Interrogating the ccm-3 Gene Network
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Interrogating the ccm-3 Gene Network

Graphical Abstract

Highlights

- Over 500 cerebral cavernous malformation (CCM) modifier genes discovered in C. elegans

- Bioinformatics methods predict 29 conserved genes in core CCM-3/CCM3 network

- Many genes exhibit ccm-3 phenotypes, including mop-25.2

- Loss of mop-25.2 homolog MO25 causes stress fiber formation in endothelial cells

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In Brief

Lant et al. use C. elegans genetics to better understand the human disease cerebral cavernous malformation (CCM). Through a whole-genome screen, and bioinformatics, they uncover a set of conserved genes that exhibit ccm-3 phenotypes. These targets may be able to inform therapeutic studies.

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SUMMARY

Cerebral cavernous malformations (CCMs) are neurovascular lesions caused by mutations in one of three genes (CCM1–3). Loss of CCM3 causes the poorest prognosis, and little is known about how it regulates vascular integrity. The C. elegans ccm-3 gene regulates the development of biological tubes that resemble mammalian vasculature, and in a genome-wide reverse genetic screen, we identified more than 500 possible CCM-3 pathway genes. With a phenolog-like approach, we generated a human CCM signaling network and identified 29 genes in common, of which 14 are required for excretory canal extension and membrane integrity, similar to ccm-3. Notably, depletion of the MO25 ortholog mop-25.2 causes severe defects in tube integrity by preventing CCM-3 localization to apical membranes. Furthermore, loss of MO25 phenocopies CCM3 ablation by causing stress fiber formation in endothelial cells. This work deepens our understanding of how CCM3 regulates vascular integrity and may help identify therapeutic targets for treating CCM3 patients.

INTRODUCTION

The nematode worm Caenorhabditis elegans contains several biological tubes, including multicellular tubes such as the intestine and germline as well as the unicellular excretory canals. Defects in the development or maintenance of these organs are often lethal to the organism or compromise its fitness. We recently developed C. elegans models of the neurovascular disease cerebral cavernous malformation (CCM) and discovered key roles for the worm CCM3 gene, ccm-3, in promoting the extension and maintenance of excretory canals (Lant et al., 2015) and germline development (Pal et al., 2017). The function of CCM-3 is dependent on its protein binding partners within the striatin interacting phosphatase and kinase (STRIPAK) complex, and loss of ccm-3 or STRIPAK genes causes defects in endocytic recycling and cytoskeletal organization (Lant et al., 2015; Pal et al., 2017). However, we lack a comprehensive understanding of how CCM-3/STRIPAK regulates these biological processes. Because of its genetic tractability and conservation of core CCM genes we took advantage of C. elegans to interrogate the CCM gene network.

CCM affects ~1 in 250 individuals and is caused by weak junctions in blood capillaries, which can leak blood into the brain parenchyma. This causes symptoms that range from mild headaches and seizures to hemorrhagic stroke, and it is not possible to predict when a lesion will bleed. The only treatment presently available for these patients is surgical resection, which can have devastating effects depending on lesion location. CCM can arise sporadically or, in approximately 40% of cases, by inheritance of mutations in one of the three CCM genes (CCM1/KRIT1, CCM2/OSM, and CCM3/PDCD10) (Fischer et al., 2013). These genes encode three distinct scaffold proteins that form a heterotrimeric complex, in which loss of any one can cause activation of the small GTPase RhoA and stimulation of Rho kinase (ROCK) (Borikova et al., 2010). This causes actin stress fiber formation, which compromises the integrity of cell-cell junctions. Although CCM1 is the most commonly mutated gene, mutations in CCM3 cause the earliest onset and most aggressive form of this disease (Shenkar et al., 2015).

A possible reason for the enhanced aggressiveness associated with CCM3 mutations may be that it functions in multiple complexes concurrently. Along with the hetero-trimeric CCM complex, CCM3 is also present in the STRIPAK complex (Kean et al., 2011). This complex contains a striatin backbone to which CCM3 binds and provides both phosphorylation and de-phosphorylation activity through germinal center kinase class III (GCKIII) kinases (MST3, MST4, and MST20) and protein phosphatase 2 (PP2A/C), respectively (Kean et al., 2011). Also bound to striatin is the MOB kinase activator (MOB3) and the striatin-interacting proteins (STRIP1/2), which themselves tether a number of proteins, including the sarcolemma-associated protein (SLMAP), IKK kinase suppressor (SIKE), and cortactin-binding protein (CTTNBP2) (Kean et al., 2011). STRIPAK has been implicated in a number of human diseases (Hwang and Pallas, 2014) and uses CCM3 as a kinase adaptor (Ceccarelli et al., 2011). CCM/GCKIII activity affects Golgi localization (Kean et al., 2011) and cytoskeletal organization by directing
Figure 1. Flowchart of Genome-wide RNAi Screen
Taking advantage of the synthetic lethal interaction previously identified between kri-1/CCM1 and ccm-3/CCM3 (Lant et al., 2015), 562 genes were identified that exhibit negative interactions (Emb. Let., embryonic lethality, sterility, or slow growth; Larv. Let., larval lethality; Syn. Let., synthetic lethality) when ablated by RNAi in kri-1 mutants. In parallel, a phenolog-like approach using GeneMANIA and STRING was used to generate a “CCM-ome” from genes known to function in human CCM signaling pathways. The 29 overlapping genes between these two sets (“tier 1 hits”) were assessed for ccm-3 phenotypes in C. elegans, such as excretory canal truncations. See also Figure S1.

The phosphorylation of a number of substrates, such as the actin-binding protein moesin (Zheng et al., 2010). Furthermore, emerging evidence indicates that CCM3 is able to act independently of the other two CCM proteins (Lant et al., 2015; Yoruk et al., 2012; You et al., 2017; Zhou et al., 2016a).

Although there have been intense efforts to understand the mechanisms by which CCM1 and CCM2 regulate vascular integrity, much less is known about the role of CCM3. Our aim in this study was to interrogate the C. elegans ccm-3 network by first conducting a whole-genome RNAi screen to identify a comprehensive set of genes that exhibit similar synthetic lethal interactions with kri-1/CCM1 as ccm-3 (Figure 1). Large-scale reverse genetics screens in C. elegans have many advantages for understanding poorly characterized genes, which often provide important insights into the biological functions of genes implicated in human diseases (Meier et al., 2014; Silverman et al., 2009; Sin et al., 2014). We uncovered more than 500 genes that exhibited strong negative genetic interactions with kri-1 and function in a range of cellular processes, including vesicle trafficking and cytoskeletal dynamics. By using a phenolog-like approach (McGary et al., 2010), we cross-referenced these C. elegans genes with a network assembled from genes previously shown to function in human CCM signaling. The resultant 29 genes (tier 1 hits) were systematically knocked down by RNAi and evaluated for ccm-3-specific phenotypes, such as excretory canal truncations. Of the 29 tier 1 genes, 14 caused strong excretory canal truncations and membrane defects when knocked down in wild-type animals. The most severe canal defects were caused by ablation of mop-25.2, a homolog of mouse embryo scaffolding protein 25 (MO25), which is involved in mediating AMPK signaling (Boudeau et al., 2003, 2004). Our analysis reveals a role for MOP-25.2, independent of its canonical binding partners, in the localization of both CCM-3 and GCK-1 to apical membranes necessary for biological tube integrity. Finally, we show that loss of MO25 phenocopies CCM3 ablation in endothelial cells by causing actin stress fiber formation. The combination of unbiased screening and bioinformatics analysis in C. elegans affords a powerful and efficient method for understanding how CCM proteins regulate biological tube development and may even uncover therapeutic targets.

RESULTS

Whole-Genome Screen to Identify Genes in the ccm-3 Pathway
We previously observed that, although loss of neither kri-1 nor ccm-3 on its own affects the survival of the worm, ablation of ccm-3 in kri-1 mutants results in synthetic lethality (Lant et al., 2015). We reasoned that other genes that cause similar negative genetic interactions with kri-1 may act in the ccm-3 pathway or cooperate with its in vivo functions. Therefore, we conducted a full-genome RNAi screen using the Source BioScience RNAi Library, which covers ~96% of annotated genes in the C. elegans genome, and identified 562 genes that reduced fitness of kri-1 mutants but not wild-type worms (Figure 1). These phenotypes, which include lethality at all stages of development, slow growth, and reduced fertility, all represent negative genetic interactions (Figure S1). Using the GeneMANIA application in Cytoscape, we constructed a network on the basis of these interactions to evaluate connectivity between our screen hits and genes known to be involved in CCM signaling (Figures S1 and S2, blue circles). Of the 562 genes identified, 237 had annotated interactions (genetic, physical, and/or predicted) with one another, while the remaining 325 “orphan” genes had only genetic interactions with kri-1 (Figure S1). For this study we focused on the 237 genes

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with documented interactions. Using Gene Ontology (GO) classification, these genes were found to be associated with a number of cellular processes, with a notable enrichment in cell localization and migration categories (including cell migration, cell motility, localization of the cell, and inductive cell migration) (Figure S2). Analysis of their functional roles indicate that a number fall into well-defined signaling cascades or cell process categories (Figure 2; Table S1). For example, we identified several cytoskeletal genes (i.e., actin, tubulin, interfilament organizer, and chaperones) and genes involved in small G protein signaling (i.e., small GTPases, GEF, and RHO-activating proteins). This is in line with previously described roles of CCM proteins in the regulation of actin polymerization and cell junction stability through Rho signaling (Borikova et al., 2010; Stockton et al., 2010; Whitehead et al., 2009). Consistent with observations in mammalian systems (Draheim et al., 2015; Zhou et al., 2015, 2016b) we also identified a number of MAPK signaling genes, such as a MEKK3 kinase, MAPK phosphatases, and a KLF transcription factor. Finally, several vesicle-trafficking proteins (including the exocyst complex, endosome-associated GTPase, and components of the adaptin complex) were identified, consistent with our recent observations on CCM-3/STRIPAK function in biological tube development (Lant et al., 2015; Pal et al., 2017).

To refine the primary network into a set of genes that are evolutionarily conserved, we used a “phenolog”-like approach which we refer to as “tier 1” hits (Figure 2, triangles, and Figure 3A; Table 1). We reasoned that genes identified in the RNAi screen that are orthologous to human genes in the CCM-ome would predict roles in CCM3 biology.

To investigate their in vivo functions, we used the C. elegans excretory canal as a model. This unicellular tube extends bi-directionally along both sides of the worm and functions as a renal system for the animal (Nelson and Riddle, 1984) (Figure 3B). Loss of ccm-3 causes significant truncations and “cyst” formation in excretory canals (Lant et al., 2015). We systematically ablated the 29 tier 1 genes by RNAi in wild-type animals and quantified excretory canal lengths. This was compared with a list of 29 randomly selected “control set” genes (Table S2). We found that the tier 1 set was significantly enriched (p = 6.9 × 10−36) for genes that caused canal truncations compared with the control set (Figures 3C, S3A, and S3B). Using the distribution of canal lengths under control conditions (Figure 3B), we denote “truncations” as canals that are <80% the length of those in wild-type animals. For a gene to cause a biologically relevant truncation when ablated, it must cause at least 5 times more canal truncations in the population of worms compared with control RNAi; a population of worms on control RNAi exhibits ~7% truncation (33 of 500), so for a gene to cause truncation, ≥35% of animals must have truncated canals. We defined a strong truncation to be similar to ccm-3 mutants (≥80% of animals with truncated canals). Analysis of individual genes within

**Figure 2. C. elegans CCM Network Map**
The network, with gene clusters showing enrichments in specific signaling pathways and cellular processes, was constructed using the GeneMANIA app in Cytoscape and gene annotations from Wormbase (www.wormbase.org). Genetic, physical, and predicted interactions are marked with green, pink, and orange lines, respectively. Negative hits from the primary screen are shown to have interaction with the CCM-related genes (blue spheres). Hits that will later be identified as “tier 1” are indicated by triangular outline. See also Tables S1 and S4 and Figure S2.
each set (Figures S3A and S3B; Tables 1 and S2) revealed that the control set had 26 genes with no truncation, 3 genes causing truncation, and no genes causing strong truncation. Conversely, the tier 1 set contained only 15 genes with no truncation, 11 genes causing truncations, and 3 genes causing strong truncations, which was a significant enrichment in genes required for canal extension compared with the control set (\( p = 0.0031 \)). Tier 1 genes causing canal truncations (Figure 3A, triangle- and diamond-outlined genes) represent a range of cellular processes, such as rab-5 (early endosome targeting GTPase), the Rho GEF ortholog ephx-1, the small GTPase cdc-42, and sec-3/sec-15 (exocyst). We also identified a number of genes that have no previously documented roles in excretory canal morphology, including the chaperonin complex subunit cct-2, the RNA lariat debranching enzyme dbr-1, polya-denylate-binding protein pab-1, and Rad21/Rec8-like cohesion protein scc-1. In addition, two uncharacterized genes, the RBM (RNA binding motif-containing) ortholog H28G03.1 and nuclear hormone receptor nhr-69, also caused canal truncations when ablated.

**Genes with CCM1/kri-1 Phenotypes**

Because the methodology of both the RNAi screen and the subsequent bioinformatic filtering do not bias against potential CCM1/kri-1 interactors, we asked whether any of the tier 1 hits would also cause kri-1 phenotypes. Loss of kri-1 causes resistance to ionizing radiation (IR)-induced germline apoptosis in the pachytest region (Ito et al., 2010) (Figure S3C, yellow asterisks). Conversely, loss of ccm-3 causes oocyte growth defects that leads to their death in the distal end of the germ line (Figure S3D, red asterisk). We quantified germ cell corpses after ablating tier 1 genes by RNAi in wild-type worms exposed to 60 Gy of IR, using kri-1 and the anti-apoptotic Bcl-2 homolog ced-9 as controls. Although ablation of some genes (cct-2, cdc-25.1, cdc-42, erfa-1, T04G9.4, ubl-1, and unc-112) caused germline defects that made it impossible to quantify apoptotic corpses, five genes (Figure S3E; dbr-1, his-13, icd-1, lin-23, and nhr-23) caused suppression of apoptosis similar to kri-1 RNAi (\( n = 50, p < 0.05 \) from control RNAi, \( p > 0.05 \) from kri-1 RNAi). If we define suppression as less than or equal to the levels of physiological apoptosis, or two corpses per gonad arm (Gumienny et al., 1999), ablation of kri-1 by RNAi caused ~50% suppression of apoptosis (24 of 50 germline arms with two or fewer corpses). Only his-13 RNAi caused a greater suppression of apoptosis (29 of 50 germline arms) than kri-1.

**mop-25.2 Is Required for Excretory Canal Integrity**

Of the tier 1 hits (Figure S3A), knockdown of MO25 ortholog mop-25.2 caused the most severe canal truncations (50 of 50 canals with canals <80% wild-type length) that were even more severe than knockdown of ccm-3 or gck-1 (\( p < 0.05 \) in both cases) (Figures 4A and 4B). Consistent with loss of ccm-3 (Lant et al., 2015), we frequently observed distended lumen and distal canal tip cysts in animals depleted of mop-25.2 (Figure 4C, middle and bottom). Interestingly, RNAi to either gck-1 or mop-25.2 significantly exacerbated truncations in ccm-3 mutants (\( p = 1.089 \times 10^{-9} \) and \( p = 2.49 \times 10^{-5} \), respectively) (Figure 4D). We also observed this with gck-1 mutant worms, in which depletion of ccm-3 or mop-25.2 significantly enhanced canal truncations (\( p = 3.30 \times 10^{-9} \), and \( p = 8.68 \times 10^{-9} \), respectively) (Figure 4D). Previously, we found that loss of ccm-3 and STRIPAK genes affected the stability of Golgi as well as subcellular localization of CDC-42 and the endocytic recycling protein RAB-11 in the canal. Knockdown of mop-25.2 also diminished puncta for these...
markers (Figure 4E; Table S3), indicating a similar role as CCM-3/STRIPAK. Collectively, these results reveal a role for MOP-25.2 in canal extension by both CCM-3/GCK-1-dependent and CCM-3/GCK-1-independent mechanisms.

Because there are three nematode MO25 paralogs in C. elegans, in which mop-25.1 and mop-25.3 are more distantly related to mop-25.2 (Chien et al., 2013), we wondered if they shared similar roles in canal development. Ablation of mop-25.1 and mop-25.3 caused weak canal truncations (Figure S4) and diminished puncta of CDC-42/recycling markers (Table S3), which concurs with previous studies showing some overlapping functions (Chien et al., 2013; Denning et al., 2012). Because MO25 regulates AMPK signaling through the MO25/STRAD/LKB1 complex (Boudeau et al., 2004; Hawley et al., 2003), we wondered whether MOP-25.2 regulates canal extension through the AMPK homolog. Therefore, we ablated both homologs of the catalytic alpha subunit of AMPK (aak-1/aak-2), the upstream LKB1 homolog (par-4), the ortholog of STRAD (strd-1), and the STRD-1 associated kinase (sad-1). Only knockdown of par-4 caused mild canal truncations (Figure S5), suggesting an alternate function for MOP-25.2 in canal extension that is distinct from its canonical binding partners.

**mop-25.2 Is Required for Germline Rachis Stability**

Loss of ccm-3 also causes sterility, whereby the germline lumen (rachis) collapses and impairs oocyte growth (Pal et al., 2017) (Figures 4F and 4G). Previously, mop-25.2 was shown to be required for embryonic development, whereby the mop-25.2(ok2073) deletion allele caused a maternal lethal effect (Chien et al., 2013), so we wondered if mop-25.2 shared any germline phenotypes with ccm-3. Using markers for both germline nuclei (mCherry::HIS-58) and membranes (GFP::PLCdelta1), knockdown of mop-25.2 caused rachis membrane collapse (Figure 4H, yellow asterisk) and small and rounded oocytes (Figure 4H, blue asterisk) in approximately 60% of animals (31 of 50). These germline defects were strikingly similar to those caused by loss of ccm-3, providing further support for MOP-25.2 playing a key role in the biological functions of CCM-3/STRIPAK.

### Table 1. Tier 1 Hits and Their Human Homologs

| Gene Name | WB Gene ID     | Truncation Rating | Human Gene Homolog | Human Gene ID     |
|-----------|----------------|-------------------|--------------------|-------------------|
| apl-1     | WBGene00000149 | no truncation     | APLP1              | ENSG00000105290   |
| cct-2     | WBGene00000378 | truncation        | CCT2               | ENSG00000166226   |
| cdc-25.1  | WBGene00000386 | truncation        | CDC25A             | ENSG00000164045   |
| cdc-42    | WBGene00000390 | strong truncation | CDC42              | ENSG00000070831   |
| dbr-1     | WBGene00000937 | no truncation     | DBR1               | ENSG00000138231   |
| ephx-1    | WBGene00019487 | truncation        | ARHGEF16           | ENSG00000130762   |
| erfa-1    | WBGene00020269 | no truncation     | ETF1               | ENSG00000120705   |
| glp-1     | WBGene0001609  | no truncation     | NOTCH2             | ENSG00000145741   |
| H28G03.1  | WBGene00019249 | truncation        | HNRNPA1            | ENSG00000135486   |
| his-13    | WBGene00001887 | no truncation     | HIST2H3D           | ENSG00000183598   |
| icd-1     | WBGene0002045  | truncation        | BTF3               | ENSG00000145741   |
| lin-23    | WBGene00003009 | no truncation     | FBXW11             | ENSG00000072803   |
| mop-25.2  | WBGene00013140 | strong truncation | CAB39              | ENSG00000135932   |
| nhr-23    | WBGene00003622 | no truncation     | RORA               | ENSG0000069667    |
| nhr-69    | WBGene00003659 | truncation        | HNF4A              | ENSG00000101076   |
| pab-1     | WBGene00003902 | no truncation     | PABPC4             | ENSG0000090621    |
| ppfr-2    | WBGene00017064 | no truncation     | PPP4R2             | ENSG00000163605   |
| rac-5     | WBGene00004268 | truncation        | RAB5B              | ENSG00000111540   |
| rc-1      | WBGene00009800 | no truncation     | ARHGAP30           | ENSG00000186517   |
| isc-1     | WBGene00004737 | truncation        | RAD21              | ENSG00000164754   |
| sdn-1     | WBGene00004749 | no truncation     | SDC1               | ENSG00000115884   |
| sec-15    | WBGene00016188 | truncation        | EXOC6              | ENSG00000138190   |
| sec-3     | WBGene00018703 | strong truncation | EXOC1              | ENSG00000090989   |
| sek-6     | WBGene00012162 | no truncation     | MAP2K4             | ENSG0000065559    |
| skr-21    | WBGene00004827 | no truncation     | SKP1               | ENSG00000113558   |
| T02G5.7   | WBGene00020166 | no truncation     | ACAT1              | ENSG0000075239    |
| T04G9.4   | WBGene00020215 | no truncation     | AASDHPPPT          | ENSG00000149313   |
| ubl-1     | WBGene00006725 | no truncation     | RPS27A             | ENSG00000143947   |
| unc-112   | WBGene00006836 | no truncation     | FERM1              | ENSG00000073712   |

Many C. elegans genes are the single homologs for multiple human genes or gene isoforms. Listed above are the human homologs on the basis of the BLASTP matches per Wormbase.org. For an expanded list of possible human homologs, see Table S4.
Previously, we showed that ablation of gck-1 in the germline caused CCM-3 to relocalize from the apical membrane to meiotic nuclei (Pal et al., 2017). Because mammalian MO25 has been shown to bind and stabilize GCKIII class kinases (Filippi et al., 2011), and a MO25-GCKIII-CCM3 ternary complex may exist (Yin et al., 2012), we asked if MOP-25.2 affects CCM-3 localization in a manner similar to GCK-1. CCM-3 localizes to apical membranes in both the canal and germline (Lant et al., 2015; Pal et al., 2017) (Figures 5 A and 5B, top). Using GFP translational reporters for CCM-3, we measured the fluorescence of GFP::CCM-3 in the excretory canal (along the lumen) and the pachytene region of the germline. Knockdown of ccm-3 caused the expected dissipation of canal GFP::CCM-3 (Figure 5A, second panels), as observed before, loss of gck-1 caused CCM-3 relocalization into pachytene nuclei (Figure 5B, third panels). However, loss of mop-25.2 significantly diminished GFP::CCM-3 signal on the lumen (Figure 5 B, fourth panel, and Figure 5D; p = 1.02 × 10^-13) but did not cause its relocalization to nuclei. Thus, although gck-1 and mop-25.2 both cause identical defects when ablated, they differentially affect CCM-3 localization in both the excretory canal and germline.

**Figure 4. mop-25.2 Is Required for Excretory Canal Extension and Germline Integrity**

(A and B) Knockdown of ccm-3, gck-1, and mop-25.2 causes significant canal truncations (B) (n = 50, *p < 0.05), with mop-25.2 significantly more truncated than either (*p < 0.05). The red asterisk denotes the distal worm measurement, and the yellow asterisk indicates the end of the canal. (C) Compared with wild-type (WT) canals, mop-25.2 RNAi causes lumen defects and distal tip cysts.

(D) Loss of either gck-1 or mop-25.2 in a ccm-3 mutant background exacerbates canal truncations, and similarly, loss of ccm-3 or mop-25.2 in a gck-1 mutant background causes more severe truncations (all n = 50, *p < 0.05).

(E) Ablation of mop-25.2 reduces endocytic recycling markers from the distal region of the canal (see Table S3).

(F) Cartoon illustrating a healthy (wild-type) C. elegans germline and a ccm-3 mutant with a damaged germline.

(G) In ccm-3 mutants, the lumen bordered by the rachis membrane collapses (white arrow), resulting in underdeveloped oocytes (blue overlays).

(H) Using fluorescent membrane and nuclear markers (GFP::PH [PLC1delta1], mCherry::his-58) to visualize the germline, knockdown of mop-25.2 phenocopies ccm-3 mutants. Rachis collapse (yellow asterisks) and small oocytes (blue asterisks) was observed in ~60% of worms treated with mop-25.2 RNAi (n = 50).

Scale bars, 50 μm (A) and 10 μm (C, E, G, H). See also Table S3 and Figures S4 and S5.

**mop-25.2 Is Required for CCM-3 Localization**

Previously, we showed that ablation of gck-1 in the germline caused CCM-3 to relocalize from the apical membrane to meiotic nuclei (Pal et al., 2017). Because mammalian MO25 has been shown to bind and stabilize GCKIII class kinases (Filippi et al., 2011), and a MO25-GCKIII-CCM3 ternary complex may exist (Yin et al., 2012), we asked if MOP-25.2 affects CCM-3 localization in a manner similar to GCK-1. CCM-3 localizes to apical membranes in both the canal and germline (Lant et al., 2015; Pal et al., 2017) (Figures 5 A and 5B, top). Using GFP translational reporters for CCM-3, we measured the fluorescence of GFP::CCM-3 in the excretory canal (along the lumen) and the pachytene region of the germline. Knockdown of ccm-3 caused the expected dissipation of canal GFP::CCM-3 (Figure 5A, second panels), as observed before, loss of gck-1 caused CCM-3 relocalization into pachytene nuclei (Figure 5B, third panels). However, loss of mop-25.2 significantly diminished GFP::CCM-3 signal on the lumen (Figure 5 B, fourth panel, and Figure 5D; p = 1.02 × 10^-13) but did not cause its relocalization to nuclei. Thus, although gck-1 and mop-25.2 both cause identical defects when ablated, they differentially affect CCM-3 localization in both the excretory canal and germline.

**mop-25.2 Is Required for GCK-1 Localization**

Because mop-25.2 knockdown dramatically affected CCM-3 localization, we next asked if it had a similar effect on GCK-1.
Figure 5. *mop-25.2* Is Required for Membrane Localization of CCM-3 and GCK-1

(A, B, E, and F, top panels) Both CCM-3 and GCK-1 localize to apical membranes of the excretory canal and germline. (A) *ccm-3* RNAi diminishes GFP::CCM-3 signal in excretory canals as expected (C; n = 50, *p* < 0.05), whereas knockdown of *gck-1* causes CCM-3 to relocalize into discrete puncta along the apical membrane (yellow asterisk). Knockdown of *mop-25.2* diminishes CCM-3 signal significantly (n = 50, *p* < 0.05). (B) GFP::CCM-3 localizes to rachis membrane in the germline (yellow arrow) around openings to the rachis (yellow asterisk) but not within the cell nuclei themselves (red asterisks). (C) *ccm-3* RNAi diminishes GFP::CCM-3 signal in excretory canals as expected (n = 50, *p* < 0.05), whereas knockdown of *gck-1* causes CCM-3 to relocalize into discrete puncta along the apical membrane (yellow asterisk). Knockdown of *mop-25.2* diminishes CCM-3 signal significantly (n = 50, *p* < 0.05). (D) Depletion of *gck-1* causes GFP::CCM-3 relocalization to germ cell nuclei, whereas *mop-25.2* RNAi reduced its levels on the rachis membrane (n = 50, *p* < 0.05). (E) GCK-1 in the canal is frequently punctate (yellow asterisks). (F) In the germline, *gck-1* RNAi completely ablates GFP::GCK-1 signal. (G) Ablation of *gck-1* or *mop-25.2* reduced GFP::GCK-1 signal (n = 50, *p* < 0.05), but knockdown of *ccm-3* did not (see E), although it reduced punctate localization by half (n = 50, *p* < 0.05). (H) Depletion of *ccm-3* or *mop-25.2* significantly reduced GFP::GCK-1 membrane localization (n = 50, *p* < 0.05). *mop-25.2* RNAi also caused perinuclear accumulation of GCK-1 (red asterisks) and patchy cytoplasmic localization of GFP::GCK-1 in oocytes (blue asterisks).
GCK-1 co-localizes in a punctate manner with CCM-3 along the apical membrane of the canal (Figure 5E, yellow asterisks) and the rachis membrane of the germline (Figure 5F, yellow arrow/asterisk). Ablation of \( gck-1 \), as expected, caused signal to disappear in both tissues (Figures 5E and 5F). In the canals, ablation of \( ccm-3 \) did not change either GCK-1::mCherry localization or its fluorescence intensity (\( p = 0.18 \)) (Figure 5E, third panel), but it did reduce the punctate distribution of GCK-1 to \(~30\%\) (16 of 50, \( p = 0.0086 \)). Similarly, in the germline, loss of \( ccm-3 \) did not cause relocalization of GCK-1 but significantly diminished its signal (\( p = 4.35 \times 10^{-16} \) along the rachis membrane (Figure 5F)). As with GFP::CCM-3, knockdown of mop-25.2 caused a significant reduction of GCK-1 signal in both the canal and germline (Figures 5E and 5F, fourth panels, and Figures 5G and 5H). Interestingly, ablation of mop-25.2 also reduced the punctate localization of GCK-1::mCherry (\( p = 0.0022 \)) in the canal and increased its perinuclear and patchy cytoplasmic localization in oocytes (\( p = 0.00028 \)) (Figure 5F, fourth panel insets, red and blue asterisks, respectively). Thus, MOP-25.2 regulates the integrity of the excretory canal and germline, most likely by controlling the localization of CCM-3 and GCK-1.

Knockdown of CCM3 and MO25 Causes Stress Fibers in Endothelial Cells

In order to assess if the role of mop-25.2 is conserved, we ablated both CCM3 and MO25 in human umbilical vein endothelial cells (HUVECs) and assessed stress fiber (SF) formation. Loss of CCM proteins cause actin stress fibers through increased RhoA/ROCK activity (Richardson et al., 2013; Stockton et al., 2010). Staining for both VE-cadherin and F-actin, we observe the change in dynamics between the two (Abu Taha and Schnittler, 2014). With control scrambled (SCR) small interfering RNA (siRNA), VE-cadherin and F-actin largely co-localize at cell junctions (Figures 6A–6C). However, siRNA to both CCM3 and MO25 lock the actin into transversal arcs that cross the nucleus and stretch across the cell (Tojkander et al., 2012). Prevalence of stress fibers in cell populations (Figure 6D) showed significant increases with siRNA to CCM3 and MO25 (\( p = 0.0083 \) and \( p = 0.018 \), respectively, \( n > 300 \)), indicating that loss of MO25 phenocopies loss of CCM3 in endothelial cells.

DISCUSSION

Here, we took advantage of the genetic tools of \( C. \) \( elegans \) to interrogate the CCM gene network, focusing on the less studied CCM3 branch. Our RNAi screen identified 562 genes that cause negative genetic interactions in \( kri-1/CCM1 \) mutants, similar to

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**Figure 6. Loss of MO25 Causes Stress Fiber Formation in Endothelial Cells**

(A–D) Pooled HUVECs transfected with control siRNA (A), siCCM3 (B), and siMO25 (C) and incubated with antibodies against VE-cadherin and phalloidin (F-actin) were quantified for the formation of transverse actin fibers (SFs) (D). Loss of both CCM3 and MO25 caused a significant increase in stress fiber formation (\( n = 3 \) independent experiments, \(* p < 0.05 \)).

\( p = 5.11 \times 10^{-09} \) and \( p = 3.24 \times 10^{-14} \), respectively) (Figures 5E and 5F, fourth panels, and Figures 5G and 5H).
ablation of ccm-3. Although we focused on the 237 genes with annotated interactions, greater data integration of the “orphan genes” in the future should provide more insights into their gene-gene interactions and biological functions. Many of the genes with well-documented interactions clustered into a limited but functionally diverse number of biological processes, including protein translation, cytoskeletal dynamics, and vesicle trafficking (Figure 2; Table S1). This is consistent with our previous work showing a role for CCM3 in these processes to control biological tube integrity (Lant et al., 2015; Pal et al., 2017). Our screen uncovered several expected hits, including genes encoding small GTPases, Notch, MAPK, and KLF (Table S1), confirming the robustness of this method (Borikova et al., 2010; Ma et al., 2007; You et al., 2013; Zhou et al., 2015). The identification of cell cycle, histone, and ion channel genes (Figure 2; Table S1), expands the number of biological processes affected by CCM signaling, which will be interesting to investigate in the future.

We were particularly interested in characterizing the functions of genes that are conserved in humans. Using the bioinformatics platforms GeneMANIA and STRING (von Mering et al., 2005; Mostafavi et al., 2008) to generate a human CCM-ome allowed us to interrogate a limited set of genes predicted to have conserved functions in CCM biology. Remarkably, of the 29 tier 1 genes, 14 caused significant excretory canal truncations when ablated. Genes that regulate canal integrity are uncommon (Buechner et al., 1999; Sundaram and Buechner, 2016), and of these 14 genes, 3 were previously known to affect the excretory canal: glp-1 (Abdus-Saboor et al., 2011), bab-5 (Mattingly and Buechner, 2011), and cdc-42 (Lant et al., 2015). The remaining 11 genes exhibit previously undocumented excretory cell phenotypes. Because the tier 1 genes might also function in the KRI-1/CCM1 pathway, we evaluated their role in germline apoptosis and found that 5 genes (Figure S3E) conferred resistance to radiation-induced germline apoptosis when ablated, similar to kri-1. Although this is not as striking as the frequency of genes that affect the excretory canal, it is possible that we have uncovered some genes in which the CCM-3 and KRI-1 pathways converge. Only ablation of his-13, exceeded “kri-1-level” apoptosis suppression, and while his-13 has not been reported to affect IR-induced germline apoptosis, histones affect the expression of many genes, so it will be interesting to determine if his-13 has a direct role in CCM signaling.

Tier 1 genes causing ccm-3-like canal truncations included a number of small GTPases and small GTPase regulators, which is not surprising given that RhoA is hyper-activated in the absence of CCM proteins (Richardson et al., 2013; Stockton et al., 2010). We previously reported that CCM-3 regulates another small GTPase, CDC-42 (Lant et al., 2015), which was also identified as a tier 1 hit in this study (Figure S3; Table 1). In addition, we identify a number of GTPase related genes (Figure 2), including the Rho GEF epx-1, which has been reported to regulate CDC-42 activity in both nematodes and human cells (Oliver et al., 2011; Walck-Shannon et al., 2016). Therefore, we predict that a more extensive network of small GTPases are involved in CCM signaling.

Consistent with our previous work showing that CCM3/STRIPAK regulates endocytic recycling (Lant et al., 2015; Pal et al., 2017), several tier 1 genes encode vesicle-trafficking proteins, including the early endosomal GTPase rab-5 and exocyst complex genes sec-3 and sec-15. This is significant because we previously discovered a link between CCM-3 and the exocyst complex gene exoc-8 (Lant et al., 2015), which was also identified in the RNAi screen. Work by Armenti et al. (2014) identified important roles for the exocyst gene sec-5 and the exocyst complex regulator ral-1 in excretory canal development, with localization of exocyst proteins to apical membrane mediated by polarity proteins. Combined, our work showing that CCM-3 is required for proper localization of the polarity proteins PAR-2 and PAR-6 (Pal et al., 2017) suggests that CCM-3 may coordinate polarity establishment and endocytic trafficking during biological tube development.

Many of the tier 1 genes had not been previously implicated in functions related to CCM-3/CCM3 signaling, and of these, the most intriguing gene to us was the MO25/CAB39 homolog mop-25.2. Not only did its ablation cause the strongest canal truncations, but it also had similar germline defects as observed in ccm-3 and gck-1 mutants (small oocytes and rachis collapse) (Figure 4). Although mop-25.2 is required for localization of CCM-3 to apical membranes, its co-ablation with ccm-3 or gck-1 caused additive effects on canal truncations (Figure 4D). We propose that MOP-25.2 functions to integrate CCM-3/GCK-1, and likely other factors, to promote canal extension and germline integrity. Although mammalian MO25 is best known to regulate AMPK signaling, its role in C. elegans tubulogenesis is independent of its canonical binding partners LKB1/STRAD and AMPK (Figure S5). Canal defects caused by knockdown of par-4 and mop-25.2, but not strd-1 and the AMPK genes, are interesting in light of observations showing that par-4 and strd-1 function independently of their canonical pathway (Narbonne and Roy, 2009; Narbonne et al., 2010).

The impact of losing mop-25.2 on CCM-3 localization at the canal lumen and germline rachis is distinct from gck-1 loss, which caused CCM-3 to localize into discrete puncta in the canal (Figure 5A) and nuclei of the meiotic germline (Figure 5B) (Pal et al., 2017). Interfering with binding to GCK-1 causes CCM-3 to relocalize to the nucleus, whereas loss of binding to CASH-1 (striatin) disrupts apical membrane localization but does not nuclear localization (Pal et al., 2017). Loss of mop-25.2 also caused GCK-1 to become dissipated from apical membranes in both tissues, showing that it has a CASH-1-like effect on localization of the CCM-3/GCK-1 complex to apical membranes. It will be interesting to determine whether MOP-25.2 interacts with the CASH-1 branch of the CCM-3 network to direct CCM-3/GCK-1 to luminal membranes. However, because we did not identify MOP-25.2 as a binding partner of the CCM-3/STRIPAK complex (Pal et al., 2017), it is possible that it might affect the synthesis or stability of CCM-3, which would subsequently alter GCK-1 localization. Although this collectively supports the co-dependency of CCM3 and GCKs in localization to apical membranes (Fidalgo et al., 2010, 2012; Ma et al., 2007; Zhang et al., 2013b), it also reveals a role for MOP-25.2 in the CCM-3/STRIPAK network. Clearly, there is much to learn about the non-canonical roles of MO25/MOP-25.2.

Because it shares a similar function with CCM3 in stress fiber formation in cells, MO25 may be an important determinant of CCM disease. Prevention of stress fiber formation by inhibiting RhoA/ROCK (Richardson et al., 2013; Stockton et al., 2010)
downstream of the hetero-trimeric complex is an attractive therapeutic strategy for managing this disease (McDonald et al., 2012; Shenkar et al., 2017). Our observations suggest that MOP-25.2/MO25 affects the stability of CCM-3/CCM3-containing complexes, and we suspect that the severity of disease in CCM3 patients may be due to its loss from multiple protein complexes. Exploiting the powerful genetics of *C. elegans* has provided insights into how CCM3 functions *in vivo* and uncovered nodes in this network that might offer therapeutic strategies for managing this disease in humans.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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**SUPPLEMENTAL INFORMATION**

Supplemental Information includes five figures and four tables and can be found with this article online at https://doi.org/10.1016/j.celrep.2018.08.039.

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**AUTHOR CONTRIBUTIONS**

B.L. performed experiments, analyzed and assembled data, contributed to project design, and wrote the manuscript. S.P. and E.M.C. performed germline experiments. E.F. performed endothelial cell experiments, and analysis with support from C.A.-R. B.Y. contributed to the generation of transgenic strains and optimized screening with B.L. B.L., S.P., B.Y., D.W., S.C., and L.Z. performed the whole-genome screen. W.B.D. conceived and directed the project and contributed to manuscript writing.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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### STAR METHODS

#### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Anti-VE-Cadherin Antibody (CD144), clone BV9 | Millipore Sigma/Merck | Cat # sc-52751 (RRID: AB_628919) |
| TRITC-conjugated phalloidin | Sigma Aldrich | FAK100 part 90228 |
| Goat anti-Mouse IgG (H+L) Highly Cross-Adsorbed Secondary Antibody, Alexa Fluor Plus 488 | Invitrogen | Cat # A32723 (RRID: AB_2633275) |
| **Bacterial and Virus Strains** |        |            |
| Caenorhabditis elegans RNAi feeding library | Source Bioscience | https://www.sourcebioscience.com/products/life-science-research/clones/mai-resources/c-elegans-mai-collection-ahringer/ |
| OP50 (E. coli) | Caenorhabditis Genetics Center | WormBase ID: OP50 |
| **Chemicals, Peptides, and Recombinant Proteins** |        |            |
| Ampicillin | Sigma-Aldrich | A1593; CAS Number 69-53-4 |
| Tetracycline hydrochloride | Sigma-Aldrich | T7660; CAS Number 64-75-5 |
| Isopropyl-β-D-thiogalactopyranoside (IPTG) | Sigma-Aldrich | I1284; CAS Number 367-93-1 |
| Tetramisole | Sigma-Aldrich | T1512; CAS Number 5086-74-8 |
| Penicillin/Streptomycin | Sigma-Aldrich | F0895 |
| Fibronectin Human Plasma | Sigma-Aldrich | F0895 |
| Rat Tail Collagen I | Corning | Product # 354236 |
| Vasculife VEGF Medium | Lifeline Cell Technology | LL-0005 |
| EBM-2 Basal Medium | Lonza | CC-3156 |
| Lipofectamine RNAi max | Life Technologies | ref. 13778-150 |
| Trypsin inhibitor | Sigma-Aldrich | T6522 |
| Mowiol | Sigma-Aldrich | 324590 |
| **Experimental Models: Cell Lines** |        |            |
| C. elegans: Strain N2 | Caenorhabditis Genetics Center | WormBase ID: N2 |
| C. elegans: Strain BK204: qpis96[Pesc-9::mCherry::cdc-42] | The Buechner Laboratory | WormBase ID: BK204 |
| C. elegans: Strain BK205: qpis97[Pesc-9::mCherry::rab-11]V | The Buechner Laboratory | WormBase ID: BK205 |
| C. elegans: Strain BK220: qpis103[Pesc-9::mCherry::GFP] | The Buechner Laboratory | WormBase ID: BK220 |
| C. elegans: Strain BK262: qpis104[Pesc-9::mCherry::GBDwsp-1] | The Buechner Laboratory | WormBase ID: BK262 |
| C. elegans: Strain OD95: unc-119(ed3) III; fts37[pAA64; pie-1::mCherry::HIS-58 + unc-119(+)] IV; fts38[pAA1; pie-1::GFP::PH(PLC1delta1) + unc-119(+)] | Caenorhabditis Genetics Center | WormBase ID: OD95 |
| C. elegans: Strain WD187: kri-1(ok1251)III; bgls312[Pvha-8::GFP] | The Derry Lab | N/A |
| C. elegans: Strain WD188: ccm-3(tm2806)/mlm1[mls14 dpy-10(e128) let-60(k838)]; bgls312[Pvha-8::GFP] | The Derry Lab | N/A |
| C. elegans: Strain WD355: onls3[Pcmm-3::GFP::CCM-3; unc-119(+)]III; unc-119(ed3) III | The Derry Lab | N/A |
| C. elegans: Strain WD419: N2; onEx62[Pesc-9::ccm-3::GFP; Pmyo-2::mCherry; Prab-8::mCherry] | The Derry Lab | N/A |
| C. elegans: Strain WD423: N2; onEx69[Pesc-9::ccm-3::GFP; Pesc-9::gck-1::mCherry; rol-6(su1006)] | The Derry Lab | N/A |

(Continued on next page)
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Dr. Brent Derry (brent.derry@sickkids.ca).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

**Caenorhabditis elegans strains and maintenance**

Worms were cultivated on lawns of *E. coli* (strain OP50) grown on NGM (nematode growth medium) plates at 20°C (unless otherwise stated) (Brenner, 1974). Wild-type (WT) refers to the *C. elegans* variety Bristol, strain N2. Compound mutant and transgenic strains were constructed according to standard *C. elegans* protocols (Fay, 2008). See KRT for strain list. Excretory canal marker strains designated ‘BK’ were a kind gift from Dr. Matthew Buechner.

**Endothelial cell growth conditions**

Pooled HUVECs were grown on collagen I coated plates in complete EBM media, supplemented with 100U/ml penicillin and 100 μg/ml streptomycin, at 37°C in a 5% CO2-humidified chamber according to the manufacturer’s instructions. HUVEC (1.5 × 10⁶ cells) were transfected twice at 24 h intervals with 20 nM siRNA and 45 μL lipofectamine RNAi max according to the manufacturer’s instructions.

**METHOD DETAILS**

**RNA interference**

RNAi was performed by feeding bacteria expressing double-stranded RNA (RNAi) to worms at the L1 stage, according to Kamath et al. (Kamath et al., 2001). Bacterial cultures expressing double-stranded RNA targeting the gene of interest were drawn from the Source BioScience RNAi library, and grown on nematode growth media (NGM) plates supplemented with antibiotics (50 μg/ml ampicillin and 12.5 μg/ml tetracycline) and 0.1 mM isopropyl-β-D-thiogalactopyranoside to induce dsRNA production. Bacteria expressing an empty RNAi vector (L4440; within competent HT115 cells) were used as controls. For excretory canals analysis, to decrease variance between worms, RNAi was concentrated 5X, and worms were grown on RNAi bacteria for two generations. In the cases where worms did not survive two full generations of RNAi, L4s were placed on the bacteria, and subsequent F1 progeny were analyzed.

**Whole Genome RNAi Screen**

We previously observed that ablation of ccm-3 and kri-1 causes synthetic lethality (Lant et al., 2015). Hence, we reasoned that other genes that cause synthetic conditions (‘sickness’) when knocked down in *kri-1(ok1251)* mutants might reveal additional components...
of the ccm-3 pathway. kri-1(ok1251) mutants were fed bacteria expressing dsRNA to genes from the Source Bioscience RNAi library in liquid culture format (96 well plates). A 96 pin-replicator was plunged into the wells of a thawed RNAi library plate (4 pins/plates per 384 well library plate), then immersed in 150μl of LB+AMP+TET, aliquoted into each well of a conical tip 96 well plate. The plates were covered in parafilm and a wet paper towel, placed in a sealed plastic bag to prevent evaporation, then incubated on a shaker overnight at 37°C. The following day bacterial cultures were induced with 6μl IPTG (0.1M) per well, and incubated for one more hour. Bacteria was pelleted by centrifugation (3,500rpm) for 5 min, the supernatant removed, and pellets re-suspended in 150μl liquid NGM per well. 50μl of this induced bacterial RNAi culture was transferred to a flat-bottom 96 well plate and 10 L1 stage kri-1(ok1251) mutant worms were added. The plates were sealed (as above) and placed on an orbital shaker at room temperature and grown 4-5 days. Sickness (embryonic lethality, larval lethality, sterility, and slow growth) of F1 progeny from kri-1 mutants was assessed and cataloged. Concurrently, N2 (WT) controls were grown in the same RNAi conditions (in separate wells). Negative genetic interactions based on synthetic lethal and sick phenotypes were categorized as those observed in kri-1 but not N2 controls.

C. elegans microscopy
Nematodes were slide mounted in a 5μl aliquot of tetramisole anesthetic (20-100 mM; higher concentration for roller strains), on a flat pad of 4% agar. Nematodes were imaged at 63X or 100X magnification using a Leica DMRA2 compound microscope equipped differential interference contrast (DIC) optics and epifluorescence. Images were captured with a Hamamatsu C472-95 digital camera using Openlab software (PerkinElmer Inc.).

Bioinformatics
A master list of 35 genes (from mammalian CCM literature was converted into gene and protein IDs via BioMart (Ensembl), and then used to generate ‘CCM-omes’ via GeneMANIA and STRING, respectively. For the gene lists a total of 500 were obtained by selecting for biological relevance using GO term sorting for GO Biological Process, GO Molecular Function, and GO Cellular Component in GeneMANIA. The amalgamated GeneMANIA list (i.e., removing multiple entries for the same genes) amounted to 769 genes. For the protein interaction list STRING was used to generate a list of 500 proteins, using its automatic weighting system. These lists were converted to worm orthologs via BioMart, resulting in 535 independent genes from the GeneMania list and 231 genes from the STRING list. Further amalgamation of these two lists produced a final worm “CCM-ome” of 653 genes. The overlapping genes between the ‘Sick’ list from the RNAi screen (562 genes) and the CCM-ome list (653 genes) were termed “Tier 1” hits, which amounted to 29 genes (‘Experimental Set’) that were examined for canal effects against 29 randomly selected (‘Control Set’) genes. The ID of Control set genes were generated by assigning a number to each gene in the Source Bioscience RNAi library, and then generating a random list of 29 numbers using the RAND function in Microsoft Excel.

Cell spreading and immunofluorescence
Transfected cells were trypsinized, treated with 1 mg/ml trypsin inhibitor, and incubated in serum-free EBM-2/1% BSA for 30 min at 37°C. Sparse HUVECs (10^4) were spread for 4h on 24-well plates slides coated with 2 μg/ml of Fibronectin (FN), in vasculife VEGF media containing 5% FN-depleted serum and then fixed with 4% PFA. Confluent HUVECs (2 × 10^5 cells) were seeded in 24-well plates on slides coated with 20 μg/ml FN and incubated for 72h in complete vasculife media. Cells were fixed with 4% PFA, permeabilized with 0.2% Triton X-100, and incubated with VE-cadherin BV9 antibody at 1/200 dilution. After rinsing, coverslips were incubated with an Alexa Fluor 488-conjugated secondary antibody and phalloidin-TRITC. The cells were mounted in Mowiol/DAPI solution and imaged on epifluorescent Axiomager microscope (Zeiss) with AxioCamMRc camera.

QUANTIFICATION AND STATISTICAL ANALYSIS
Significance of the Tier 1 set (29 genes) of hits generated from overlapping genes in the informatics set (653 genes) and the genome screen set (562 genes) was validated through a hypergeometric analysis and was found to be significant (p = 0.012). Additionally, randomizing sets of numbers (n = 10), the average overlapping values for comparisons of 653 and 562 numbers resulted in 18.2 ± 4.6 (mean ± standard deviation), wherein our value of 29 is outside of two standard deviations.

Excretory canal lengths were measured from excretory cell to the canal tip. These values were normalized by a ‘total’ length measurement of the worm (excretory cell to anus), with the ‘relative’ canal length expressed as a percentage according to Lant et al., 2015. Individual canal measurements were plotted in GraphPad Prism, with mean values marked by a red bar. Germline apoptotic corpse counts were performed 24 hr post IR, following a 60 gy dosage at the young adult stage. Germlines were observed and counted directly at 630X magnification using a Leica DMRA2 compound microscope. Individual germline arm corpse counts were plotted in GraphPad Prism, with mean values marked by a red bar. Values were accumulated from three or more experimental datasets. P values, relative to ‘control RNAi’ were determined using a two-tailed Student’s t test, assuming equal variance. Significance levels (P value) and sample size (N) are marked in each corresponding figure legend.

Comparison of Control and Experimental sets of data, cumulatively, using a two tailed Student’s t test, assuming equal variance, showed the two sets to be significantly different (p = 6.92762E-36). Comparison of Control and Experimental sets, as collections of genes with a rating of no truncation, truncation, or strong truncation was conducted using the CHI square test and showed the two groups to have significantly different distributions in these categories (p = 0.0031).
Puncta counts and fluorescent signal measurements were obtained using ImageJ software. Puncta counts were measured (Lant et al., 2015) using an RGB profile plot on a stretch of canal (~100μm in length) at or near the posterior distal tip. Each representative image shows a single plane, of one canal, with the maximal amount of puncta present. Relative intensity of GFP/RFP signals were assessed by measuring the requisite RGB channel signal, and normalizing with RGB signal from surrounding tissue (canals), or within the pachytene cells (germline rachis) in which a neutral ‘non-signal’ holds a value of 1. In each case, values have been compared from the given RNAi type to a control (empty vector) RNAi.

For quantification of the number of cells displaying transversal actin fibers (stress fiber) above their nucleus, more than 100 cells per phenotype in 3 independent experiments were counted, and the relative frequency of stress fiber presence was normalized for control siRNA numbers.