported by RNAseq or RT-PCR data, so may be a computational artifact. *C54A12.4.1* contains a long 5'UTR without the CUGCUG repeat. The 5'UTR does contain the sequences necessary to generate the hypothetical “minimal DRN-1 protein” that so closely aligns with the mammalian Di-Ras subfamily members.

2.10. SSR-2

ssr-2 encodes the *C. elegans* ortholog of mammalian Dexamethasone-induced Ras protein 1/RasD1/Dexras1 and Dexamethasone-induced Ras protein 2/RasD2/Dexras2/RHES. RasD1/2 are widely expressed, and expression is

regulated by diverse hormones and stress response (Thapliyal et al., 2014). No functional data for **SSR-2** have been reported. Because it is more distantly related, we did not include **SSR-2** in alignments.

3. The Rho family

Rho-family GTPases in *C. elegans* and other systems have roles in cytoskeletal organization, cell polarity, cell migration, and cell membrane protrusion. Classic studies from the laboratory of Alan Hall, using mammalian cell culture, established that activated Rho promotes formation of stress fibers and focal adhesions (Ridley and Hall, 1992), activated Rac promotes lamellipodia and membrane ruffles (Ridley et al., 1992), and activated Cdc42 promotes spike-like filopodia (Nobes and Hall, 1995). The discovery of enormously diverse GEFs for the Rho family (Rossman et al., 2005), plus many subsequent analyses of their functions, identified the importance of tight spatial regulation of Rho, Rac, and Cdc42 function by combinatorial activity of their GEFs for proper control of cytoskeletal activities and consequent morphogenetic actions (Goicoechea et al., 2014; Lawson and Burridge, 2014). Like the GEFs for the Rho family, GAPs for the Rho family are also quite diverse, but are under-studied.

Structurally, Rho family members are distinguished by the “Rho family insert” between β -strand 5 and α -helix 4 (Valencia et al., 1991). The *C. elegans* genome encodes seven members of the Rho family of GTPases, and effector-binding regions are typically highly conserved, with some exceptions (Figure 10). Canonical members of the Rho-family GTPases include Rho (*rho-1*), Rac (*ced-10*), and Cdc42 (*cdc-42*). The *rac-2* gene is nearly identical to *ced-10*, but it is unclear if *rac-2* is a functional gene or a non-functional duplication of the *ced-10* locus, although RNAi of *rac-2* enhances *ced-10* (Struckhoff and Lundquist, 2003). *mig-2* encodes an Mtl (MIG-2-like) GTPase, a family found in invertebrates with similarity to both Rac and Cdc42. Functionally, **MIG-2** is similar to mammalian RhoG (deBakker et al., 2004). Two additional atypical Rho-like GTPases are **CHW-1**, which is similar to the Chp/Wrch family of GTPases in vertebrates (Kidd et al., 2015), and **CRP-1**, a Cdc42-like protein with a divergent effector-binding region (Jenna et al., 2005) (Figure 10).

Cdc42	TNKFPSE <u>YVPTVFDNY</u> AVTVMIGGE
TCL	NDAFPEE <u>YVPTVFDHY</u> AVTVTVGGK
TC10	NDAFPEE <u>YVPTVFDHY</u> AVSVTVGGK
CDC-42	TNKFPSE <u>YVPTVFDNY</u> AVTVMIGGE
CRP-1	VFDNWAV <u>SVHIDNKNY</u> SVHIDNKNY
Wrch1	TNGYPTE <u>YIPTAFDNF</u> SAVSVVDGR
Chp	CNGYPARY <u>RPTALDTF</u> SVQVLVDGA
CHW-1	TNVYPHN <u>YVPTAFDNF</u> SVVVLVDKK
Rac1	TNAFPGE <u>YIPTVFDNY</u> SANVMVDGK
Rac2	TNAFPGE <u>YIPTVFDNY</u> SANVMVDGK
Rac3	TNAFPGE <u>YIPTVFDNY</u> SANVMVDGK
CED-10	TNAFPGE <u>YIPTVFDNY</u> SANVMVDGK
RAC-2	TNAFPGE <u>YILTVFDTY</u> STNVMVDGR
RhoG	TNAFPKE <u>YIPTVFDNY</u> SAQSAVDGR
MIG-2	TDSFPVQ <u>YVPTVFDNY</u> SAQMSLDGN
RhoA	KDQFPEV <u>YVPTVFENY</u> VADIEVDGK
RhoB	KDEFPEV <u>YVPTVFENY</u> VADIEVDGK
RhoC	KDQFPEV <u>YVPTVFENY</u> IADIEVDGK
RHO-1	KDQFPDV <u>YVPTVFENY</u> VADIEVDGK

Figure 10. Effector-binding region of human vs. *C. elegans* Rho family members. The entire effector-binding region of human and *C. elegans* proteins was aligned. The core sequence, thought to govern direct protein-protein interactions, is underlined. Core residues not conserved in the subfamily are bolded and blue. Human orthologs are listed above *C. elegans* orthologs. *C. elegans* protein names have a dash and number. Some human proteins have alternate names (see Table 1).

Rho GTPases are key regulators of actin cytoskeleton dynamics, cell shape, and cell polarity. Based on four parameters of biochemical interaction with GTP—loading, GTP??S competition, hydrolysis, and exchange—the Rho GTPases were clustered according to functional similarity (Caruso et al., 2005). When all four traits were considered together, **RHO-1**, **CDC-42**, and **CED-10** were in one functional cluster, with **CDC-42** and **CED-10** being closer together, and **MIG-2** and **CRP-1** were in another, suggesting that **CRP-1** and **MIG-2** have similar biochemical properties. By similarity of sequence, **MIG-2** was closer to **CED-10** and **CDC-42**, followed by **RHO-1** and **CRP-1**. **CHW-1** was not included in this analysis. Specific roles of the different Rho GTPases are described below.

3.1. RHO-1

Loss of *rho-1* leads to embryonic lethality, and *rho-1* function has been ascertained using RNAi and by transgenic expression of constitutively active and dominant negative forms of *rho-1*. Conditional perturbation of *rho-1* function in adults results in dysfunction in numerous neuronal and non-neuronal functions and leads to death (McMullan and Nurrish, 2011), highlighting the central role of the RHO-1 GTPase in many biological processes throughout the life of the organism.

3.1.1. Embryonic polarity and cytokinesis

RNAi-mediated knockdown of *rho-1* activity resulted in early embryonic arrest, often with a failure of cytokinesis (Jantsch-Plunger et al., 2000; Spencer et al., 2001; Bringmann and Hyman, 2005; Morita et al., 2005; Motegi and Sugimoto, 2006). Embryos that arrested later in development displayed severe defects in tissue morphogenesis. These phenotypes could be explained by defects in Rho-mediated actin cytoskeleton organization (e.g., actin forms the contractile ring during cytokinesis and is involved in cellular migration and morphogenesis) (Schonegg and Hyman, 2006). Other mutations that affect embryonic morphogenesis and cytokinesis identified other genes that act with *rho-1* in this process. *let-502* encodes a Rho kinase (mammalian ROCK or ROK), *mel-11* encodes a myosin phosphatase (Wissmann et al., 1997; Wissmann et al., 1999; Piekny and Mains, 2002), and RHO-1 acts with the actin and myosin binding protein Anillin (ANI-1) in cytokinesis (Fotopoulos et al., 2013). These molecules likely affect actin-myosin cytoskeletal dynamics of the contractile ring during cytokinesis in response to Rho signaling (Spiga et al., 2013). Furthermore, RHO-1 is involved in establishing and maintaining the position of the PAR complex proteins in early embryonic polarity (Jenkins et al., 2006; Schonegg and Hyman, 2006; Anderson et al., 2008; Canman et al., 2008; Nakayama et al., 2009).

GAPs and GEFs also regulate RHO-1 in polarity and cytokinesis. The *cyk-4* gene, identified in a screen for mutations affecting embryonic cytokinesis, encodes a Rho GAP that likely modulates RHO-1 activity in establishment of embryonic polarity (Jantsch-Plunger et al., 2000; Jenkins et al., 2006; Canman et al., 2008). The Rho GAPs RGA-3, RGA-4, and PAC-1, and the RHGF-2 GEF act with RHO-1 in cytokinesis (Schmutz et al., 2007; Schonegg et al., 2007; Anderson et al., 2008; Chan et al., 2015). The ECT-2 GEF (*Drosophila* Pebble) acts with RHO-1 in embryonic polarity and cytokinesis (Morita et al., 2005; Motegi and Sugimoto, 2006).

3.1.2. Embryonic elongation

LET-502 and MEL-11 also control embryonic elongation. After epiboly and ventral enclosure, hypodermal cells that surround the embryo contain circumferential rings of actin that contract, resulting in a “squeezing” of the embryo and a subsequent increase in length. LET-502 and MEL-11 have opposing roles in this process: LET-502 normally enhances contraction and MEL-11 normally inhibits contraction. (Wissmann et al., 1997; Wissmann et al., 1999) (Figure 11). The RhoGAP RGA-2 negatively regulates LET-502 in this process (Diogon et al., 2007). RHGF-2 is the corresponding Rho GEF: even though both *rga-2* and *rhgf-2* alone are lethal, the double mutant is viable (Chan et al., 2015). Interestingly, the Mtl Rac GTPase MIG-2 and the UNC-73 Trio Rac/Rho GEF act in the MEL-11 pathway in this process. UNC-73 Trio acts as a GEF for RHO-1, CED-10, and MIG-2 (Steven et al., 1998; Spencer et al., 2001; Wu et al., 2002). These results indicate that MIG-2 Mtl Rac and RHO-1 might have opposing roles in embryonic elongation (Piekny et al., 2000) (Figure 11). The Rac CED-10, the Rac GEF PIX-1, and PAK-1/PAK regulate muscle hemidesmosome formation, which mediate the latter part of elongation after hypodermal contraction (Zhang et al., 2011). PAK-1 acts in parallel to LET-502 (Gally et al., 2009), and PAK-1 might be regulated by PIX-1 during elongation (Martin et al., 2014).

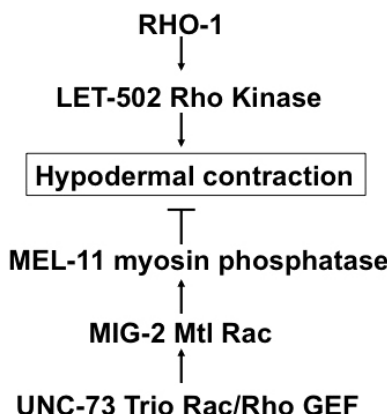


Figure 11. The opposing roles of RHO-1 and MIG-2 Rac in hypodermal contraction during embryonic elongation. Circumferential actin bundles in the hypodermis contract to drive embryonic elongation. RHO-1 stimulates hypodermal contraction via the LET-502 Rho-binding kinase, and MIG-2/Mtl/Rac inhibits hypodermal contraction via MEL-11 myosin phosphatase in response to UNC-73 Trio Rac/Rho GEF.

3.1.3. P-cell migration

Later in development, *rho-1* controls the ventral migrations of the P cells (Spencer et al., 2001). The P cell nuclei are born in sublateral locations and migrate ventrally to align at the ventral midline. RHO-1 controls migration of the P cells along with the GEF UNC-73/Trio and LET-502 (Steven et al., 1998; Spencer et al., 2001). Interestingly, the Rac GTPases CED-10 and MIG-2 control P-cell migration in a parallel redundant manner with RHO-1 (Spencer et al., 2001). ECT-2 activation of RHO-1 also controls migration of P cells as well as in induction of the 1° vulval cell fate in vulval patterning (Canevascini et al., 2005; Morita et al., 2005; Mitin et al., 2007; Cook et al., 2014).

3.1.4. Neuronal morphogenesis and function.

RHO-1 also controls neuronal cell shape after the establishment of the normal axon and dendrite morphology of the neuron. Expression of dominant-negative RHO-1(T19N) in the ASE sensory neuron resulted in an expanded cell body morphology and ectopic neurite initiation (Zallen et al., 2000). This phenotype is similar to that caused by loss of function of the SAX-1 Ndr kinase (Zallen et al., 2000), a known downstream target of Rho activity in other systems. RHO-1 and SAX-1 might act together to regulate neuronal morphogenesis. The RhoGEF RHGF-2 inhibits neurite outgrowth when over-expressed in neuroblastoma cells, similar to Rho constitutive activation (Lin et al., 2012). RHO-1 also regulates neurotransmitter release (Steven et al., 2005; Hiley et al., 2006; McMullan et al., 2006; Hu et al., 2011). The RhoGEF OSG-1 sensitizes neurons to oxidative stress, which might be part of the aging process (Duan and Sesti, 2015). RHO-1 may also cooperate with a non-canonical LET-60/Ras function in response of the rectal epithelium to pathogen (McMullan et al., 2012).

3.2. CDC-42

CDC-42 is required for an array of developmental events involving the cytoskeleton, cell polarity, and protrusion. The roles of CDC-42 in some of these events are described below. Two themes emerge from these studies: a conserved signaling module of CDC-42 with the polarity proteins PAR-3/PAR-6/PKC-3 is iteratively used in multiple developmental events, and CDC-42 acts upstream of the Rac GTPases CED-10 and MIG-2 in protrusive events in a Rho GTPase hierarchy of signaling.

3.2.1. Embryonic polarity

cdc-42-directed RNAi caused defects in embryonic cytokinesis similar to *rho-1(RNAi)* (Kay and Hunter, 2001). *cdc-42(RNAi)* also perturbed the polarity of the single-celled zygote and resulted in defects in anterior-posterior axis formation and mitotic spindle orientation (Gotta et al., 2001; Kay and Hunter, 2001). Upon fertilization, sperm entry results in a rearrangement of the zygote cytoplasm such that anterior cortex accumulates PAR-3 and PAR-6 and the posterior cortex accumulates PAR-2, which together define the anterior-posterior axis of the organism (Levitin et al., 1994; Etemad-Moghadam et al., 1995; Watts et al., 1996; Hung and Kemphues, 1999). The PAR proteins associate with the actin cytoskeleton at the cell cortex, and the actin cytoskeleton is required for their localization. As a result of PAR gene localization, the first cell division generates a larger anterior cell and a

smaller posterior cell, whose spindle rotates 90° in relation to the anterior cell (Kemphues et al., 1988). Mutations in the *par* genes disrupt zygotic polarity and result in the symmetric division of zygote and defects in spindle orientation. RNAi of *cdc-42* results in a similar loss of zygotic polarity and an associated failure to localize the PAR proteins to their proper domains (Gotta et al., 2001; Kay and Hunter, 2001). *cdc-42* controls the localization of the PAR proteins via acto-myosin contractility along with RHO-1 (Schonegg and Hyman, 2006). **CDC-42** physically associates with **PAR-6**, a CRIB- and PDZ-domain-containing component of a conserved complex including **PAR-3**, and **PKC-3**, a protein kinase C ortholog, that defines the anterior cortical domain in the zygote (Gotta et al., 2001; Aceto et al., 2006). The **CDC-42** GEF **CGEF-1** and the **CDC-42** GAP **CHIN-1** act with **CDC-42** in embryonic polarity (Kumfer et al., 2010; Beatty et al., 2013). A conserved **CDC-42/PAR-3/PAR-6/PKC-3** module acts in multiple cellular events beyond embryonic polarization, including neuronal morphogenesis and cell migration (Welchman et al., 2007).

As cell division proceeds, blastomeres distinguish between inner surfaces, characterized by cell-cell contact, and outer surfaces, indicated by lack of cell-cell contact. This process is known as radial polarity. **CDC-42** activity is inhibited by the RhoGAP **PAC-1** at sites of blastomere contact with one another (Anderson et al., 2008). The RhoGEFs **CGEF-1** and **ECT-2** activate **CDC-42** at the cortex of contact-free blastomere surfaces (Chan and Nance, 2013). This results in **PAR-6** recruitment to the cortex at no-contact sites, and exclusion from contact sites, to maintain radial polarity.

3.2.2. Ventral hypodermal enclosure

During *C. elegans* gastrulation, the dorsally located hypodermal cells migrate toward the ventral midline, where they meet and completely surround the embryo in a process known as ventral enclosure (Sulston et al., 1983; Williams-Masson et al., 1997; Simske and Hardin, 2001). *C. elegans* **WSP-1** is the homolog of the human Wiskott-Aldrich syndrome protein (WASP). WASP is a downstream effector of Cdc42 that activates the Arp2/3 complex, an actin nucleating and branching complex. **wsp-1** mutants display a low-penetrance embryonic lethality due to a failure of migration of the hypodermal cells and ventral enclosure (Withee et al., 2004). The **CDC-42** interacting FBAR proteins **TOCA-1** and **TOCA-2** also regulate embryonic morphogenesis (Giuliani et al., 2009).

3.2.3. Axon guidance and neuronal morphogenesis

UNC-6/Netrin and its receptors **UNC-40/DCC** and **UNC-5** are a conserved dorsal-ventral axon guidance system that regulates both attractive guidance ventrally, and repulsive guidance dorsally. In ventrally directed axons attracted to **UNC-6**/Netrin, **UNC-40/DCC** promotes neuronal protrusion from cell bodies and growth cones (Norris and Lundquist, 2011; Demarco et al., 2012). **CDC-42** is required for **UNC-40/DCC**-mediated protrusion, and also affects the guidance of axons attracted to **UNC-6**/Netrin (Demarco et al., 2012). In protrusion, **CDC-42** acts upstream of the Rac GEF **TIAM-1** and the Rac GTPases **CED-10** and **MIG-2** (Demarco et al., 2012), which modulate protrusion via the Arp2/3 complex and the Arp2/3 complex activators **WAVE-1** and **WSP-1**, respectively (Shakir et al., 2008; Norris et al., 2009).

Consistent with a role in neuronal protrusion, constitutive activation of **CDC-42** in neurons results in ectopic protrusive structures dependent upon Rac GTPases and the Arp2/3 complex (Alan et al., 2013). PI3 kinase signaling (e.g., **AGE-1/PI3K** and **AKT-1/AKT**), and **RICT-1/mTORC2** were also required for protrusion, whereas mTORC1 signaling components (**DAF-15/RAPTOR** and **DAF-16/FOXO**) were not (Alan et al., 2013). **CDC-42** regulation of protrusion utilizes a complex network of signaling pathways involving Rac GTPases as well as PI3K and mTORC2 signaling (Figure 12).

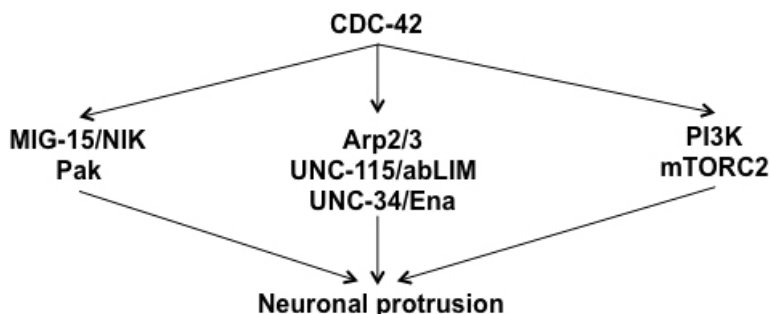


Figure 12. *CDC-42* in neuronal protrusion. *CDC-42* utilizes multiple pathways to stimulate neuronal protrusion, including the actin-interacting molecules Arp2/3, **UNC-34/Enabled**, and **UNC-115/abLIM**, the kinases **MIG-15/NIK** and **PAK-1**, and PI3K signaling via the mTORC2 complex.

3.2.4. Apoptotic cell corpse engulfment by muscles

Apoptotic cells are cleared by phagocytosis by neighboring cells. Signals from dying cells induce the formation of phagocytic protrusions from neighboring cells, which involve the Rac GTPases **CED-10** and **MIG-2** (see Section 3.3). **CDC-42** is also required for cell corpse engulfment (Hsieh et al., 2012; Neukomm et al., 2014). In muscles, the α integrin **PAT-2** acts with **CED-1/CD91**, **CED-6/GULP**, and **CED-7/ABC transporter** transmembrane protein engulfment module (Neukomm et al., 2014). **CDC-42** acts downstream of **PAT-2** in engulfment, and upstream of **CED-10/Rac** (Hsieh et al., 2012; Neukomm et al., 2014). The GEF **UIG-1** acts with **CDC-42** in engulfment (Hsieh et al., 2012; Neukomm et al., 2014). Interestingly, **UIG-1** and **CDC-42** are also required for the structure of integrin-containing dense bodies that attach the myofilament lattice to the muscle cell plasma membrane (Hikita et al., 2005).

3.2.5. Retrograde endocytic recycling

CDC-42 is involved in trafficking from recycling endosomes to the trans-Golgi network in retrograde endocytic recycling (Bai and Grant, 2015). This also involves the PAR complex (e.g., **PAR-6** and **PKC-3**) and **TOCA-1,2** as well as the Arp2/3 activator **WVE-1**. Components of Wnt secretion utilize this pathway, and thus **CDC-42**, can affect Wnt signaling via retrograde endocytic recycling (Bai and Grant, 2015).

3.2.6. Notch signaling

LIN-12/Notch is necessary and sufficient for 2° cell fate specification during developmental patterning of the vulva. Genetically, **CDC-42** acts as a negative regulator of **LIN-12** function in VPC fate specification (Choi et al., 2010), although the molecular nature of this interaction is unclear.

3.3. Rac-like GTPases **CED-10** and **MIG-2**

ced-10 encodes a Rac1-like molecule, and *mig-2* encodes an Mtl Rac (Zipkin et al., 1997; Reddien and Horvitz, 2000; Lundquist et al., 2001). **MIG-2** has similarity to both Cdc42 and Rac molecules, and has an N-terminal predicted myristylation sequence not found in Cdc42 or Rac. In cell corpse engulfment, **MIG-2** is the functional equivalent of vertebrate RhoG (deBakker et al., 2004). *ced-10* mutations were identified in mutant screens for defective apoptotic cell corpse phagocytosis (Ellis et al., 1991), and *mig-2* mutations were identified in screens for cell migration-defective mutants (Zipkin et al., 1997). The *rac-2/3?* locus is an apparent tandem duplication, and two deletions (*ok326* and *gk281*) each affect only one copy. *rac-2/3* loss-of-function induced by RNAi had no apparent phenotypic consequence (Lundquist et al., 2001; Struckhoff and Lundquist, 2003). It is possible that the *rac-2/3* locus is non-functional. However, *rac-2/3* RNAi did enhance *ced-10* and *mig-2*, suggesting it might have some function. This section will focus on the roles of **CED-10** and **MIG-2**. While *ced-10* and *mig-2* mutations alone affect some developmental processes (e.g., phagocytosis and cell migration), analysis of pairwise *Rac* double mutant combinations revealed processes in which *ced-10* and *mig-2* have overlapping and compensatory roles (i.e., in which they act redundantly). Therefore, the roles of **MIG-2** and **CED-10** will be discussed together.

3.3.1. Cell corpse phagocytosis

ced-10 mutations alone cause failure of phagocytosis of cell corpses after programmed cell death (Ellis et al., 1991; Reddien and Horvitz, 2000). Upon programmed death of a cell, neighboring cells extend actin-based plasma membrane protrusions to engulf and metabolize the cell corpse. *ced-10* mutants fail in this process, resulting in the persistence of unengulfed cell corpses. *ced-10* might be required to organize the actin cytoskeleton underlying the extension of cellular protrusions of the engulfing cell. Other engulfment mutations identified the *ced-1*, *ced-2*, *ced-5*, *ced-6*, *ced-7*, and *ced-12* genes (Ellis et al., 1991). These genes can be placed into two pathways based upon redundancy of function in engulfment: *ced-1*, *-6*, and *-7* function in parallel to *ced-2*, *-5*, *-10*, and *-12*. Double mutant combinations between groups cause enhanced engulfment defects, whereas double mutant combinations within each group do not enhance. This observation suggests that two parallel pathways exist to control phagocytosis, one of which involves *ced-10 Rac*.

Molecular and genetic analyses revealed other **CED-10** pathway members in phagocytosis (Figure 13), including **CED-2**, a CrkII-like receptor adapter protein (Reddien and Horvitz, 2000), and **CED-5**, a DOCK/CDM-family Rac GEF (Wu and Horvitz, 1998). Furthermore, **CED-12** is an ELMO-like protein, which cooperates with CDM proteins in Rac GTP exchange (Gumienny et al., 2001; Zhou et al., 2001; Lu et al., 2004). Indeed, in vertebrate cells and *C. elegans*, **MIG-2** and its exchange factor **UNC-73/Trio** act upstream of

CED-5, which in turn activate CED-10 (deBakker et al., 2004). The CED-10/Rac GAP SRGP-1 negatively regulates CED-10/Rac in engulfment (Neukomm et al., 2011), and PDR-1/Parkin negatively regulates CED-10/Rac in phagocytosis possibly via ubiquitylation and targeting CED-10 for degradation (Cabello et al., 2014).

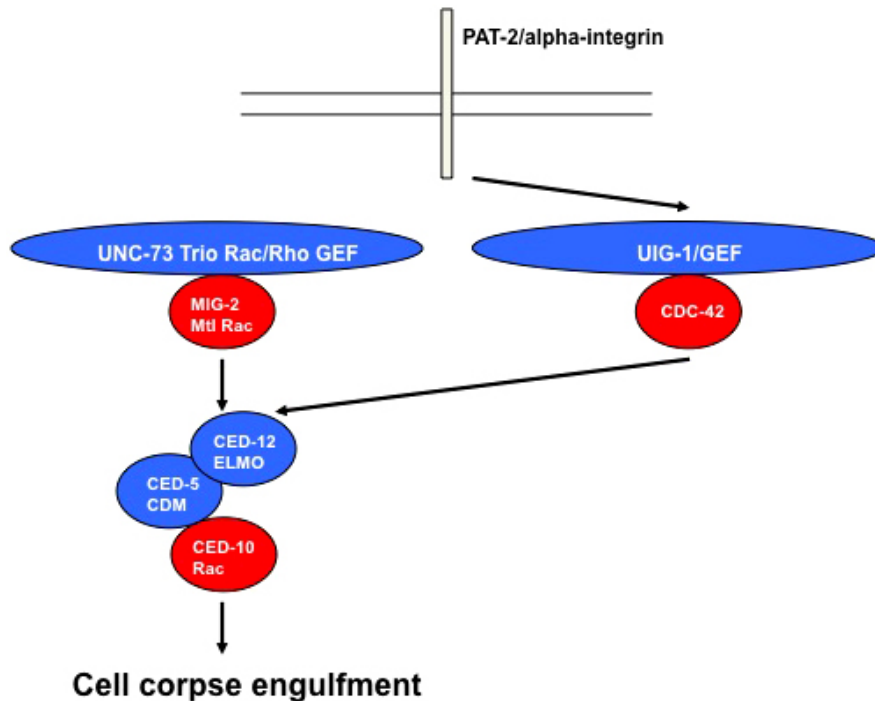


Figure 13. Rho GTPases in programmed cell death corpse engulfment by muscles. CED-10/Rac acts downstream of CDC-42 and the alpha-integrin PAT-2 in extension of cell protrusions in corpse engulfment. MIG-2/RhoG also acts upstream of CED-10/Rac along with the UNC-73/Trio GEF. This pathway likely acts in parallel to the CED-1/6/7 complex.

The CED-1/(CD91/LRP/SREC)/CED-6/(hCED-6/GULP)/CED-7(ABCA1) complex acts in parallel to the CED-10 pathway to regulate actin dynamics underlying phagocytic protrusion (Kinchen et al., 2005). Both pathways might converge on the nucleoside diphosphate kinase NDK-1 in this process (Fancesalszky et al., 2014). A third parallel pathway involves the ABL-1/Abl tyrosine kinase (Hurwitz et al., 2009), illustrating the complex signaling network used to regulate protrusion during phagocytosis.

3.3.2. Cell migration

As key regulators of cytoskeletal dynamics and protrusion, *ced-10* and *mig-2* mutations affect the migration of the gonadal distal tip cells (DTCs), which results in misshapen gonad arms (Lundquist et al., 2001). The dorsalward migration of the DTCs involve UNC-6/Netrin and the UNC-5 receptor (Su et al., 2000). Many of the molecules that act with CED-10 and MIG-2 in phagocytosis are conserved in DTC migration, including the CED-5/CED-12 module (but not CED-2/CrkII), NDK-1, UNC-73/Trio, and PDK-1/Parkin (Gumienny et al., 2001; Lundquist et al., 2001; Neukomm et al., 2011; Cabello et al., 2014). The p21-activated kinase MAX-2 might act downstream of CED-10 in DTC migration (Lucanic et al., 2006; Lucanic and Cheng, 2008; Peters et al., 2013), and the BED domain-containing protein MIG-39 acts with MIG-2 and CED-10 to halt migration of the DTCs (Kikuchi et al., 2015). Furthermore, CACN-1/Cactin acts with MIG-2 in parallel to CED-10 in DTC migration and cessation of migration (Tannoury et al., 2010).

CED-10 and MIG-2 also regulate the migration of the Q neuroblasts and their descendants, along with the NIK kinase MIG-15 and UNC-34/Enabled (Shakir et al., 2006). MIG-15/NIK controls the direction and ability of the Q neuroblasts to extend protrusions and to migrate (Chapman et al., 2008). CED-10 and MIG-2 and their GEFs have both unique and redundant roles in Q migration. MIG-2 and CDC-42, along with the Rac GEFs UNC-73/Trio and PIX-1/PIX are required for robust protrusions of the Q cells preceding migration, whereas CED-10 is required to limit extent of protrusion, similar to its role in DTC migration (Dyer et al., 2010). Double mutant analysis indicates redundancy between MIG-2 and CED-10 in extension of robust protrusions, and that PIX-1/Pix might act in the CED-10 pathway in parallel to UNC-73/Trio and MIG-2 (Dyer et al., 2010). MIG-2 activity correlates with faster and longer Q descendant cell migration due to its role in maintaining persistent cell and cytoskeletal

polarization in the direction of migration (Ou and Vale, 2009). The migration of the CAN neurons is also redundantly controlled by **MIG-2** and **CED-10**, along with the GEF **UNC-73/Trio** (Zipkin et al., 1997; Lundquist et al., 2001).

3.3.3. Gastrulation and morphogenesis

Null alleles of ***ced-10*** cause maternal-effect embryonic lethality with defects in early epidermal cell movements in gastrulation, resulting in failure of the endoderm to be surrounded by the ectoderm (the **gut on the exterior** (**Gex**) phenotype) (Lundquist et al., 2001; Soto et al., 2002). Mutations in ***gex-1***, ***gex-2***, and ***gex-3*** cause a similar maternal-effect lethal phenotype (Soto et al., 2002). **GEX-2** encodes an Sra-1-like molecule, and **GEX-3** encodes a Hem2-like molecule. Sra-1 and Hem2, along with Nck, form a complex with the Arp2/3 complex activator WAVE (Eden et al., 2002). WAVE-1/WVE acts with **GEX-2** and **GEX-3** in epidermal gastrulation movements, likely downstream of **CED-10** (Patel et al., 2008). This complex leads to the subcellular enrichment of F-actin at apical or basolateral regions in different epidermal cells (Patel et al., 2008).

CED-10 and **MIG-2** act redundantly in multiple aspects of vulval development and morphogenesis. ***mig-2*** and ***ced-10*** single mutant vulvae are largely wild-type, whereas ***mig-2*; *ced-10*** double loss-of-function mutants display defects in the orientation of asymmetric divisions of the 1° and 2° vulval cells, indicating that ***mig-2*** and ***ced-10*** might redundantly control spindle orientation (Kishore and Sundaram, 2002). ***mig-2*; *ced-10*** doubles also display a failure in the migrations of 2° vulval cells toward the 1° vulval cells to form a functional vulva (Kishore and Sundaram, 2002). **CED-10** might act downstream of the Semaphorin guidance cue **SMP-1** and its receptor **PLX-1/Plexin** in parallel to **UNC-73/Trio** and **MIG-2** (Dalpe et al., 2005).

In the male tail, **CED-10** and **MIG-2** are involved in the positioning of the ray 1 sensillum relative to other ray sensilla. The guidance cues **SMP-1/Semaphorin**, **UNC-6/Netrin**, and **PVF-1/VEGF** interact in ray 1 placement, and **PVF-1** inhibits **CED-10** but not **MIG-2** activity during ray 1 placement (Dalpe et al., 2012; Dalpe et al., 2013). The ray 1 response to **SMP-1/PLX-1** signaling is governed by level of **MIG-2** and **CED-10** activity, whereby the attractive response at normal **CED-10** and **MIG-2** activity is converted to repulsion by reduced activity (Dalpe et al., 2004).

3.3.4. Cytokinesis

RhoA activity in formation and function of the actin contractile ring during cytokinesis is governed by the central spindle and the centalspindin complex of proteins. The Rho GAP **CYK-4** is a component of centalspindin, and one of its roles is to inhibit **CED-10/Rac** activity during cytokinesis (Canman et al., 2008). This model is consistent with a general theme of opposing roles of Rho and Rac in the cell, with Rho controlling cell contraction and retraction, and Rac controlling cell protrusion and migration (Burridge and Doughman, 2006).

3.3.5. Endosomal recycling and apical-basolateral polarity

Recycling endosomes are critical for maintenance of apico-basolateral polarity in intestinal epithelial cells. In the absence of the 14-3-3 scaffolding protein **PAR-5**, apical recycling endosomes ectopically accumulate at basolateral positions dependent upon **CED-10** and **RHO-1** (Winter et al., 2012). The **CED-5/CED-12/CED-10** regulatory module also promotes progression of basal recycling endosomes to the plasma membrane by downregulating the **RAB-5** GTPase (see Section 4.4) and allowing exit from the early endosome (Sun et al., 2012).

3.3.6. Synaptic vesicle clustering

The conserved **CED-5/CED-12/CED-10** complex is required for the clustering of synaptic vesicles at active zones in response to the guidance cue **UNC-6/Netrin** and its receptor **UNC-40/DCC** (Stavoe and Colon-Ramos, 2012). Active zone positioning itself was not affected by **CED-10**. The functional consequence of aberrant synaptic vesicle clustering was revealed by pharmacological studies in ***ced-10*** mutants (sensitivity to the GABA(A) receptor antagonist pentylenetetrazole, and to the acetylcholinesterase inhibitor aldicarb) (Locke et al., 2009).

3.3.7. Wnt signaling

CED-10/Rac participates in Wnt-dependent developmental events that regulate polarity and the cytoskeleton (Cabello et al., 2010). The **CED-5/CED-12/CED-10** regulatory module acts downstream of the Wnt receptor **MOM-5/Frizzled**, **GSK-3** kinase, and **APR-1/APC** in cell corpse engulfment and DTC migration (Cabello et al., 2010). In orientation of mitotic spindle polarity in the early embryo, **MOM-5**, **GSK-3**, and **WRM-1/β-catenin** regulate **CED-10/Rac** independently of downstream canonical transcriptional components **LIT-1/NLK** and **POP-1/TCF** (Cabello et al., 2010).

3.3.8. Axon pathfinding

ced-10 and *mig-2* act redundantly in the pathfinding of axons (Lundquist et al., 2001; Wu et al., 2002). Single mutants display weak defects, which become synergistically stronger in the double mutant. Axon pathfinding in the dorsal, ventral, anterior, and posterior directions are affected, suggesting that **CED-10** and **MIG-2** act with multiple guidance cues. The double mutant displayed defects in axon outgrowth, axon guidance, and axon branching, indicating that Racs control multiple aspects of axon pathfinding. Furthermore, *Rac* double mutants displayed ectopic axon branches, suggesting that Racs also control the pruning of spurious axon branches (Struckhoff and Lundquist, 2003). Constitutively active **CED-10** and **MIG-2** perturb axon pathfinding (Zipkin et al., 1997; Struckhoff and Lundquist, 2003; Shakir et al., 2006; Shakir et al., 2008). *Rac(G12V)* molecules induce the formation of ectopic lamellipodia and filopodia on neurons that resemble those found on the growth cone during axon outgrowth (Knobel et al., 1999).

Genetic and epistasis analysis indicated that the GEFs **UNC-73/Trio** (Lundquist et al., 2001; Wu et al., 2002; Struckhoff and Lundquist, 2003) and **TIAM-1** (Demarco et al., 2012) act with **CED-10** and **MIG-2** in axon pathfinding (Figure 14). **CED-5/CDM** also acts with **CED-10** and **MIG-2** (Lundquist et al., 2001), suggesting that the conserved **CED-5/CED-12** GEF is involved. Downstream, the Arp2/3 activators **WVE-1/WAVE** and **WSP-1/WASP** act in the **CED-10** and **MIG-2** pathways, respectively. WAVE molecules are known to act with Racs. WASP molecules were shown to act with **CDC-42**, and this work illustrates an interaction of **WSP-1** in the **MIG-2** pathway. This interaction was confirmed in polarization and intercalation of embryonic epithelial cells, where **WSP-1** also acts with **MIG-2** (Walck-Shannon et al., 2015). The WASP and WAVE regulators **GEX-2/Sra-1** and **GEX-3/Hem2** act in both pathways (Shakir et al., 2008). The actin-binding protein **UNC-115/abLIM** also acts downstream of **CED-10** and **MIG-2** in axon pathfinding (Struckhoff and Lundquist, 2003). Thus, **CED-10** and **MIG-2** might utilize at least two distinct pathways to modulate the growth cone actin cytoskeleton: the Arp2/3 complex and **UNC-115/abLIM**. Indeed, lamellipodial and filopodial protrusions on growth cones are dependent upon Arp2/3 and **UNC-115/abLIM** (Norris et al., 2009) (Figure 14). The 7WD-repeat receptor of activated C kinase **RACK-1** might mediate **CED-10** and **MIG-2** interaction with **UNC-115/abLIM** (Demarco and Lundquist, 2010), and the conserved 7WD-repeat protein **SWAN-1** might inhibit **CED-10** and **MIG-2** activity in axon pathfinding (Yang et al., 2006). In addition to initial axon guidance, **CED-10** also mediates axon regrowth after severing (Gabel et al., 2008).

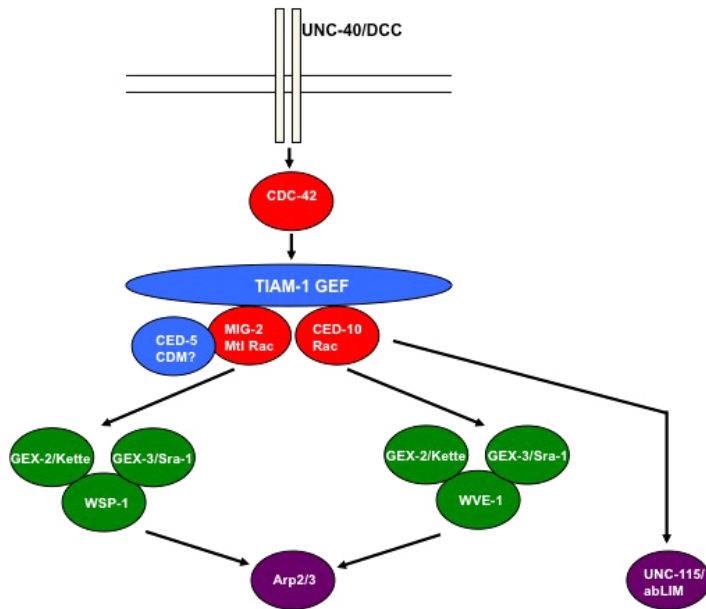


Figure 14. Rho GTPases in attractive axon guidance and neuronal protrusion. **CED-10** and **MIG-2** act redundantly downstream of **CDC-42** and the guidance receptor **UNC-40/DCC** in attractive axon guidance and neuronal protrusion. The Arp2/3 activator complex acts in both pathways, with **WSP-1/WASP** acting in the **MIG-2** pathway and **WVE-1/WAVE** acting in the **CED-10** pathway. The GEF **TIAM-1** regulates **CED-10** and **MIG-2** in attractive axon guidance and neuronal protrusion.

Growth cones detect guidance information using transmembrane guidance receptors. Indeed, **CED-10** and **MIG-2** and the **UNC-115** actin-binding protein act downstream of the netrin guidance receptor **UNC-40 DCC** in

ventral axon pathfinding (Gitai et al., 2003; Demarco et al., 2012). Furthermore, **CED-10**, **MIG-2**, and **RAC-2** act with the **MIG-15** NIK kinase downstream of the integrin complex in VD/DD motor axon circumferential navigation (Poinat et al., 2002). Asymmetric activation of **CED-10** by **UNC-6**/Netrin signaling results in the asymmetric accumulation of the **MIG-10**/Lamellipodin that defines axon initiation in attractive guidance (Quinn and Wadsworth, 2008). Furthermore, **MIG-2** is inhibited by an **UNC-6**-independent role of **UNC-40/DCC** mediated by the SYD-1C RhoGAP-like protein (Xu et al., 2015).

3.3.9. Growth cone protrusion

Genetic analyses described above implicate **MIG-2** and **CED-10** in regulation of axon guidance, and downstream cytoskeletal molecules in growth cone protrusive behavior during outgrowth has revealed a role of **CED-10** and **MIG-2** in regulating protrusion downstream of guidance receptors. The **UNC-6**/Netrin molecule regulates dorsal-ventral guidance by two distinct receptor complexes: **UNC-40/DCC** homodimers mediate attractive ventral directed growth, and **UNC-5/UNC-40** heterodimers mediate repulsive dorsal growth (Figure 15). Expression of a constitutively activated **UNC-40** cytoplasmic domain (MYR::UNC-40) in attracted or repelled neurons had distinct effects. In neurons with axons attracted to **UNC-6**/Netrin, MYR::UNC-40 led to excessive growth cone protrusion involving lamellipodial and filopodial structures (Gitai et al., 2003; Norris and Lundquist, 2011; Demarco et al., 2012) whereas in repelled growth cones, MYR::UNC-40 inhibited growth cone protrusions (Norris and Lundquist, 2011). These data indicate that the **UNC-6**/Netrin receptors differentially regulate growth cone protrusion, with the attractive **UNC-40/UNC-40** receptor stimulating protrusion, and the repulsive **UNC-5/UNC-40** receptor inhibiting protrusion, possibly in a gradient across the growth cone resulting in directed outgrowth.

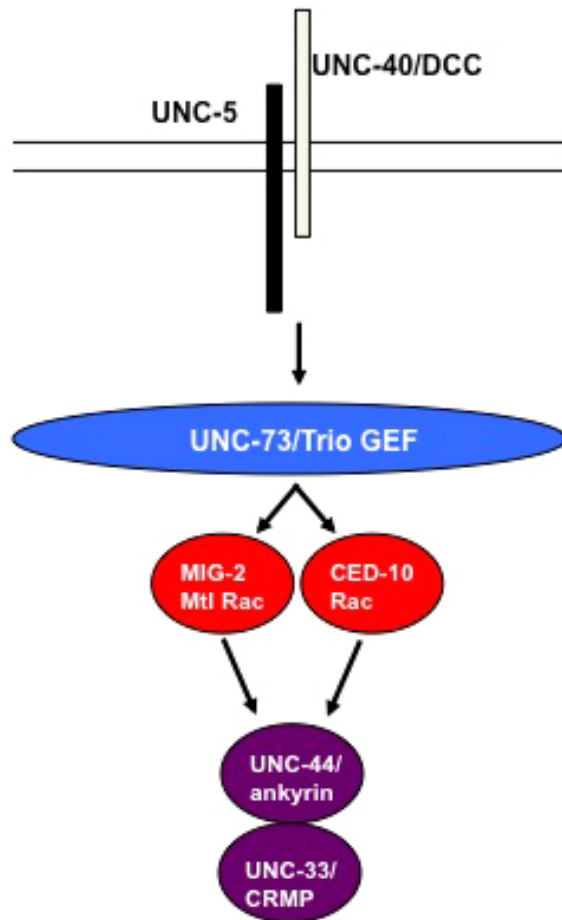


Figure 15. Rac GTPases in repulsive axon guidance and inhibition of growth cone protrusion. Downstream of the repulsive guidance receptor heterodimer of **UNC-40/DCC** and **UNC-5**, **CED-10** and **MIG-2** act redundantly to inhibit growth cone protrusion in repulsive axon guidance via the microtubule-interacting protein **UNC-33/CRMP** and **UNC-44/CRMP**. The **UNC-73/Trio GEF** regulates **CED-10** and **MIG-2** in inhibition of protrusion.

CED-10 and **MIG-2** were required for ectopic protrusion caused by **MYR::UNC-40** in neurons with axons attracted to **UNC-6/Netrin** (Gitai et al., 2003; Demarco et al., 2012). The GEF **UNC-73/Trio** was not required, but the GEF **TIAM-1** was, indicating that **TIAM-1** but not **UNC-73** regulate **MIG-2** and **CED-10** in protrusion. These data along with those discussed above suggest that **UNC-40/DCC** stimulates growth cone protrusion via **TIAM-1**, **CED-10/MIG-2**, and the downstream cytoskeletal effectors **UNC-115/abLIM** and **Arp2/3**.

In growth cones repelled from **UNC-6/Netrin**, **MYR::UNC-40** inhibited protrusion, resulting in small growth cones with little or no filopodial protrusions (Norris and Lundquist, 2011; Norris et al., 2014). Consistent with this, loss of repellent **UNC-5** receptor resulted in larger-than-normal growth cones with more filopodial protrusion. Using epistasis analysis, a pathway required for growth cone inhibition downstream of **UNC-40/UNC-5** repulsive receptor was described, including **UNC-73/Trio**, **MIG-2** and **CED-10**, and the cytoskeletal regulators **UNC-44/Ankyrin** and **UNC-33/CRMP** (collapsin response mediating protein) (Norris et al., 2014) (Figure 15). **UNC-33/CRMP** is a putative microtubule-interacting protein whose localization is dependent upon **UNC-44/Ankyrin** (Maniar et al., 2012). Activated **CED-10** or **MIG-2** expression in these neurons mimicked **MYR::UNC-40** and led to small growth cones with few protrusions (Figure 16). **UNC-44/Ankyrin** and **UNC-33/CRMP** were required, indicating that they are downstream effectors of **CED-10** and **MIG-2** (Figure 17). Thus, **CED-10** and **MIG-2** might inhibit growth cone protrusion downstream of **UNC-40/UNC-5** by regulating the microtubule cytoskeleton.

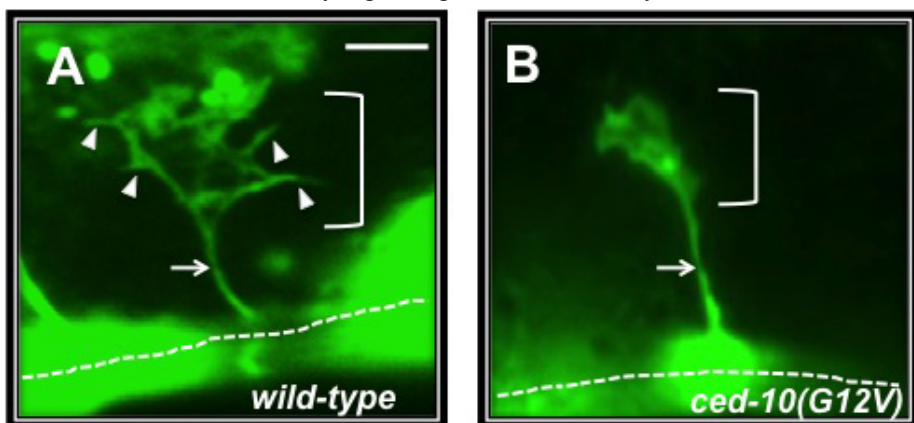


Figure 16. Activated CED-10(G12V) inhibits VD growth cone protrusion. Fluorescent micrographs of *Punc-25::gfp* expression in the growth cones of VD neurons in early L2 animals are shown. A) A *wild-type* VD growth cone displayed multiple robust filopodial protrusions. B) A VD growth cone expressing activated **CED-10(G12V)** displayed no robust filopodial protrusions and was reduced in size relative to the wild type. The scale bar in A represents 1 μ m. Dorsal is up, and anterior is to the right. The ventral nerve cord is indicated by a dashed line. Arrows indicate the VD axon, and the growth cone is bracketed. Arrowheads indicate filopodial protrusions. Micrographs: Maheka Gujar from the Lundquist lab.

3.3.10. Modularity of CED-10 and MIG-2 signaling

In *C. elegans*, 22 RhoGEFs of the DH-PH family regulate six Rho-family GTPases. Studies of **CED-10** and **MIG-2** in growth cone protrusion indicate that they can both stimulate and inhibit protrusion (Gitai et al., 2003; Norris and Lundquist, 2011; Demarco et al., 2012; Norris et al., 2014). The data suggest that different modules of **CED-10** and **MIG-2** associated with distinct GEFs and downstream cytoskeletal effectors mediate these opposing responses (e.g., **TIAM-1** and the **Arp2/3** complex in stimulation of protrusion, and **UNC-73/Trio** and **UNC-33/CRMP** in inhibition) (Figure 17).

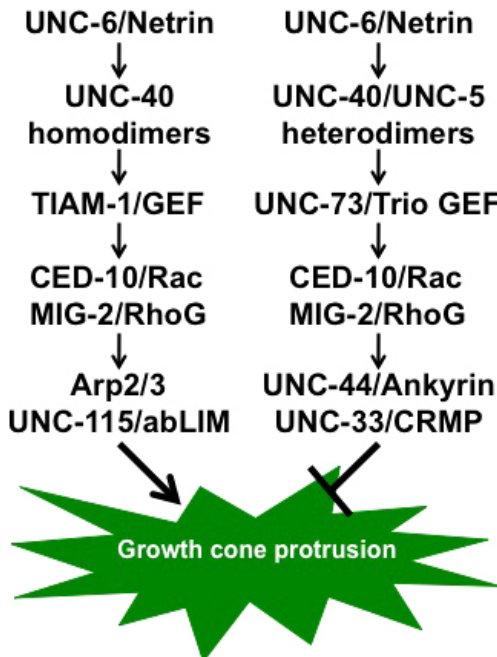


Figure 17. Modularity of Rac signaling in axon pathfinding. Rac GTPases MIG-2 and CED-10 control both stimulation and inhibition of neuronal protrusion in attractive versus repulsive UNC-6/Netrin-mediated axon guidance. The Rac act downstream of distinct receptor complexes (UNC-40 homodimers in attractive/stimulatory, and UNC-40/UNC-5 heterodimers in repulsive/inhibitory). Different GEFs regulate the Racs in each (TIAM-1 in attractive/stimulatory and UNC-73/Trio in repulsive/inhibitory). The stimulatory pathway influences the actin cytoskeleton via Arp2/3 and UNC-115/abLIM, and the inhibitory might modulate the microtubule cytoskeleton via UNC-33/CRMP and UNC-44/Ankyrin.

3.3.11. New, conserved Rho GTPase activating or dominant-negative mutations

Point mutations that alter GTPase activity or the interactions of Rho GTPases with GEFs, GAPs, and downstream effectors can lead to constitutive activation of the molecules or dominant inactivation of the pathways involving the molecules (e.g., the activating G12 and Q61 mutations found in human tumors, Rac and Ras numbering) (Fernandez-Medarde and Santos, 2011; Forbes et al., 2011). Genetic analysis of *C. elegans* axon guidance identified new activated and/or dominant-negative mutations in CED-10 and MIG-2 (Shakir et al., 2006). *ced-10(lq20)P29L*, *ced-10(n3246)G60R*, and *mig-2(lq13)S75F* do not behave as loss-of-function mutations and have dominant effects in transgenic expression assays not observed with the wild-type proteins (Shakir et al., 2006). Strikingly, the equivalent P29L mutation in *ced-10(lq20)* was recently identified in human *Rac1* as a driver of melanoma tumorigenesis (Hodis et al., 2012; Krauthammer et al., 2012), and G60R is synonymous with the *let-60(ay75)* temperature-sensitive gain-of-function mutation (Schutzman et al., 2001) (see Section 2.1). Biochemical analysis indicated that Rac1P29L was an activating mutation that enhanced interaction with the p21 binding domain (PBD) of PAK kinase. Further analysis will indicate if CED-10(G60R) and MIG-2(S75F) are activating or dominant negative, and if their mechanism is conserved in other organisms including humans.

3.4. Non-canonical Rho GTPases

3.4.1. CHW-1

chw-1 encodes an ortholog of mammalian proteins Chp/Wrch2/RhoV and Wrch-1/RhoU. Mammalian Wrch-1 was discovered as a gene whose expression is upregulated in response to oncogenic Wnt signaling. Wrch-1 is required for Wnt-dependent transformation and activated Wrch-1 transforms similarly to Wnt (Tao et al., 2001). This subfamily shares most homology with the Cdc42 subfamily. Like Cdc42, Wrch-1 can bind and activate Pak and Par6/aPKC and stimulate JNK (Tao et al., 2001; Chuang et al., 2007; Brady et al., 2009).

The CHW-1 protein structure departs from its Chp/Wrch subfamily members in three ways. First, the position 18 residue (12 by Ras and Rac numbering) is an Ala instead of the Gly that is nearly universal in Ras superfamily proteins. Second, CHW-1 contains a short 7 residue N-terminal extension that contains no prolines, compared to the 45 and 28 residue proline-rich Wrch-1/RhoU and Chp/Wrch-2/RhoV N-terminal extensions, respectively. These Proline-rich N-terminal extensions are thought to mediate auto-inhibitory functions that are relieved by binding of

SH3 domain-containing proteins like Grb2 (Shutes et al., 2004). The third difference that clearly distinguishes **CHW-1** from its subfamily relatives is the lack of a C-terminal membrane-targeting sequence. 3' RACE, RT-PCR, and subsequent RNAseq failed to identify missing 3' coding sequences that could contain the distinctive C-terminal CAAX sequence that mediates lipid modification (D.Reiner, unpublished results). Like many small GTPase C-termini, the **CHW-1** C-terminus is enriched with basic residues that could mediate an electrostatic switch, but whether **CHW-1** is membrane-targeted like its mammalian relatives is unclear. Notably, all of these **CHW-1** sequence features are highly conserved in **CHW-1** in related Caenorhabditids *C. briggsae* and *C. remanei*.

Animals mutant for the *chwg(ok697)* null deletion allele are superficially wild type. This allele also deletes the 5' end of the overlapping gene **F22E12.13**. A newly identified *chwg* exon 3 (of 4) nonsense allele from the Million Mutation Project (Thompson et al., 2013), *gk731043*, should disrupt *chwg* function without impacting **F22E12.13** function, but has not yet been analyzed. Ectopic expression of wild-type **CHW-1** in the distal tip cells of the somatic gonad weakly disrupted DTC migration (Kidd et al., 2015). Similar experiments with wild-type **CED-10/Rac** and **CDC-42** caused no defect on DTC migration, presumably because the activity of most small GTPases are tightly regulated by GEFs and GAPs. Ectopic DTC-specific expression of mutationally activated **CED-10/Rac** and **CDC-42** strongly disrupted DTC migration, consistent with affecting positional information during migration (Peters et al., 2013). Mutating **CHW-1** atypical A18 to G (Ras and Rac numbering A12G) abolished migration defects from ectopic expression, while the A18V (Ras and Rac numbering A12V) or Q69L (Ras and Rac numbering Q61L), both predicted to maximally activate **CHW-1**, increased activation over the wild-type **CHW-1** (Kidd et al., 2015). These results suggest that **CHW-1** is partially constitutively active, unlike most small GTPases. Genetic experiments suggest that **CHW-1** functions non-autonomously in the vulva to equalize contributions of **LIN-18/Ryk** and **LIN-17/Fz** receptors in **P7.p** polarity (Kidd et al., 2015). Thus, like its mammalian counterpart **Wrch-1**, **CHW-1** might act with Wnt signaling.

3.4.2. CRP-1

CRP-1 is most similar to **CDC-42** (Caruso et al., 2005; Jenna et al., 2005) and has no obvious mammalian counterpart based on sequence, although it remains to be seen if **CRP-1** has a functional counterpart in mammals, as does **MIG-2** (deBakker et al., 2004). Like other Rho GTPases, **CRP-1** has a C-terminal CAAX prenylation sequence but lacks the Rho insert domain found in other Rho GTPases (Jenna et al., 2005). The **CRP-1** effector-binding region is highly diverged, suggesting that **CRP-1** engages effectors other than canonical Cdc42 effectors. **CRP-1** also has an unconventional nucleotide-binding P-loop structure, with the highly conserved Gly 12 changed to a Thr in **CRP-1**. Indeed, **CRP-1** is largely deficient in GTP hydrolysis activity, and has a low affinity for GTP (Caruso et al., 2005; Jenna et al., 2005). Thus, while similar to Rho GTPases, GTP hydrolysis and interaction might not be central to **CRP-1** function. In fibroblasts, **CRP-1** protein localized to the trans-Golgi network and recycling endosome, and affected trafficking in epithelial cells in *C. elegans* (Jenna et al., 2005).

4. The Rab family

As in other systems, Rab GTPases are conserved regulators of multiple aspects of intracellular membrane trafficking and dynamics. The 31 *C. elegans* Rab-like genes are listed in Table 1 (Section 9) (Gallegos et al., 2012). While genetic analyses in *C. elegans* have delineated the functions of many of the Rab genes in diverse cellular and developmental events, the roles of others remain mysterious.

Descriptions of the Rab molecules in diverse cellular and developmental events are summarized below. This review is intended as a compendium of the effects of specific *rab* genes and not a comprehensive review of the processes affected by Rabs in *C. elegans*. Rab molecules in *C. elegans* with known roles affect membranous organellar formation, maturation, and trafficking, in particular the endocytic and late/recycling endosome (see Wormbook chapter *C. elegans* as a model for membrane traffic). Rabs are involved in clathrin-dependent and independent receptor-mediated endocytosis (RAB-5, 10, and 35), in intra-Golgi and endosome to Golgi retrograde transport (RAB-6.1), recycling from endosomes to the plasma membrane (RAB-10 and RAB-11.1), and maturation and function of the autophagosome in autophagy and the phagosome in cell corpse degradation (RAB-2, 7, and 14) (see Wormbook chapters Autophagy in *C. elegans* and Programmed cell death). Rabs are also key regulators of lysosomal biogenesis (RAB-6.1 and GLO-1). In the nervous system, Rabs are involved in synaptic vesicle trafficking (RAB-1, 3, 5, 27, and GLO-1), dense core vesicle secretion (RAB-2, 10, and 27), and post-synaptic neurotransmitter receptor recycling via the endocytic pathway (RAB-6.2 and 10). Some other diverse roles of Rabs involve innate immunity (RAB-1), ethanol sensitivity (RAB-3), cortical granule secretion (RAB-6.1), and sensory cilia development (RAB-8).

That different Rabs have distinct effects suggest that some specificity in function resides in the Rabs themselves and not their regulators (as is the case with Rho GTPases). There is also likely much redundancy of function among the Rabs that have yet to be described, such as **RAB-3** and **RAB-27** in synaptic vesicle trafficking (Iwasaki et al., 1997; Iwasaki and Toyonaga, 2000; Mahoney et al., 2006). Additionally Rabs act sequentially, such as **RAB-2** and **RAB-14** acting together to regulating phagosome maturation and upstream of **RAB-7**, which mediates fusion with the lysosome (Guo et al., 2010; Guo and Wang, 2010). Rabs also act sequentially with other Rho-family GTPases, such as downstream of **CED-10/Rac** in endocytic recycling (Sun et al., 2012), and upstream of **ARF-6** in endosomal trafficking (Nilsson et al., 2008; Shi and Grant, 2013). These themes of redundancy and sequential activity are described in the vignettes below.

4.1. **RAB-1**

RAB-1 controls trafficking of the GPCR odorant receptor **ODR-10** into sensory cilia (Chen et al., 2014) and **CAV-1/caveolin** trafficking in oocytes (Sato et al., 2006). **RAB-1** is involved in innate immunity, which consists of the expression and secretion of anti-microbial peptides, the neuropeptide-like proteins (NLPs), and the caenacins (CNCs) (Couillault et al., 2004). **RAB-1** acts with **TIR-1**, a Toll/IL/IR-domain protein similar to vertebrate SARM, in this process (Couillault et al., 2004).

4.2. **RAB-2/UNC-108**

Mutations in *rab-2* were first identified by uncoordinated locomotion, and the locus was called *unc-108*. Here we will refer to it as **rab-2**. **RAB-2** controls a variety of vesicle trafficking and maturation events in neurons and other cells. In neurons, **RAB-2** and the RabGAPs **TBC-2** and **TBC-8** regulate dense core vesicle maturation and sorting (Sumakovic et al., 2009; Hu et al., 2011; Hannemann et al., 2012; Ailion et al., 2014). **RAB-2** regulates post-synaptic endocytosis of the **GLR-1** receptor to the recycling endosome, but not the multivesicular body degradation pathway (Chun et al., 2008), and trafficking of the GPCR odorant receptor **ODR-10** to sensory cilia (Chen et al., 2014). **RAB-2** also controls the maturation and function of the phagosome involved in the degradation of apoptotic cell corpses after phagocytic engulfment (Lu et al., 2008; Mangahas et al., 2008; Guo et al., 2010; Guo and Wang, 2010; Lu et al., 2012).

4.3. **RAB-3**

RAB-3 localizes to synaptic vesicles and controls their trafficking to the presynaptic active zone (Nonet et al., 1997; Mahoney et al., 2006; Gracheva et al., 2008). **RAB-27** acts with **RAB-3**, and both are regulated by the Rab GEF **AEX-3** in this process (Iwasaki et al., 1997; Iwasaki and Toyonaga, 2000; Mahoney et al., 2006). **UNC-10/Rim** might act with **RAB-3** to control vesicle fusion (Koushika et al., 2001; Gracheva et al., 2006) and with **RAB-3/AEX-3** to mediate **UNC-57/Endophilin** association with SVs to promote fusion (Bai et al., 2010).

RAB-3 loss results in sensitivity to ethanol and other inhalants (Kapfhamer et al., 2008; Davies et al., 2012), and this effect appears to be independent of **RAB-3**'s role in synaptic vesicle secretion (Johnson et al., 2013). **RAB-3** might affect other neurosecretory events such as dense core vesicle secretion, or have some other as-yet unknown role in ethanol sensitivity.

4.4. **RAB-5**

RAB-5 regulates the early to late endosome maturation process along with **RAB-7** (Chotard et al., 2010; Poteryaev et al., 2010). In the nervous system, this involves the earliest stages of synaptic vesicle formation (Sann et al., 2012). The GAP **TBC-2** negatively regulates **RAB-5** and is recruited to the early endosome as it matures, removing **RAB-5** to allow later-acting Rabs, such as **RAB-7** (degradation) and **RAB-10** (recycling), to function. **RAB-10** directly recruits **TBC-2**, with help from **CED-10** and **AMPH-1** (Chotard et al., 2010; Sasidharan et al., 2012; Sun et al., 2012; Liu and Grant, 2015). In addition, **RAB-5** is involved in endocytosis of yolk (Grant and Hirsh, 1999; Balklava et al., 2007), and the GEF **RME-6** mediates association with clathrin-coated pits (Sato et al., 2005).

RAB-5 is involved in a number of other trafficking and membrane-associated events. In neurons it is involved in dense core vesicle release along with **RAB-10** (Sasidharan et al., 2012) and sensory ciliary transport (Kaplan et al., 2012). **RAB-5** is also required for phagosomal maturation involved in degradation of engulfed cell corpses (Li et al., 2009; Kinchen and Ravichandran, 2010; Li et al., 2012). In the early embryo, **RAB-5** regulates the association of

the polarity protein **PAR-6** with the cell cortex and affects actin organization (Hyenne et al., 2012). Along with RAB-11, **RAB-5** is involved in plasma membrane repair after damage by bacterial pore-forming toxins (Los et al., 2011). **RAB-5** also has a role in maintaining the structure of the endoplasmic reticulum, which seems to be independent of its role in endosomal trafficking (Audhya et al., 2007). The **RAB-5** GEF, **RABX-5**, which contains a zinc finger and Vps9 domain, acts with **RAB-5** and the **RAB-5** effector **RABN-5** in multiple events including endosomal trafficking and synaptic vesicle release (Poteryaev et al., 2010; Sann et al., 2012).

4.5. **RAB-6.1**

In the oocyte after fertilization, **RAB-6.1** is required for fusion of the cortical granules to the oocyte plasma membrane by recruiting the separase fusion protein to the cortical granules (Kimura and Kimura, 2012). **RAB-6.1** associates with **VPS-52**, a component of the Golgi-associated GARP complex that functions in retrograde transport within the Golgi and in Golgi-to-endosome transport (Luo et al., 2011). *rab-6.1; rab-6.2* double knockdown treatment results in sterility, suggesting that these molecules act redundantly in the germline (Audhya et al., 2007).

4.6. **RAB-6.2**

In neurons, **RAB-6.2** is involved in the retrograde recycling of the **GLR-1** AMPA-type glutamate receptor along with the retromer complex (Zhang et al., 2012). The grinder is an extracellular structure in the pharynx required to crush bacteria as they are eaten. **RAB-6.2** and the GAP **EAT-17** are required for secretion and formation of the pharyngeal grinder (Straud et al., 2013).

4.7. **RAB-7**

RAB-7 controls late endosomal maturation along with **RAB-5** (Poulin and Ahringer, 2005; Poteryaev et al., 2007; Sato et al., 2008; Chotard et al., 2010). In autophagy, **RAB-7** controls the maturation of the autophagosome (Jenzer et al., 2014; Manil-Segalen et al., 2014) and the phagosome in cell corpse degradation (Kinchen et al., 2008; Yu et al., 2008; Li et al., 2009; Guo et al., 2010; Guo and Wang, 2010; He et al., 2010; Lu et al., 2011) along with CED-10/Rac (Nieto et al., 2010). Other trafficking events controlled by **RAB-7** include formation of gut lysosomes (Delahaye et al., 2014), and late endosomal trafficking that antagonizes *let-23*/EGFR signaling in the vulva (Skorobogata and Rocheleau, 2012). **RAB-7** might also play a role in α -synuclein lethality in a transgenic model of α -synuclein overexpression (Kuwahara et al., 2008).

4.8. **RAB-8**

In ciliated sensory neurons, **RAB-8** is required for membrane delivery to ciliary and periciliary domains, balanced against clathrin and dynamin-mediated removal of material from these domains. (Mukhopadhyay et al., 2008; Kaplan et al., 2010; Kaplan et al., 2012). Maintenance of ciliary and periciliary domains is disrupted in Bardet-Biedl syndrome (Avidor-Reiss and Leroux, 2015).

4.9. **RAB-10**

RAB-10 is generally involved in endocytic recycling. It regulates endosomal phosphatidylinositol-4,5-bisphosphate (PI(4,5)P₂) levels, and via the Arf GAP **CNT-1**, downregulates the small GTPase **ARF-6**, allowing interaction of a series of other downstream effectors involved in vesicle budding (e.g., **RME-1** and **SPDN-1**) (Nilsson et al., 2008; Shi and Grant, 2013). It acts with exocyst in endocytic regulation (Jiu et al., 2012), as well as with actin-binding protein **EHBP-1** and **NUM-1/Numb** in endosomal transport in intestine (Nilsson et al., 2008; Shi et al., 2010). **NUM-1** acts between **RAB-10** and **RME-1** but is not a direct target of **RAB-10** as is **EHBP-1**. A clathrin-independent role of **RAB-10** is involved in the recycling of basolateral regions via a novel endosomal compartment in intestinal cells (Chen et al., 2006; Shi et al., 2010; Chen et al., 2014). In neurons, **RAB-10** regulates dense core vesicle secretion along with **RAB-5** (Sasidharan et al., 2012) and glutamate receptor recycling in post-synaptic regions along with **NUM-1/Numb** (Glodowski et al., 2007).

4.10. **RAB-11.1**

In epithelial cells **RAB-11.1** is positioned by the polarity molecule **PAR-5** to mediate apical-basolateral polarity (Winter et al., 2012). **RAB-11.1** vesicles localize to the cleavage furrow and mid-body during cell division downstream of separase and the receptor for activated C kinase **RACK-1** (Ai et al., 2009; Bembenek et al., 2010), necessary for proper spindle alignment (Zhang et al., 2008). After fertilization, **RAB-11.1** is required for secretion of chondroitin proteoglycans (Sato et al., 2008).

4.11. RAB-14

In engulfment of cell corpses, RAB-14 controls phagosome maturation along with RAB-7 and RAB-2/UNC-108 (Guo et al., 2010; Guo and Wang, 2010). RAB-14 and RAB-2/UNC-108 act redundantly in phagosome maturation, and RAB-7 mediates fusion with the lysosome in a sequential action of Rab GTPases in this process.

4.12. AEX-6

AEX-6 was identified as a mutant with defects in defecation, and was later found to regulate neuromuscular synaptic transmission underlying defecation and to correspond to RAB-27. AEX-6 is controlled by the AEX-3 GEF and the effector RBF-1/Rabphilin might act downstream to modulate SNARE function in synaptic vesicle release (Mahoney et al., 2006; Tanaka et al., 2008). AEX-6 also has a role in dense core vesicle release (Feng et al., 2012) and possibly insulin secretion involved in aging and Dauer larva formation (Ailion and Thomas, 2003).

4.13. GLO-1

GLO-1 Rab GTPase is required for lysosomal organization in the formation of gut granules—birefringent lysosome associated organelles—in the intestine (Hermann et al., 2005; Delahaye et al., 2014). Furthermore, GLO-1 is required downstream of RPM-1/PHR in synaptic organization and axon termination. The RCC GEF GLO-4 regulates GLO-1 in this process, which likely involves a late endosomal pathway that acts in parallel to FSN-1, which acts with RPM-1 in the E3 ubiquitin ligase pathway (Grill et al., 2007).

4.14. RAB-35

RAB-35 controls accumulation of yolk receptors at the cortex of oocytes, which are required for clathrin-based receptor-mediated endocytosis of yolk proteins (Sato et al., 2008). RME-4 is a DENN-domain containing Rab GEF that might link RAB-35 to clathrin-coated pits. This complex might act upstream of RAB-7 and synergistically with RAB-11.

4.15. RAB-39

RAB-39, along with its binding partner RASF-1/RASSF protect against oxidative stress (Takenaka et al., 2013). RASF-1 interacts specifically with the GTP-bound form of RAB-39, but the basis of resistance to oxidative stress mediated by this complex is unclear.

4.16. RSEF-1

RSEF-1 is a divergent Rab-like GTPase similar to RASEF, with a calcium-binding EF hand domain in addition to the GTPase domain. RSEF-1 is required for the development of the uterine seam, a syncytial structure that connects the uterus to the body wall of the animal (Ghosh and Sternberg, 2014). RSEF-1 affects nuclear migration and process outgrowth in the uterine seam.

5. The Arf/Sar family

The *C. elegans* genome encodes 14 proteins in the Arf family. The Arf (ADP-ribosylation factor) family gets its name from their ability to stimulate ADP-ribosylation of G α subunits of heterotrimeric G proteins in collaboration with cholera toxin; ADP-ribosylation is a form of post-translational modification of proteins and is also the mechanism of action of various bacterial toxins. Arfs are generally involved in membrane/vesicle trafficking and cytoskeletal dynamics. Notably, Arfs are generally lipid modified by a N-terminal myristoyl group rather than the C-terminal farnesyl, geranylgeranyl, or palmitoyl modification found in most Ras superfamily members. Like other members of the Ras superfamily, Arfs are tightly regulated by GEFs and GAPs and have characterized effectors in various systems. Based on sequence SAR1 is considered to be part of the Arf group. Arls are typically similar to Arfs, but their functions are broader (Donaldson and Jackson, 2011).

The Arf family is divided into three classes: the Arfs, the Arf-like proteins (Arls), and Sar1. *C. elegans* expresses a single Class I Arf, ARF-1.2 (human Arf1 and Arf3), a single Class II Arf, ARF-3 (human Arf4 and Arf5), and a single class III Arf (human Arf6). ARF-1.1 is an outlier with no clear membership in a particular class, and may be a *C. elegans*-specific Arf. ARL-1 is the Arl protein most closely related to the Arf subfamily (Skorobogata et al., 2014).

5.1. ARF-1.1

arf-1.1 encodes an Arf-like gene that does not fall into a specific class or align with a specific canonical Arf type. ARF-1.1 proteins can be found in Caenorhabditis but not more distant nematode relatives. ARF-1.1 has been shown to repress meiotic maturation (Govindan et al., 2006).

5.2. ARF-1.2

arf-1.2 encodes an ARF1/3 ortholog required for embryonic development and proper secretion in multiple tissues. ARF-1.2 is broadly expressed throughout development, particularly in embryos. ARF1.2 functions in endoplasmic reticulum dynamics, endocytic sorting of proteins and cleavage furrow formation (Grant and Hirsh, 1999; Skop et al., 2004; Poteryaev et al., 2005; Sato et al., 2006). ARF-1.2, ARF-3, and ARF-6 may collaborate to regulate asymmetric cell division (Teuliere et al., 2014), caspase-independent cell extrusion (Denning et al., 2012), and microRNA regulation (Parry et al., 2007).

During patterning of the vulva (described in Section 2.1 and Section 2.7), EGFR must be localized to the basolateral plasma membrane of VPCs to be activated by EGF. The LIN-2/-7/-10 complex mediates basolateral localization. In the absence of this complex, or upon deletion of the EGFR C-terminal PDZ domain recognition peptide recognized by the LIN-2/-7/-10, EGFR is mislocalized to the apical surface of VPCs, which abrogates proper signaling (Simske et al., 1996; Kaech et al., 1998; Whitfield et al., 1999).

AGEF-1 is an ArfGEF: genetic and homology data are consistent with AGEF-1 promoting exchange on ARF-1.2 and ARF-3. AGEF-1, ARF-1.2, and ARF-3 antagonize EGFR signaling, perhaps at the level of re-localizing EGFR to the apical membrane (Skorobogata et al., 2014). A hypomorphic mutation in *agef-1* suppresses a *lin-2* mutation, thus blocking the re-localization of LET-23/EGFR to the apical plasma membrane compartment. AGEF-1 also impacts secretion from other cell types and, unsurprising for a key trafficking protein, is an essential gene. Arfs are involved in recruiting clathrin-coated vesicles (Stamnes and Rothman, 1993; Traub et al., 1993; Robinson, 2004). The AP-1 clathrin adaptor complex plays a role in LET-23 localization (Lee et al., 1994; Shim et al., 2000). AGEF-1, ARF1.2, and ARF-3 may function together to localize LET-23/EGFR to the apical membrane (Skorobogata et al., 2014).

CNT-2/AGAP, an ArfGAP containing a G-protein-like domain, Ankyrin-like repeat, and a PH domain, regulates apoptotic fate and daughter size from the Q.p neuroblast asymmetric division (Singhvi et al., 2011). CNT-2 functions in endocytosis. ARF-1.2 (formerly ARF-1) functions in the same process, and *arf-1.2(RNAi)* suppressed the CNT-2/AGAP mutant phenotype, consistent with CNT-2/AGAP GAP activity directly inhibiting ARF1.2. The putative ArfGEFs, GRP-1/cytohesin, EFA-6, and BRIS-1, also regulate Q neuroblast asymmetric division (Teuliere et al., 2014).

5.3. ARF-3

ARF-3 expression specifically in touch receptor neurons is under transcriptional control of MEC-3 (Zhang et al., 2002), though ARF-3 is probably ubiquitously expressed. The joint interactions of multiple Arf family members suggests functional redundancy between them, or multiple layers of vesicle regulation requirements in many cell biological events. Also see joint role of ARF-3 with ARF-1.2 in inhibiting EGFR signaling (in Section 5.2).

5.4. ARF-6

ARF-6 is strongly expressed in coelomocytes and generally expressed elsewhere. ARF-6 co-localizes with MTM-6/myotubularin and they function together in an endocytic pathway and regulation of phosphatidylinositol regulation (Dang et al., 2004). The ArfGAP CNT-1/ACAP, in mammals an Arf6-specific GAP, is a binding partner of RAB-10. In the intestine, RAB-10 and CNT-1 co-localize with ARF-6 on recycling endosomes to control synthesis of phosphatidylinositol-4,5-bisphosphate [PI(4,5)P2] from phosphatidylinositol-4-phosphate [PI(4)P] by the type I phosphatidylinositol-4-phosphate 5 kinase, an ARF-6 effector. RAB-10 thus activates CNT-1/GAP to inhibit ARF-6 dependent trafficking of PI(4,5)P2 to the plasma membrane (Shi et al., 2012).

The ArfGAP CNT-1/ACAP is also cleaved by CED-3/Caspase during apoptosis, and the N-terminal fragment of CNT-1/ACAP blocks Akt activation by binding and sequestering phosphatidylinositol (3,4,5)-triphosphate at the plasma membrane, thus removing the Akt pro-survival activity independently of ARF-6 (Nakagawa et al., 2014).

Also see the joint role of **ARF-6** with **ARF-1.2** and **ARF-3** in inhibiting Q neuroblast asymmetrical division (in Section 5.2).

5.5. ARFRP1

Y54E10BR.2 encodes the *C. elegans* ortholog of mammalian Arf-related protein 1 (ARFRP1). In mammals ARFRP1 is thought to function in membrane trafficking, but the specifics are not well understood. Phenotypes for **Y54E10BR.2** mutants have not been described.

5.6. SAR-1

sar-1 encodes the SAR1 ortholog, which is present in all eukaryotes. SAR1 proteins have conserved functions in formation of COPI and COPII secretory vesicles. Not surprisingly, **SAR-1** has come up in several RNAi screens for diverse biological functions, but **SAR-1** has not yet been studied in detail. **SAR-1** RNAi depletion was found to impact endoplasmic reticulum stress (Caruso et al., 2008), consistent with its described cell biological role in other systems.

5.7. ARL-1

arl-1 encodes an Arl1 ortholog. *Drosophila* Arf27A, an ARL-1-like protein, is required for RNAi silencing, and this observation was validated in *C. elegans* by manipulating **ARL-1** (Saleh et al., 2006). **ARL-1** is faintly expressed in the cytoplasm of male tail hypodermal cells before male tail morphogenesis in the L4 stage. **ARL-1::GFP** becomes punctate during male tail whip retraction, and *arl-1(RNAi)* caused defective retraction of the tail whip (Nelson et al., 2011). Expression in male tail neurons persists in adulthood. Male tail **ARL-1::GFP** expression in neurons is under positive control of **DMD-3** and/or **MAB-3** transcription factors, but **ARL-1::GFP** is negatively regulated by **DMD-3** and **MAB-3** in hyp10. Initial whole-genome RNAi screens suggest that **arl-1** is an essential gene.

5.8. EVL-20

evl-20, also known as **arl-2**, encodes an ARL2 ortholog. Mutations in **evl-20** disrupt many morphogenetic events, most notably in the vulva, male tail, and gonad. Cytokinesis is also disrupted in the **evl-20** mutant (Antoshechkin and Han, 2002). The **evl-20** mutant phenotypes were rescued by transgenic expression of human ARL2 under the control of the **evl-20** promoter, but not ARL1 or ARL3-7.

Transcriptional and translational GFP fusions were expressed broadly during embryogenesis, particularly in migrating hypodermis, and in neurons throughout life. **EVL-20** was expressed strongly in the vulva during vulval morphogenesis in the L4 larva, and also expressed in uterus, spermatheca, distal tip cells, sheath cells of the gonad, and other tissues whose development was disrupted in the mutant. Putative **EVL-20** maternal product was detected in early embryonic blastomeres by antibody, and was localized to the cell cortex. Maternal **EVL-20** is also required for embryogenesis, suggesting that early events in the **evl-20** mutant are maternally rescued. Depletion of maternal **EVL-20** also causes catastrophic defects in the mitotic spindle, highly aneuploid cells, and defective cytokinesis (Antoshechkin and Han, 2002).

5.9. ARL-3

arl-3 encodes the *C. elegans* ARL3 ortholog. **ARL-3** and **ARL-13** coordinate intraflagellar transport (Li et al., 2010). **ARL-3** localizes to the ciliated endings of sensory neurons. The **arl-3** mutant is superficially wild type and does not impact sensory ending structure or intraflagellar transport markers. Cilia-specific expression of constitutively activated **ARL-3** (Q72L, Q61L in Ras numbering) disrupted sensory cilia function. **ARL-3** is thought to inhibit cilogenesis through **HDA-6**/HDAC6-mediated regulation of microtubule acetylation. The **hda-6** mutant rescues the **arl-13** sensory neuron defect to the same degree as the **arl-3** mutant, and confers a sensory ending defect when over-expressed in ciliated sensory neurons. There is no additivity between **hda-6** and **arl-3** mutants, suggesting that they function in a linear pathway (Li et al., 2010).

5.10. ARL-5

arl-5 encodes the ortholog of mammalian Arl5. Whole-genome RNAi screens determined that **ARL-5** is required for viability and fertility, but **ARL-5** has not been studied specifically.

5.11. ARL-6

arl-6 encodes an Arl6 ortholog that is a component of cilia and undergoes intraflagellar transport (Fan et al., 2004). Studies of *arl-6* in *C. elegans* represent a novel and powerful application of model organism genetics to human disease. Bardet-Biedl syndromes (BBSs) are multisystemic disorders of blindness, polydactyly, obesity, and cognitive impairment (Katsanis, 2004). Mutations at multiple loci can cause BBS, but the cellular and molecular mechanism of the disease is unclear. Interestingly, all BBS gene homologs in *C. elegans* are expressed in ciliated neurons (Li et al., 2004), indicating that the disease might affect ciliary function. This fact was used to identify an additional human gene responsible for BBS3 (Fan et al., 2004). *C. elegans* genes expressed in ciliated neurons contain a DAF-19 RFX binding X-box in their promoters (Ansley et al., 2003; Li et al., 2004). All *C. elegans* genes with X-boxes in their promoters were identified, which included *arl-6*, along with their counterparts in the human genome. Those human genes that localized to the genetically defined interval of BBS3 were sequenced in BBS3 patients. By this method, human *arl-6* was found to harbor mutations in multiple BBS3 families (Fan et al., 2004).

5.12. ARL-8

arl-8 encodes the *C. elegans* ortholog of mammalian Arl8. ARL-8 regulates axonal transport of presynaptic vesicle cargo (Klassen et al., 2010). Animals harboring *arl-8* mutations show excessive accumulation of synaptic vesicle precursors, suggesting that ARL-8 functions to repress precursor aggregation. Viable reduced-function mutations were isolated in this study. ARL-8 and a JNK MAP kinase cascade act in opposition during clustering of presynaptic components, with ARL-8 binding directly to the UNC-104/Kif1A kinesin motor protein (Wu et al., 2013).

Maternally provided ARL-8 is required for proper embryogenesis. Lethality was rescued by human Arl8 as well as wild-type and gain-of-function Q75L (Q61L by Ras numbering) in *C. elegans* ARL-8, but not by putative dominant-negative T34N (T17N by Ras numbering) ARL-8 (Nakae et al., 2010). In coelomocytes ARL-8 is localized primarily to lysosomes, and ARL-8 is involved in endosome-lysosome transport by promoting late endosome-lysosomal fusion. The failure of animals mutant for *arl-8* to fuse late endosomes to lysosomes is consistent with defects in transporting endocytosed macromolecules to the lysosome. Consistent with these observations, ARL-8 is involved in phagolysosome formation and is required for efficient removal of apoptotic cells in the germline (Sasaki et al., 2013). As with late endosomes in coelomocytes (Nakae et al., 2010), in *arl-8* mutant germline sheath cells phagosomes fail to fuse with lysosomes, consequently delaying degradation of apoptotic cells. ARL-8 physically binds to the VPS-41 component of the fusion complex.

ARL-8 also functions in endogenous RNAi pathways (Fischer et al., 2013). A non-null ARL-8 mutant has decreased 26G siRNAs in ERGO-1 and ALG-3/4 pathways, and thus may function in 26G siRNA biosynthesis in an unknown manner.

5.13. ARL-13

arl-13 encodes an Arl13 ortholog required for ciliogenesis. Human ARL13B is mutated in Joubert syndrome (Cantagrel et al., 2008), which is a disorder of cilia function. The Zebrafish scorpion (*sco*) mutant encodes ARL13b, which is required for cilia formation in the kidney; *sco* disruption causes cystic kidneys (Duldulao et al., 2009). ARL13b protein is highly enriched in cilia (Cevik et al., 2010).

Mutants for *arl-13* have shortened and malformed cilia. ARL-13 coordinates intraflagellar transport (Cevik et al., 2010; Li et al., 2010); in these mutant animals association between the IFT-A and IFT-B complexes is disrupted. RNAi-mediated depletion of *arl-3* (see Section 5.9) partially suppresses the *arl-13* mutant phenotype by re-establishing IFT-A and -B association, suggesting that ARL-3 and ARL-13 play opposing roles in IFT complex association. ARL-13 is expressed only in ciliated sensory neurons; early studies suggest that ARL-13::GFP is specifically located to the ciliated endings (Blacque et al., 2005; Li et al., 2010), but later analysis suggests specific localization to a small compartment (Brear et al., 2014). Double mutant analysis suggests that *hda-6* and *arl-3* are antagonistic (Warburton-Pitt et al., 2014); these differences may rely on assay. Cilia-specific expression of constitutively activated ARL-13 caused more severe disruption of sensory cilia function than did the *arl-13* loss-of-function mutant.

Mutants for *arl-13* also interacted additively with other sensory cilia mutants, such that ciliary neuron structure was worse in double mutants compared to single mutants suggesting the genes act in parallel (Cevik et al., 2010). Sequence analysis suggested that all ARL-13 homologs are palmitoylated, and ARL-13 and human Arl13b both undergo palmitoylation and are located to a proximal ciliary compartment.

Mutations in *arl-13* disrupted localization of the GFP-tagged G-protein coupled receptor STR-163 specifically in the ASK sensory neurons but not other neurons, and did not impact localization of other receptors, suggesting that **ARL-13** functions in a highly selective localization scheme (Brear et al., 2014). The *C. elegans* SUMO-conjugating enzyme **UBC-9** interacts with and SUMOylates the C-terminus of **ARL-13**. Mutant **ARL-13** that cannot be SUMOylated can still support ciliogenesis, but cannot localize certain sensory receptors to endings, and sensory functions mediated by those receptors are abrogated (Li et al., 2010). Constitutively SUMOylated **ARL-13** performs all **ARL-13** functions.

ARL-13 and mammalian **ARL13B** are localized to the inversinC-like subciliary compartment (Cevik et al., 2013). **ARL-13** palmitoylation and the **ARL-13** protein sequence motif, RVVP, prevent distal cilium and nuclear localization. **ARL-13** and **UNC-119** are partially redundant with **NPHP-2**/InversinC and **KLP-11**/kinesin for ciliogenesis (Warburton-Pitt et al., 2014), and together they precisely pattern microtubule structures in the cilia.

5.14. **ARC-1**

arc-1 encodes the ortholog of mammalian **ARD1/TRIM23**. **ARC-1** is a member of the Arf-like family of proteins, but it lacks the ability to promote ADP-ribosylation of substrates. **ARC-1**, and mammalian relatives, contain an N-terminal RING domain, two BBOX's, a BBC superfamily motif, and the C-terminal Arf GTPase domain, so the protein is definitely not a typical Arf family member. Mammalian **ARD1** has been shown to cooperate with **HIF-1** in hypoxic signaling (Mazure et al., 2004) and in NF- κ B signaling in anti-viral response (Poole et al., 2009; Arimoto et al., 2010). The function or expression pattern of **ARC-1** in *C. elegans* is unknown.

6. **Ran**

Eukaryotes generally encode a single Ran ortholog. The classic function of Ran is to regulate the cycle of nuclear import and export. A gradient of activated Ran is set up by GEFs and GAPs, with elevated Ran•GTP inside the nucleus and elevated Ran•GDP outside the nucleus (Clarke and Zhang, 2008; Cavazza and Vernos, 2015). Upon nuclear envelope breakdown accompanying cell division, a Ran•GTP gradient, driven by high RCC1/RanGEF proximity to chromosomes, promotes and maintains spindle assembly (Kalab et al., 2002; Kalab et al., 2006).

The *C. elegans* Ran ortholog is *ran-1*. Consistent with studies in other systems in which Ran controls import/export, nuclear membrane organization, and spindle organization, *ran-1(RNAi)* embryos display failure of nuclear membrane reassembly after mitosis (Askjaer et al., 2002; Bamba et al., 2002). Additionally, *ran-1(RNAi)* causes defects in mitotic spindle formation consistent with a defect in kinetochore association with microtubules rather than a defect in microtubule organization per se. RNAi of genes associated with Ran signaling cause similar defects, including the RanGAP/**RAN-2**, the Ran-binding protein **RanBP2/NPP-10**, the RanGEF **RCC1/RAN-3**, and **NTF2/RAN-4**. *ran-1(RNAi)* does not impact meiosis.

The Ran system maintenance of import/export, nuclear membrane structure, nuclear pores, and spindle assembly is important for many other processes that intersect with the nuclear membrane (Fernandez and Piano, 2006; Updike and Strome, 2009). Because of its broad impact, the Ran system arises repeatedly in screens for cell biological effects. Yet it is difficult to separate direct from indirect effects of such a fundamental molecular system.

7. The Rag family

Rag GTPases are not technically members of the Ras superfamily. While Rags are functionally GTPases, their structure diverges slightly from that of classic Ras superfamily members and they are not membrane-targeted through C-terminal prenylation. Their GEFs and GAPs are also atypical, consisting of the Ragulator complex (GEF) for RagA/B, the GATOR1 RagA/B GAP complex (Jewell et al., 2013; Bar-Peled and Sabatini, 2014), and the Folliculin RagC/D GAP complex (Petit et al., 2013; Tsun et al., 2013). Rags were first discovered in yeast, with *S. cerevisiae* Gtr1 being required for growth (Bun-Ya et al., 1992). We include the Rags because they have emerged as major regulators of TORC1 and anabolic metabolism. Where Rheb (and perhaps Ral) mediate global inputs of nutritional state in TORC1 and anabolism, via growth factor and insulin signals, Rags mediate local inputs into TORC1 at organelles. For example, Rag activity is high at the lysosome when levels of amino acids are elevated (Sancak et al., 2008).

Rag proteins come in two subfamilies: *C. elegans* **RAGA-1** corresponds to mammalian RagA and RagB. **RAGC-1** corresponds to mammalian RagC and RagD. Rag proteins heterodimerize. RagA/B•GTP binds with RagC/D•GDP to activate TORC1, while RagA/B•GDP binds with RagC/D•GTP when dissociated from TORC1 (Jewell et al., 2013; Bar-Peled and Sabatini, 2014).

7.1. RAGA-1

RAGA-1 is similar to mammalian RagA and RagB. Loss of **RAGA-1** prevents behavioral and neuromuscular decline associated with aging, perhaps through decreased activation of TORC1-dependent anabolism and attendant decreased lifespan and healthspan (Schreiber et al., 2010). Consistent with this model, *raga-1* loss ameliorates the life-shortening effect of high concentrations of food.

7.2. RAGC-1

Based on mammalian studies, **RAGC-1** is expected to be functionally equivalent to **RAGA-1**. However, **RAGC-1** has not been extensively analyzed.

8. Summary

In this chapter, we have described the roles of small GTPases in the many aspects of development and cell biology. As molecular switches their roles in specific events are determined by combinations of distinct GEFs and GAPs (e.g., in the case of the relatively few Rho-family GTPases), or proliferation of gene family members (e.g., in the case of the Rab-family). Questions remain about how GTPase activities are localized or restricted to particular subcellular compartments. For example, **CED-10/Rac** both stimulates and inhibits protrusion in growth cones, with each activity controlled by distinct GEFs. Future studies will unravel the combinations of regulators, GTPases, and their subcellular localizations that result in specific Rho-family GTPase activities.

9. Table 1. Ras superfamily GTPases in *C. elegans*.

C. elegans gene designation ¹	C. elegans gene name ²	Human/mouse orthologs ³
Ras family		
ZK792.6	let-60	H, N, KRas ⁴
C44C11.1	ras-1	R-Ras, R-Ras2/TC21
F17C8.4	ras-2	M-Ras/R-Ras3
C54A12.4	drn-1	Di-Ras1/Rig, Di-Ras2, Di-Ras3/Noey2/ARHI ⁵
C27B7.8	rap-1	Rap1A, B
C25D7.7	rap-2	Rap2A, B, C
C08F8.7	rap-3	Atypical Rap
Y53G8AR.3	ral-1	RalA, B
F54C8.5	rheb-1	Rheb, Rheb2/RhebL1
C14A11.7	ssr-2	Dexamethasone-induced Ras-related protein
Rho family		
Y51H4A.3	rho-1	RhoA, B, C
R07G3.1	cdc-42	Cdc42, TC10/RhoQ, TCL/RhoJ
C09G12.8	ced-10	Rac1, 2, 3
K03D3.10	rac-2	Atypical Rac ⁶
C35C5.4	mig-2	Invertebrate Mtl (MIG-2-like) family, functional similarity to mammalian RhoG
F22E12.2	chw-1	Chp/Wrch2/RhoV, Wrch1/RhoU
Y32F6B.3	crp-1	Atypical Cdc42

C. elegans gene designation ¹	C. elegans gene name ²	Human/mouse orthologs ³
Rab family		
<i>C39F7.4</i>	<i>rab-1</i>	Rab1
<i>F53F10.4</i>	<i>rab-2/unc-108</i>	Rab2
<i>C18A3.6</i>	<i>rab-3</i>	Rab3
<i>F26H9.6</i>	<i>rab-5</i>	Rab5
<i>F59B2.7</i>	<i>rab-6.1</i>	Rab6
<i>T25G12.4</i>	<i>rab-6.2</i>	Rab6
<i>W03C9.3</i>	<i>rab-7</i>	Rab7
<i>D1037.4</i>	<i>rab-8</i>	Rab8
<i>T23H2.5</i>	<i>rab-10</i>	Rab10
<i>F53G12.1</i>	<i>rab-11.1</i>	Rab11
<i>W04G5.2</i>	<i>rab-11.2</i>	Rab11
<i>K09A9.2</i>	<i>rab-14</i>	Rab14
<i>Y92C3B.3</i>	<i>rab-18</i>	Rab18
<i>Y62E10A.9</i>	<i>rab-19</i>	Rab19
<i>T01B7.3</i>	<i>rab-21</i>	Rab21
<i>Y87G2A.4</i>	<i>rab-27</i>	Rab27
<i>Y11D7A.4</i>	<i>rab-28</i>	Rab28
<i>Y45F3A.2</i>	<i>rab-30</i>	Rab30
<i>R07B1.12</i>	<i>glo-1</i>	Rab32/38
<i>F43D9.2</i>	<i>rab-33</i>	Rab33
<i>Y47D3A.25</i>	<i>rab-35</i>	Rab35
<i>W01H2.3</i>	<i>rab-37</i>	Rab37
<i>K02E10.1</i>		Rab37
<i>C56E6.2</i>		Rab6
<i>F11A5.3</i>		Rab2
<i>ZK669.5</i>		Rab23
<i>F11A5.4</i>		Rab2
Arf family		
<i>B0336.2</i>	<i>arf-1.2</i>	Arf1
<i>F57H12.1</i>	<i>arf-3</i>	Arf5
<i>Y116A8C.12</i>	<i>arf-6</i>	Arf6
<i>F45E4.1</i>	<i>arf-1.1</i>	Arf1
<i>Y54E10BR.2</i>		ARFRP1
<i>ZK180.4</i>	<i>sar-1</i>	Sar1A,B
<i>F54C9.10</i>	<i>arl-1</i>	Arl
<i>F22B5.1</i>	<i>evl-20</i>	Arf2
<i>F19H8.3</i>	<i>arl-3</i>	Arf3
<i>ZK632.8</i>	<i>arl-5</i>	Arf8
<i>C38D4.8</i>	<i>arl-6</i>	Arf6
<i>Y57G11C.13</i>	<i>arl-8</i>	Arl8A,B

C. elegans gene designation ¹	C. elegans gene name ²	Human/mouse orthologs ³
<i>Y37E3.5</i>	<i>arl-13</i>	Arf2
<i>ZK1320.6</i>	<i>arc-1/arl-4</i>	TRIM23
Ran family		
<i>K01G5.4</i>	<i>ran-1</i>	Ran

¹The *C. elegans* gene designation as noted on Wormbase.
²The three-letter *C. elegans* gene designation. Some genes have not yet been assigned a three-letter designation.
³Similarities are as noted in Wormbase.
⁴LET-60, by virtue of its polybasic hyper-variable C-terminus, is most similar to the KRas splice variant KRas4B.
⁵Assuming longer-than-annotated N-terminus and conserved GTPase domain based on alternative CUG initiator codons.
⁶Possibly a pseudogene, based on non-abundant RNaseq reads.

10. Acknowledgements

Thanks to N. Rasmussen for vulval images in Figure 6, M. Gujar for the growth cone images in Figure 16, K. Monahan and V. Muñiz-Medina for sharing unpublished results, J. Alan for helpful discussion, and all of the members of the Lundquist and Reiner labs who contributed to studies of small GTPases in development. Work in the Lundquist lab has been supported by NIH grants R01 NS040945 and R21 NS070417 to E.A.L., the Kansas Infrastructure Network for Biomedical Research Excellence (NIH P20 GM103418), and the Center for Molecular Analysis of Disease Pathways (NIH P20 GM103638). Work in the Reiner lab is supported by NIH grant R01 GM085309 to D.J.R.

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