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A combined binary interaction and phenotypic map of C. elegans cell polarity proteins

Thijs Koorman1, **Diana Klompstra**2,3,* , **Monique van der Voet**1,4,* , **Irma Lemmens**5, **João J. Ramalho**1, **Susan Nieuwenhuize**1, **Sander van den Heuvel**1, **Jan Tavernier**5, **Jeremy Nance**2,3, and **Mike Boxem**1,6

¹Division of Developmental Biology, Department of Biology, Faculty of Science, Utrecht University, 3584 CH, Utrecht, The Netherlands ²Helen L. and Martin S. Kimmel Center for Biology and Medicine at the Skirball Institute of Biomolecular Medicine, NYU School of Medicine, New York, New York 10016, USA ³Department of Cell Biology, NYU School of Medicine, New York, New York 10016, USA ⁵Department of Medical Protein Research, VIB, and Department of Biochemistry, Faculty of Medicine and Health Sciences, Ghent University, 9000 Ghent, Belgium

Abstract

The establishment of cell polarity is an essential process for the development of multicellular organisms and the functioning of cells and tissues. Here, we combine large-scale protein interaction mapping with systematic phenotypic profiling to study the network of physical interactions that underlies polarity establishment and maintenance in the nematode *Caenorhabditis* elegans. Using a fragment-based yeast two-hybrid strategy, we identified 439 interactions between 296 proteins, as well as the protein regions that mediate these interactions. Phenotypic profiling of the network resulted in the identification of 100 physically interacting protein pairs for which RNAi-mediated depletion caused a defect in the same polarity-related process. We demonstrate the predictive capabilities of the network by showing that the physical interaction between the RhoGAP PAC-1 and PAR-6 is required for radial polarization of the C. elegans embryo. Our network represents a valuable resource of candidate interactions that can be used to further our insight into cell polarization.

⁶Correspondence should be addressed to M.B. (; Email: m.boxem@uu.nl)

4Present address: Department of Human Genetics, Donders Institute for Brain, Cognition and Behaviour, Radboud university medical center, 6500 HB Nijmegen, The Netherlands.

Competing Financial Interests

The authors declare no competing financial interests.

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^{*}These authors contributed equally

Author Contributions

M.v.d.V. contributed to the cloning and Y2H screens. D.K. performed the PAC-1 rescue experiments. I.L. performed the MAPPIT experiments. J.J.R. and S.N. contributed to the RNAi screens. M.B. performed the computational analyses. All other experiments were performed by T.K. S.v.d.H. and T.K. contributed to the design of the study. J.N. contributed to the design of the PAC-1 PAR-6 experiments. J.T. contributed to the design of the MAPPIT experiments. T.K. and M.B. wrote the manuscript. M.B. guided all aspects of the study.

Introduction

The ability to polarize is a fundamental cellular property, required for processes such as cell migration and asymmetric cell division, and for the specification of functionally distinct domains. A series of cortically localized proteins has been identified that can drive the establishment of cell polarity. The PAR3-PAR6-aPKC and Crumbs-SDT-PATJ complexes together promote apical domain identity 1^{-3} , while the LGL-SCRIB-DLG proteins, originally identified as *Drosophila* tumor suppressors⁴⁻⁶, promote basolateral identity². These cortical polarity complexes act together with a number of other components, such as Rho family GTPases, junctional components, and cytoskeletal linkers of the ERM family, in establishing polarity.

Although it is clear that mutual exclusion is a key mechanism by which cortical polarity regulators establish polarity^{7,8}, we still lack a detailed mechanistic understanding of how cortical polarity regulators are segregated into distinct domains. Moreover, we know little of the mechanisms through which cortical polarity is integrated with cellular events such as cytoskeletal rearrangement, organization of a polarized trafficking machinery, and functional specialization of membrane domains. A full understanding of polarity establishment will require a comprehensive knowledge of the proteins involved in this process and the molecular interactions between them.

Here, we study the network of physical interactions that underlies polarity establishment in the nematode *Caenorhabditis elegans* using a combination of large-scale yeast two-hybrid screens and phenotypic profiling. We identified a polarity interaction network of 439 interactions, and mapped the protein regions mediating these interactions. Phenotypic profiling by RNAi revealed 100 protein pairs that exhibited a phenotype in the same polarity related process. These pairs are strong candidates for a functional interaction in vivo. We studied the interaction between PAR-6 and PAC-1 ARHGAP21 in detail, and demonstrate that this physical interaction is important for radial polarization of the C. elegans embryo. Our data provides a resource for future studies into cell polarity, and should contribute to our understanding of this essential process. A searchable web interface of all interactions and fragments identified is available at [http://www.projects.science.uu.nl/interactome/.](http://www.projects.science.uu.nl/interactome/)

Results

Identification of the C. elegans polarity interaction network

To generate a map of interactions underlying polarity establishment in C. elegans, we selected 69 proteins that control cell polarity in C . elegans, or are homologous to known polarity regulators in other organisms (Fig. 1a and Supplementary Table 1). For each protein, a yeast two-hybrid (Y2H) bait construct was generated by cloning the full-length open reading frame (ORF) into a Gal4::DB vector. We previously showed that a greater number of interactions can be detected when multiple fragments of a protein are tested in the Y2H system⁹. Therefore, up to 12 additional fragment bait constructs were cloned for each protein (Fig. 1b and Supplementary Table 1). We successfully generated 65 full-length bait clones, and 338 fragment bait clones, which together represent all 69 polarity regulators (Supplementary Table 1).

Each of the bait constructs was used to screen two Gal4::AD libraries: a library that contains full-length and fragment clones of 749 genes that are essential for early embryonic development (AD-Fragment library)⁹, and a mixed-stage *C. elegans* AD-cDNA library (Fig. 1c). We eliminated auto-activators that arose during the screening process^{10, 11}, and interactions where the AD-ORF fusion was out of frame. To further increase the accuracy, we only included AD-Fragment library-derived interactions identified in 2 or more yeast colonies. The AD-cDNA library is more complex, and many valid interactions may only be identified in a single yeast colony. Hence, we experimentally retested all interactions identified only once, retaining those that retested positively (Fig. 1c). The final C elegans polarity interaction network (CePIN) contains 439 interactions between 296 proteins (Fig. 1d and Supplementary Table 2). Most interactions (359/439) were detected only using nonfull-length bait constructs, confirming that the use of protein fragments increases the detectability of interactions⁹. The network contains 54 interactions that have previously been reported, including 19 interactions that have been studied in detail (Supplementary Table 3).

Quality assessment and experimental validation of the polarity interaction network

To assess the quality of the CePIN we examined whether interacting protein pairs shared other characteristics that indicate a functional association. We found that interacting protein pairs were enriched for similar GO terms, as well as for presence in WormNet, which predicts functional linkages between C. elegans genes¹² (Fig. 2a,b and Supplementary Fig. 1). In addition, interactions identified from AD-cDNA library screens were highly enriched for similar mRNA expression profiles (Fig. 2c and Supplementary Fig. 1d).

We next retested a random sample of 33 interactions by co-affinity purification from Human Embryonic Kidney (HEK) 293 cells. Each interaction was tested using both a full-length prey protein and a shorter prey protein fragment containing the minimal region of interaction. We successfully reproduced the interaction between 16 of the 33 pairs tested (48%) (Fig. 2d, Supplementary Fig. 2, and Supplementary Table 4). This is comparable to published validation rates of binary interactions in orthogonal assays^{13,14} and to our own previous observations using this approach¹⁵. For 12 of the validated interactions, both the full-length and fragment prey were expressed. Of these, 6 were detected only using the fragment prey protein. Thus, the use of fragments provides an advantage in co-affinity purifications as well.

Finally, we retested 93 randomly selected protein pairs using the MAmmalian Protein-Protein Interaction Trap (MAPPIT), which detects interactions between proteins in living cells based on restoration of the signaling capacity of a signaling-deficient cytokine receptor^{16,17} (Supplementary Table 4). We examined the fraction that tested positively at a range of assay stringency thresholds, and compared the results with three previously published *C. elegans* MAPPIT analyses¹⁸: a positive reference set of 46 low-throughput literature-curated interactions (CePRS), a set of 86 random protein pairs that serves as a negative control (CeRRS), and a set of 87 interactions detected in a large scale Y2H screen $(WI-2007)^{18}$. The retest rates of the *CePIN*, *CePRS* and WI-2007 were statistically indistinguishable (Fisher's exact tests), while scoring significantly higher than the CeRRS control pairs (Fig. 2e). Together with the identification of previously described interactions,

the computational and experimental validations confirm the quality of the polarity interaction network.

Identification of minimal regions of interaction (MRIs)

For each interaction, we defined the MRI as the smallest region shared by all interacting protein fragments (Supplementary Table 5 and Supplementary Fig. 3). The average length of all MRIs identified was 408 amino acids or 60% of the respective full-length protein (Fig. 3a,b and Supplementary Fig. 4). To evaluate the accuracy of the MRIs, we compared them with previously described interaction domains (Fig. 3c, Supplementary Fig. 4g, and Supplementary Table 5). Out of 29 MRIs for which corresponding interaction domains were previously identified, 27 are consistent with the domain described (93%). In the two cases where the MRI differed from the described domain, the published interaction involved a mammalian homolog of the *C. elegans* protein. These may reflect a difference between the mammalian and C. elegans proteins. Further evidence of the accuracy of the MRIs comes from the co-affinity purification experiments, where 10/19 interactions for which the MRI was tested scored positively. Thus, the MRIs identified by Y2H were able to mediate the interaction in an orthogonal binary interaction assay as well.

While some MRIs are a near exact match to the known interaction site, others span a larger protein region (Fig. 3c). One explanation is that shorter clones were not identified or are not present in the library. For example, the LIN-10 MRI that binds LIN-2 was defined from ADcDNA clones, which can only define the N-terminal MRI boundary. Alternatively, the interaction may be mediated by a short, linear motif that needs to be presented as part of a larger, folded polypeptide. For example, the HMP-2 MRI covered a much larger region than the C-terminal 4 residues known to mediate the interaction with MAGI-1 $[ref^{19}]$. Especially in these cases, the MRIs identified by Y2H can provide a starting point for experiments requiring expression of an interacting fragment.

Phenotypic profiling of the polarity interaction network

Further evidence of a functional relationship can be obtained by integrating protein interaction data with phenotypic data^{20_23}. We therefore performed phenotypic profiling by RNA-mediated interference (RNAi). Polarity regulators may act only in certain tissues or play highly distinct roles in polarity establishment. Therefore, we examined the effects of RNAi in nine different strains expressing fluorescently-tagged proteins involved in several polarity related processes (Supplementary Table 6 and Fig. 4a–i). To examine defects in early embryonic polarity, we used a strain expressing mCherry::PAR-6, which localizes anteriorly in the one cell embryo and to the contact free surfaces in young embryos. In the larval stages, we examined three different epithelial cell types: seam cells, intestinal cells, and the excretory canal. The seam cells exhibit both apical-basal polarity and anteriorposterior tissue polarity, and undergo a series of asymmetric stem cell-like divisions during larval development. The seam cell epithelium was examined in a strain expressing a GFP::PH domain fusion that outlines the cell membrane, and a GFP::H2B fusion that marks the $DNA²⁴$. Intestinal polarity was followed using a strain expressing PEPT-1::DsRed, which marks the apical domain, and GFP::RAB-11, which marks apically enriched recycling endosomes²⁵. In both these tissues we also examined the integrity of cell junctions using a

DLG-1::GFP expressing strain. The excretory canal is a tubular epithelium formed by a single cell, and therefore highly distinct from the multicellular seam and intestinal epithelia. We examined defects in excretory canal development using a strain expressing VHA-5::GFP. In addition to these epithelial tissues, we examined two processes that depend on cell polarity. Uptake of yolk proteins by oocytes is an endocytic process that requires the activity of the par-³, par-⁶, pkc-3 and cdc-42 genes, and we examined this process in a strain expressing the yolk-protein fusion VIT-2:: $GFP^{26,27}$. All genes that showed a phenotype with this strain were rescreened in a strain expressing the fluorescently tagged VIT-2 receptor RME-2::GFP, to determine if the observed defects were due to mislocalization of the receptor. Finally, neurons represent one of the most highly polarized cell types in the body. We examined the correct polarization of the axon in PVD neurons, in a strain expressing the fluorescently tagged presynaptic molecules mCherry::RAB-3 and SAD-1:: GFP^{28} . All genes that showed a defect in localization of these markers following RNAi were re-screened in a strain expressing myristoylated GFP in the PVD neuron, to examine changes to the extensive arborisation pattern of the PVD dendritic tree.

We first screened each of the 69 bait proteins for a total of 40 possible defects across the nine marker strains (Supplementary Table 6). For 44 bait proteins, downregulation by RNAi resulted in a detectable defect in at least one of the marker strains (Fig. 4j and Supplementary Table 6). Certain genes, e.g. par-6, $kin-19$, lit-1 and $dlg-1$, showed defects in most tissues examined, while others, such as *lgl-1* or *rheb-1* affected only a single tissue. This matches our expectation that many genes will play tissue-specific roles in polarity establishment, although it may also reflect incomplete inactivation by RNAi. Next, the binding partners of these 44 bait proteins were screened specifically in those strains in which a defect was detected. For 100 protein pairs, RNAi of the corresponding genes resulted in a defect in the same polarity-related process (Fig. 5a and Supplementary Table 6). We found no bias among these 100 pairs for interactions found using full-length or fragment baits.

To evaluate whether an overlap in phenotype predicts functional association, we first used hierarchical clustering to group the bait genes by phenotypic similarity (Fig. 4j). Several genes known to act together also clustered together, including the par genes with pkc-3 and $cdc-42$, the $lin-2,7,10$ genes, whose protein products form a complex, and the genes encoding the interacting proteins AJM-1 and DLG-1. Next, we determined whether an overlap in phenotype correlates with protein interaction, with GO similarity, and with prior description of an interaction. We found that the interaction network was significantly enriched for an overlap in phenotype compared to all possible non-interacting pairs of proteins in the network (2.3 fold, $p = 3.0 \times 10^{-15}$). To examine the correlation with GO term similarity, we used semantic similarity scores calculated with the HRSS algorithm²⁹. We defined a score of $\,0.1$ as low similarity and >0.9 as high similarity. When examining all possible pairs of proteins in the network, we found that pairs that overlap in phenotype are enriched for a high GO similarity score (1.6-fold, $p = 0.001$) and are depleted of a low GO similarity score (1.8-fold, $p = 3.0 \times 10^{-20}$). When analyzing only protein pairs that physically interact, we find that these pairs are already highly enriched for a high GO similarity score, which was not further enriched in the subset with an overlap in phenotype. However, this subset did show a significant depletion of pairs with low GO similarity (1.8-fold, $p = 0.006$). Thus, an overlap in phenotype positively correlates with high GO similarity, and negatively

correlates with low GO similarity. Finally, we examined interacting proteins that overlap in phenotype were more likely to have been described in the literature. We found that 9% (9/100) of interactions with an overlap in phenotype had previously been described, compared to 2% (5/232) of the remaining interactions (Supplementary Table 6). Taken together, these data indicate that the phenotypic profiling is able to capture interactions relevant in vivo and to cell polarity.

A strength of unbiased interaction mapping is that it may produce leads into the function of proteins. Indeed many of the interactions with an overlap in phenotype involve proteins not known to act in a common process. For example, we identified an interaction between SYS-1, a C. elegans β-catenin homolog, and HIPR-1, the C. elegans homolog of mammalian Huntingtin-interacting protein 1 (HIP1) and HIP1-related (HIP1R), which are thought to function in the endocytic pathway³⁰. Inactivation of either sys-1 or hipr-1 resulted in spreading of SAD-1 and RAB-3 vesicles into the dendrites of the PVD neuron (Fig. 5a panels 1–6). A neuronal defect is consistent with previous observations that hipr-1 modulates the presynaptic activity of C. elegans neurons, and with reported localization of HIP1 and HIPR1 to dendritic structures^{30, 31}. However, SYS-1 has only been described to function as a transcriptional co-activator with POP-1 $TCF^{32,33}$. Interestingly, SYS-1 also interacted with UNC-11, a homolog of the AP180 clathrin adapter, and RNAi against both sys-1 and unc-11 resulted in defects in yolk endocytosis (Supplementary Table 6). Not all phenotypes overlapped closely. For example, we identified an interaction between the Ezrin-Radixin-Moesin homolog ERM-1 and the uncharacterized protein C30B5.4. RNAi for both genes resulted in defects in the seam epithelium, though the nature of the phenotype differed (Fig. 5a panels 10–12). C30B5.4 contains an RNA-binding motif, and is homologous to human RBMX2, which is annotated as a spliceosome component. As a final example, we identified an interaction between DLG-1 Discs Large and LST-1, a nematode-specific protein. RNAi for both *dlg-1* and *lst-1* resulted in irregular morphology of the intestinal lumen (Fig. 5a panels 13–15), and a reduction in the subapical accumulation of RAB-11 vesicles (Fig. 5a panels 16–18). Consistent with an intestinal defect, LST-1 is expressed in the intestine³⁴. For each of these interactions, the physical association and overlap in phenotype suggest a functional association, even though it is not immediately obvious what the functional consequences of this association might be.

The phenotypic profiling of the 69 bait proteins led to the identification of several phenotypes not previously described. For example, inactivation of $lgl-1$, the C. elegans homolog of *Drosophila lethal giant larvae*, caused spreading of RAB-3 vesicles into the dendrites of the PVD neuron (Fig. 5b panel 2). C. elegans IgI-1 was shown to act redundantly with par-2 in early embryonic polarity establishment^{35, 36}, but a par-2independent function for $Igl-1$ had not yet been described. A dramatic effect on the PVD neuron was observed upon inactivation of the TCF transcription factor pop-1, which resulted in the formation of numerous additional PVD cell bodies and spreading of RAB-3 vesicles into dendrites (Fig. 5b panel 4). This may reflect additional cell divisions or cell-fate alterations in the V5 lineage that produces the PVD neuron, rather than a polarity defect, as pop-¹ is known to be required for cell fate decisions. Inactivation of SYS-1, which binds POP-1, also resulted in spreading of RAB-3 and duplication of the cell body (Fig. 5b panel 5).

We identified 9 genes whose inactivation resulted in defects in the excretory canal. Three of these, erm-1, par-6, and ral-1 had previously described roles in excretory canal formation^{37_39}. We also identified defects in canal appearance for *par-2* and *par-4*, which indicates a broader involvement of cell polarity regulators in excretory canal formation (Fig. 5 panels 7,8). Finally, mes-1 and vang-¹ showed a defect exclusively in the excretory canal (Fig. 5b panels 10,11). MES-1 is a receptor tyrosine kinase-like protein required for accurate positioning of the mitotic spindle in the P2 and EMS cells of the early embryo^{40,41}. VANG-1 is the C. elegans homolog of the planar cell polarity regulator Strabismus-Van Gogh^{42_44}. The identification of roles in excretory canal formation for mes-1 and vang-¹ may provide insights into the formation of this organ.

Interaction between PAR-6 and PAC-1 is required for radial polarization

As a final demonstration of the validity of our approach to identify functionally relevant interactions, we focused on the interaction between PAR-6 and PAC-1. In addition to the physical interaction, the phenotypic profiles of par- $6(RNAi)$ and pac- $1(RNAi)$ clustered together (Fig. 4j). PAC-1 ARHGAP21 is a RhoGAP protein required for radial polarization of the *C. elegans* embryo⁴⁵. In wild-type embryos, PAR-6 localizes to the outer, contact-free cell surface beginning at the late 4-cell stage. Cortical recruitment of PAR-6 is dependent on the Rho-family member CDC-42, which localizes along the entire cortex^{45,46}. PAC-1 localizes to cell contact sites, and is thought to trigger radial polarization by locally inactivating CDC-42, limiting the recruitment of PAR-6 by CDC-42 to the outer cell surfaces^{45,46}. Although PAR-6 and PAC-1 do not appear to co-localize in fully polarized cells, they are both found at cell contacts as polarity is initially established, and our observations suggest that a physical interaction between PAR-6 and PAC-1 contributes to their functions.

Additional fragments of PAR-6 and PAC-1 were tested by Y2H to narrow down the interaction sites. The PDZ domain of PAR-6 (amino acids 155 – 248) was sufficient to mediate binding to PAC-1 (Fig. 6a,c). A PAC-1 fragment lacking the final 7 residues was still able to interact with PAR-6, indicating that the PAR-6 PDZ domain interacts with an internal motif of PAC-1 (Fig. 6b,c). A 100 amino acid sequence just downstream of the PAC-1 RhoGAP domain (amino acids 1221 – 1328) was able to mediate binding to PAR-6 (Fig. 6b,c). We confirmed the interaction between PAR-6 and PAC-1 by co-affinity purification from mammalian HEK293 cells. A C-terminal fragment of PAC-1b (aa 1221 – 1604), as well as the minimal PAR-6 binding domain (aa $1221 - 1328$), co-purified with a PAR-6a fragment containing the PDZ domain (aa 133 – 265) (Fig. 6d). Moreover, when coexpressed in HeLa cells, PAC-1 and PAR-6 co-localize in a punctate pattern that was clearly distinct from the localization pattern of PAC-1 or PAR-6 expressed individually (Fig. 6e–g). These observations strongly support the presence of a physical interaction between PAC-1 and PAR-6.

To explore the functional relevance of the interaction between PAC-1 and PAR-6, we determined whether PAC-1 lacking the PAR-6 binding domain can functionally substitute for the wild-type protein. As previously demonstrated⁴⁵, expression of a GFP::PAC-1 fusion protein rescues the localization of PAR-6 in a pac-1 mutant embryo (Fig. 6j). We deleted the

PAR-6 binding domain (PBD) (aa 1221 – 1328) from the GFP::PAC-1 encoding plasmid and expressed GFP::PAC-1(PBD) in pac- $1(xn6)$ mutant embryos. GFP::PAC-1(PBD) was expressed at similar levels as GFP::PAC-1 (Fig. 6m) and localized to the inner cell surfaces (Fig. 6k'). However, GFP-PAC-1(PBD) failed to restore the radial localization pattern of PAR-6 (Fig. 6k and l). Thus, the direct interaction of PAC-1 with PAR-6 is necessary for PAC-1 to establish radial polarization.

Discussion

In this study we identified a *C. elegans* polarity interaction network of 439 interactions between 296 proteins, and validated the quality of the network using a combination of computational and experimental approaches. The experimental identification of the protein regions that mediated these interactions provides additional insight into the proteins in the network, as well as a starting point for studies needing to express functional protein domains.

Of the 439 interactions we identified, 385 had not been previously identified. Furthermore, out of 19 interactions studied in detail, only 12 had previously been identified in large-scale studies. These results confirm the ability of a smaller scale targeted approach to identify additional relevant interactions not present in high-throughput interaction datasets.

100 protein pairs physically interacted and, upon RNAi-mediated depletion, showed a defect in the same polarity-related process. An overlap in phenotype correlated with the presence of a physical interaction, with GO similarity, and with prior description of an interaction in the literature, confirming that the combination of interaction data with phenotypic data better predicts a functional relationship than presence of an interaction alone. The marker strains we used for the phenotypic profiling highlight only a subset of polarity-related processes. Thus, examining the interacting protein pairs using other marker lines should result in the identification of additional pairs with overlapping phenotypes.

The CePIN provides many putative polarity-related interactions, which can be pursued using detailed *in vivo* analyses that can identify more subtle or tissue-specific roles in cell polarity. Many of the mechanisms that drive polarity establishment are conserved between C . elegans, flies, and human, and interactions between conserved proteins can be examined in other model systems as well. Thus, the CePIN will be a valuable resource for future studies aiming to gain novel insights into the mechanisms that control cell polarity.

Methods

Primers and Linkers

All primers and oligonucleotide linkers are listed in Supplementary Table 7.

Y2H Vectors

Vectors pMB28 and pMB29 were derived from pPC97 and pPC86 respectively $47,48$. Both encode a flexible linker (GGGG) and AscI or NotI restriction sites, inserted as an oligonucleotide linker (linker 1) into the parent vectors digested with SmaI and SacI. pMB28

also contains the CAN1 gene, inserted into the ApaI site as a PCR product amplified from yeast genomic DNA (primers CAN1_F and CAN1_R). pMB29 contains the CYH2 gene, cut from Clontech vector pAS2-1 with EcoRV and ligated into vector digested with Acc65I and treated with Klenow + dNTPs to generate blunt ends.

Generating DB-ORF bait clones

Full-length clones were amplified from cDNA with KOD polymerase (Novagen), using primers with AscI and NotI extension tails. PCR products were gel purified, digested with AscI or NotI, and ligated into linearized pMB28. Clones were verified using internal gene specific primers and primers DB and TERM. Fragment clones were designed based on the size of the gene and the protein domain architecture as predicted by Pfam ([http://](http://pfam.sanger.ac.uk/) pfam.sanger.ac.uk/) and SMART (<http://smart.embl-heidelberg.de/>). Fragments were PCR amplified using full-length sequence verified clones as template. For 7 genes for which we were unable to obtain a full-length clone, cDNA was used as template. For each DBfragment, six clones with a correct insert size were pooled before transformation into yeast.

Yeast two hybrid analysis

DB-ORF (bait) clones were transformed into *S. cerevisiae* strains Y8930 (MAT α ⁴⁹ using the Te-LiAc method⁵⁰. Bait strains able to activate reporter genes in the absence of ADplasmid were discarded as auto-activators. Yeast two-hybrid screens were done as previously described⁹. Two Y2H AD libraries were screened: a library carrying fragments of 749 genes required for early embryogenesis⁹, and a mixed-stage cDNA library (a gift from X. Xin and C. Boone, University of Toronto). To generate the AD mating libraries, yeast strain Y8800 (MAT**a**) was transformed with 30 μg of each library. After three days of growth at 30°C, colonies were harvested in YEPD + 20% glycerol, and frozen in 1 ml aliquots. Counterselection on plates containing cycloheximide was used to eliminate auto-activators that arose during the screening process.

Identification of prey protein identity

Gal4-AD vector inserts were PCR amplified from yeast using primers AD and TERM as described⁹ , and DNA sequencing was performed by Macrogen Europe ([http://](http://dna.macrogen.com) α dna.macrogen.com). Sequences were analyzed by phred⁵¹, using the default settings and the –trim_alt flag. Bases with quality scores below the default threshold of 0.5 were eliminated. Vector ends were clipped from the traces. Sequences where no 5′ vector end could be identified were eliminated. The identity of the ORF was determined by DNA BLAST analysis against a database containing all protein coding genes in Wormbase release WS235. To determine whether the ORF was in frame with the Gal4::AD sequence, we determined the identity of the ORF by protein BLAST using the translated protein sequence. A trace was considered in frame if the DNA and protein BLAST identities match, or when no stop codon was encountered in the first 100 codons. Out of frame sequences were eliminated.

Verification of interactions

Protein pairs identified a single time from the AD-cDNA library were retested by isolating the Gal4-AD plasmid from the original yeast clone, transforming the isolated plasmid into

yeast strain Y8800, and mating with the corresponding Y8930 bait strain. HIS3 reporter activity was assayed on –Leu –Trp –His plates, and protein pairs that failed to activate the reporter were not included in the dataset. For interactions identified multiple times, we confirmed the bait protein identity by PCR amplification and sequencing of the bait construct from a representative yeast colony.

PubMed searches

PubMed searches for interactions between C. elegans proteins were guided by automated searches of the Textpresso database ([http://www.textpresso.org/nematode/\)](http://www.textpresso.org/nematode/) for the cooccurrence within the same publication of two interacting proteins. Publications were then read to determine if an interaction was described. We also manually searched PubMed for interactions between homologous proteins in other organisms.

Comparison with IntAct

To identify interactions present in IntAct ([https://www.ebi.ac.uk/intact/\)](https://www.ebi.ac.uk/intact/), we queried IntAct release 27. To identify orthologous interactions, we compiled a list of potential homologs from Homologene build 67 (<http://www.ncbi.nlm.nih.gov/homologene>), Inparanoid release 7 [\(http://inparanoid.sbc.su.se/cgi-bin/index.cgi\)](http://inparanoid.sbc.su.se/cgi-bin/index.cgi), Orthomcl version 5 ([http://orthomcl.org/](http://orthomcl.org/orthomcl/) [orthomcl/\)](http://orthomcl.org/orthomcl/), and Ensembl release 73 ([http://www.ensembl.org/index.html\)](http://www.ensembl.org/index.html).

Control networks used for computational analysis

To determine whether the cePIN was enriched in similar GO terms, similar expression profiles, or presence in WormNet, we compared the interaction network with 100,000 control networks. We screened two very different AD libraries with correspondingly different search spaces: a genome-wide cDNA library and the early embryogenesis AD-Fragment library, which contains only 749 proteins that already have very similar expression profiles and gene ontology descriptions. AD-cDNA and early embryogenesis AD-Fragment derived interactions were therefore analyzed separately. The control networks were generated by replacing the prey names of the interaction network with protein names randomly selected from the proteins present in the Gal4-AD library screened. This maintains the topology of the original network. Homodimers were excluded from all analyses.

GO term analysis

We used the Hybrid Relative Specificity Similarity based on Gene Ontology (HRSS) software²⁹ in combination with GO database release 201310 (<http://www.geneontology.org/>) to calculate GO semantic similarity scores, limiting our analysis to the Biological Process domain. We compared the distribution of similarity scores for interacting protein pairs with the average distribution of scores of protein pairs in the corresponding control networks. Statistical significance is the fraction of the 100,000 control networks that displayed the same fraction of pairs with a particular similarity score as the actual interaction network.

Comparison with WormNet

The WormNet v3 dataset¹² was queried for the presence of gene pairs, and to extract the loglikelihood score (LLS) of a true functional linkage. Enrichment values are the fraction of

interactions identified by Y2H that are present in WormNet divided by the average fraction of the control network pairs present in WormNet. Statistical significance was calculated as the fraction of the 100,000 control networks that displayed the same or higher fraction of pairs present in WormNet as the actual interaction network.

Gene expression profiling comparison

We calculated pair-wise Pearson Correlation Coefficients between all gene pairs across all C. elegans microarray data collected in Wormbase release WS236. Data were downloaded from the SPELL website (<http://spell.caltech.edu:3000>). For each dataset, we generated normalized Fisher z transformed pair-wise correlation scores as described 22 . We excluded datasets with fewer than 5 different conditions, as well as datasets where >10% of genes had exactly the same expression levels in all conditions. The average of all z correlations was calculated, and converted back to PCC values. Gene pairs which had data in less than 50% of the datasets were excluded. The final compendium contains data for 17,287 proteincoding genes compiled from 110 datasets and 2394 conditions. All calculations and plotting of PCC value distributions were done in the R language⁵³. Statistical significance was calculated as the fraction of the 100,000 control networks that had the same or higher average PCC value as the actual interaction network.

Co-affinity purification

Co-affinity purifications were done as described¹⁵. As bait, the protein fragment which most frequently detected the interaction by Y2H was used. To clone ORFs into vectors AvimCherry-C1 and pEGFP-C1, sequences were PCR amplified from a cDNA using Hot Start KOD polymerase (Novagen) and primers with appropriate restriction enzyme tails. PCR products were gel-purified, digested, and cloned into linearized vectors. Enzyme combinations used were BglII+Acc65I, XhoI+HindIII, SalI+Acc65I, SalI+BamHI and SmaI +XmaI. HEK-293 cells were originally obtained from the ATCC ([http://www.lgcstandards](http://www.lgcstandards-atcc.org/)[atcc.org/\)](http://www.lgcstandards-atcc.org/). Their identity was validated by light microscopic observation. Cells were regularly tested for mycoplasma infection.

MAPPIT

Experiments were performed in HEK-293T cells, which were authenticated in Lin *et al.*⁵⁴, and regularly tested for Mycoplasma infection. We generated Gateway entry clones of the bait and prey ORFs by PCR amplification and BP cloning into vector pDONR223. For both bait and prey constructs, we cloned the fragment that was found most frequently in the Y2H screens. The entry clones were transferred into the two MAPPIT vectors, and MAPPIT assays were performed as described \degree . The fraction of positives was scored over an experiment to control ratio (ECR) threshold range of 1 through 19.

Defining Minimal regions of interaction

Minimal regions of interaction (MRIs) for each interaction were defined as the region of a protein that is present in all of the fragments found to interact. When multiple splice variants were predicted for a gene, the splice variant we identified most frequently was used. In cases where multiple fragments of a protein without any overlap were found to interact, no MRI was defined.

C. elegans strains and culture conditions

C. elegans strains were cultured under standard conditions⁵⁵. Only hermaphrodites were used. A list of strains is available as Supplementary Table 8.

RNAi analysis

RNAi clones were derived from the Vidal RNAi library⁵⁶, the Ahringer RNAi library⁵⁷, or generated in house by cloning of a PCR product amplified from cDNA into a modified L4440 RNAi feeding vector containing AscI and NotI restriction sites (generated by inserting linker L4440_link into BglII and KpnI digested L4440). A list of RNAi clones is available in Supplementary Table 9. All RNAi clones corresponding to bait proteins were sequence verified using primers L4440_F and L4440_R. All RNAi experiments were performed at 20°C, except for experiments with strain JH2647, which was maintained at 25°C to increase mCherry::PAR-6 expression levels. Bacteria were cultured overnight at 37° C in Lysogeny Broth (LB) supplemented with 100 μg/ml ampicilin (Amp) and 2.5 μg/ml tetracyclin (Tet). 1 h before harvesting, 1 mM Isopropyl β-D-1-thiogalactopyranoside (IPTG) was added. Bacterial cultures were concentrated 5x by centrifugation. Nematode Growth Media (NGM) agar plates supplemented with $100 \mu g/ml$ Amp, $2.5 \mu g/ml$ Tet, and 1 mM IPTG were seeded with 250 μl of bacterial suspension, and kept at room temperature (RT) for 48 h before use.

L4 hermaphrodites were placed on RNAi plates, and phenotypes were analyzed in L4 stage F1 progeny, with exception of the mCherry::PAR-6 marker line, which was screened at the embryonic stage, and the yolk trafficking strain, which was screened at the young adult stage. Larval phenotypes for genes that caused an embryonic lethal phenotype were examined by starting the RNAi at L1 and examining the L4 or adult stages of the same generation. Yolk protein trafficking phenotypes for genes that caused severe germline defects were examined by starting the feeding procedure in the L4 stage and analyzing adults of the same generation.

Scoring of all RNAi experiments was done blind, with the observer not knowing the gene being analyzed. In all experiments a positive RNAi control was included: par-⁶ and par-³ for early embryonic polarity, pop-1 for the seam cell epithelium, par-5 for the intestinal epithelium, erm-1 for the excretory canal, ajm-1 for epithelial junctions, chc-1 and cdc-42 for yolk protein trafficking, unc-33 for axon-dendrite specification, and pop-¹ for PVD neuron morphology. In the prey interactor RNAi screens, RNAi targeting the bait was always included, to compare the bait and prey RNAi phenotype and to confirm the originally observed bait defects. In all experiments, empty L4440 vector was used as a negative control. For each RNAi clone, 30–50 animals of the developmental stage indicated above were screened at 10x magnification, and defects were characterized in detail and scored at 63x or 100x magnification. Severity of defects was scored on a scale of $0 - 5$ based on the number of worms affected and the strength of the effect in comparison to the positive control.

Fluorescence microscopy and live imaging

For live imaging of L3 and young adult hermaphrodites, animals were sedated with 10 mM Levamisole M9 and mounted on 5% agarose. To perform imaging on embryos, gravid adults were dissected in M9 and mounted on 5% agarose. Two main microscopes were used. First, a wide-field Zeis Axioplan2 upright microscope equipped with 25x – 0.8 NA, 63x – 1.4 NA, and 100x – 1.4 NA objectives, and an Axiocam MRm CCD monochrome camera. Zeiss filter sets used were set 34 for DAPI, set 13 for GFP, and set 31 for mCherry. The microscope and camera were controlled by Zeiss Axiovision 4.x software. Second, an Andor spinning disc platform controlled by MetaMorph software and consisting of a Nikon Ti-U inverted microscope with 60x – 1.4 NA and 100x – 1.4 NA oil objectives, a Yokogawa CSU-X1 spinning disk unit, 488 nm and 561 nm lasers, Semrock 512 – 23 + 630 – 91, 525 – 30, and 617 – 73 emission filters, and an Andor iXON DU-885 camera. Images in Figure 6 were captured using a Zeiss AxioImager, $40x - 1.3$ NA objective, and Hammamatsu Orca-R2 camera. Images of fixed embryos were deconvolved using AxioVision software, and are shown as maximum intensity projections of 3–5 adjacent planes spaced 0.3 μm apart. Images were cropped, rotated, and levels were adjusted in ImageJ and Adobe Photoshop.

Phenotype clustering

RNAi phenotypes were manually scored on a scale of 0 (no abnormal phenotype) to 5 (severely abnormal phenotype). For each gene, a binary phenotypic profile was generated consisting of the highest severity score for each of the 6 tissues analyzed, adding defects in localization of junctional DLG-1 as a 7th data point. Clustering was done using the Cluster 3.0 software [\(http://bonsai.hgc.jp/~mdehoon/software/cluster/software.htm\)](http://bonsai.hgc.jp/~mdehoon/software/cluster/software.htm), using Spearman rank correlation distance measure and the complete linkage hierarchical clustering method. Clustering results were visualized using Java Treeview [\(http://sourceforge.net/](http://sourceforge.net/projects/jtreeview/) [projects/jtreeview/\)](http://sourceforge.net/projects/jtreeview/).

Correlation of phenotype overlap with interaction, GO, and PubMed

We examined all possible pairs of proteins in the network where the effect of inactivation had been determined in at least 1 overlapping tissue. Excluding homodimers, we evaluated 332 interacting proteins pairs, and 8373 pairs for which we did not detect a physical interaction. A phenotype was considered to overlap if both genes showed a detectable defect in at least 1 overlapping tissue. To examine the correlation between phenotype overlap and physical interaction, we compared the fraction of pairs with a phenotypic overlap between the 332 interacting protein pairs and the 8373 non-interacting pairs. A Fisher's exact test was used to calculate the probability of the observed enrichment. To examine the correlation with GO terms, we used the HRSS scores described above. We defined a high similarity score as an HRSS score > 0.9 , and a low similarity score as an HRSS score $\quad 0.1$. A Fisher's exact test was used to calculate the probability of observed enrichments. To examine the correlation between phenotype overlap and prior characterization of an interaction in a publication, we examined the 19 PubMed papers listed in Supplementary Table 3, excluding homodimers.

Yeast two-hybrid analysis of PAC-1 – PAR-6

Fragments indicated in Figure 6c were cloned into the Y2H vectors pMB28 and pMB29, and transformed into yeast strains Y8930 and Y8800 respectively. Yeast pairs were then mated on YEPD plates, transferred to –Leu –Trp plates for selection of mated yeast, and transferred to –Leu –Trp –His plates to assay the interaction.

Generation of pac-1 transgenic lines

A PAC-1 Gateway clone lacking amino acids 1221-1328 was produced by PCRmodification of a full-length *pac-1* cDNA entry clone⁴⁵, then recombined into destination vector pID3.01B⁵⁸ to create fusions with *GFP*. Ppie-1::GFP::pac-1(PBD) was integrated into *unc-119(ed3)* worms using biolistic transformation⁵⁹.

Antibody staining of embryos

Embryos were freeze-fractured, fixed in methanol, and stained as described previously 45 . The following primary antibodies were used: rabbit α -PAR-6 1:20,000^(ref. 60), rat α -GFP 1:1,000 (Nacalai Tesque GF090R).

Quantifying polarity index

Polarity index was quantified in 8–12 cell stage embryos. Polarity index measurements were obtained by determining the ratio of the average intensity of a forty-pixel line along the contact-free surface of an AB lineage cell versus half of the average intensity of a forty-pixel line at that cell's contact with a neighboring AB cell. ImageJ was used for intensity measurements.

Quantifying expression levels of GFP::PAC-1 and GFP::PAC-1(PBD)

Total fluorescence values were obtained in a central focal plane of 4-cell embryos expressing each GFP fusion. Using equivalent camera exposures, average background fluorescence (wild-type 4-cell embryos) was subtracted from the average fluorescence value for each strain. Signals were integrated over the area of the embryo.

Co-localization in HeLa cells

Avi-mCherry-C1and pEGFP-C1 constructs generated for co-affinity purification were transfected in HeLa Kyoto cells grown in DMEM-Ham's F10 (50/50%) medium containing 10% FCS and 1% penicillin + streptomycin at 37° C and 5% CO₂. Two days before transfection, nearly confluent cells were plated at 1:10 in 12-well plates on 24 mm glass coverslips. Cells were transfected with 1 μg of Avi-mCherry-bait and 1 μg of pEGFP-C1 prey with polyethyleminine (PEI). After overnight incubation, cells were washed with PBS, fixed for 10 min with 4% paraformaldehyde in PBS at room temperature, washed with PBS and subsequently mounted on slides in Vectashield mounting medium with DAPI. The original source of the HeLa Kyoto cells is not known and cells were not authenticated.

Statistical analysis

Statistical tests used and sample sizes are indicated in the figure legends and in the methods section. No statistical method was used to predetermine sample sizes. No samples or animals

were excluded from analysis. The experiments were not randomized, and the investigators were not blinded to allocation during experiments and outcome assessment, with the exception of the RNAi screens, where investigators were unaware of the gene being targeted during scoring. Statistics regarding the enrichment of protein pairs in the interaction network sharing certain characteristics were based on comparisons with random control networks, with p-values corresponding to the number of times a value as extreme as the actual network was found in 100,000 control networks. This statistical approach was chosen to reduce bias by factors such as the connectivity degree of the proteins in the network. A two-sided Fisher's exact test was used to evaluate the significance of a higher fraction of interacting pairs retesting positively by MAPPIT than control random pairs. The test is appropriate to examine whether there is a significant association between the retest result by MAPPIT and prior identification as an interacting protein pair by an interaction assay. A Mann-Whitney U test was used to determine the significance in differences in polarity index and fluorescence intensity levels between embryos of different genotypes, as a normal distribution of the data could not be confidently determined from the sample values (Anderson-Darling normality test).

Reproducibility of experiments

Co-affinity purifications in Figure 2d and Supplementary Figure 2 were performed once. Localization of fluorescent marker proteins in wild-type background (Figure 4a–i) was observed in >30 animals. All RNAi phenotypes (Figure 5 and Supplementary Table 6) were observed in >5 animals. Yeast two-hybrid analyses in Figure 6c were performed 3 times. Coaffinity purifications in Figure 6d were performed twice. PAC-1 and PAR-6 localization patterns in HeLa cells (Figure 6e–g) were observed in > 30 cells. Embryo staining patterns in Figure 6h–k were observed 58/58 (h), 62/62 (i), 50/50 (k) and 61/61 (k) embryos.

Re-analysis of published datasets

In Figure 2e, we re-analyzed the MAPPIT results from $3 \, C$. elegans protein interaction datasets previously analyzed by $MAPPIT¹⁸$.

Code availability

All code that was generated in-house is available upon request. Code may not be further distributed or used for purposes other than to re-analyze the data presented here without prior permission from the authors.

Data availability

The protein interactions from this publication have been submitted to the IMEx ([http://](http://www.imexconsortium.org) www.imexconsortium.org) consortium through the IntAct database⁶¹ and assigned the identifier IM-24524. A searchable web interface of all interactions and fragments identified is available at<http://www.projects.science.uu.nl/interactome/>

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.

Identification and validation of a C. elegans polarity interaction network (CePIN) (**a**) Schematic representation of the different classes of protein selected as baits. Numbers indicate number of proteins in each class. See Supplementary Table 1 for details. (**b**) Example of the design of bait fragments. Fragments were manually designed based on the size of the gene and the predicted protein domain composition. Protein domains are indicated. Grey lines represent the fragments cloned. Red and blue arrows represent forward and reverse amplification primers, respectively. (**c**) Overview of the yeast two-hybrid screening and computational validation pipeline used to establish the C. elegans polarity interaction network. (**d**) Network graph of the identified protein interactions. Colored nodes are bait proteins, and the color indicates the classification in panel (a). Grey nodes are prey proteins. Black edges indicate interactions with a phenotype overlap (see below).

Figure 2.

Validation of the CePIN (**a,b**) Enrichment of similar GO terms (a) and presence in WormNet (b) for interacting protein pairs compared to control networks generated by replacing prey proteins with random proteins from the search space. Interactions from AD-cDNA and AD-Fragment libraries were plotted separately as the search space was highly different. Control network bars represent the mean of 100,000 control networks \pm s.d. Statistical significance is the fraction of control networks that displayed the same or higher fraction of pairs with a high GO similarity or pairs present in WormNet as the actual interaction network. (**c**) Average Pearson correlation coefficient (PCC) score of mRNAs corresponding to protein pairs in the interaction dataset (red arrow), compared with the distribution of average PCC scores of 100,000 control networks (blue line). Only AD-cDNA derived pairs were analyzed, as proteins pairs in the AD-Fragment library already have very similar expression profiles (Supplementary Fig. 1d). Statistical significance is the fraction of control networks that displayed an average PCC score identical or higher than the actual interaction network. (**d**) Experimental validation by co-affinity purification. Bait proteins were expressed as AvimCherry fusions together with the bacterial biotin ligase BirA, which biotinylates the Avi sequence. Prey proteins were expressed as EGFP fusions. Bait proteins were purified from cell lysates using streptavidin-coated beads. Protein pairs tested are indicated above the blots. For prey proteins, f.l. indicates full length protein, and frag. a shorter fragment. Lanes 1 and 2 are controls for non-specific binding of bait to EGFP or prey to Avi-mCherry. The \sim 35 kDa band in lanes 2 is due to cross-reactivity of the anti-GFP antibody with AvimCherry. Asterisks indicate bands of expected molecular mass. Expression of tagged protein in input lysates is shown in Supplementary Fig. 2. Purifications were performed once. Unprocessed scans are shown in Supplementary Figure 5. (**e**) Fraction of interactions from indicated datasets that test positive by MAPPIT at increasing assay stringency. CePRS, CeRRS, and WI-2007 data are reproduced from Simonis et al.¹⁸. Shading indicates standard

error of the proportion. At a scoring threshold of 12, p < 1×10−5 for each interaction dataset compared to CeRRS, two-sided Fisher's exact test.

Figure 3.

Identification and validation of minimal regions of interaction (**a**) Distribution of the size of the identified MRIs as a percentage of the full-length protein. Interactions identified only as full-length are indicated separately (orange bar) (**b**) Distribution of the length of the identified MRIs in absolute residues. See Supplementary Figure 6a–f for separate size distributions of MRIs identified from bait proteins, from AD-Fragment clones, and from AD-cDNA clones. (**c**) Examples of MRIs for interactions where the binding site had already been described in the literature. See Supplementary Figure 6g for the complete set of MRIs for interactions with known interactions sites.

Figure 4.

Phenotypic analysis of the bait proteins (**a**–**i**) Marker lines used for phenotypic profiling. All scale bars represent 10 μm. (**a**) Intestinal expression of PEPT-1::dsRed (apical domain) and GFP::RAB-11 (apical recycling endosomes)²⁵. (**b**) Seam epithelium with cell outline marked by GFP fused to a pleckstrin homology domain, and nuclei marked by a Histone 2B GFP fusion. (**c**) Expression of the junctional protein DLG-1::GFP (shown here in intestine). (**d**) Uptake of the yolk protein VIT-2::GFP in oocytes²⁶. (**e**) Cortical localization of RME-2::GFP, the receptor for VIT-2, in oocytes. (**f**) Asymmetric distribution of mCherry::PAR-6 at the two-cell stage. (**g**) Localization of the presynaptic molecules mCherry::RAB-3 and SAD-1::GFP in the PVD axon (arrow). SAD-1::GFP is also observed at low levels in the cell body and dendrites near the cell body. (**h**) Expression of myristoylated GFP under control of a PVD-specific promoter, to examine the extensive arborisation pattern of the PVD dendritic tree. (**i**) Excretory canal outline by expression of VHA-5::GFP. Shown is the head region where the excretory canal cell body is located. (**j**) Hierarchical clustering of bait proteins based on observed phenotypes. Green colour indicates presence of a defect, and colour intensity indicates severity of the defect on a scale of 1–5. Phenotypes scores were based on examination of >15 animals.

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Figure 5.

Examples of RNAi phenotypes. In all panels, the gene inactivated and the marker protein shown are indicated. EV: empty vector control RNAi. (**a**) Examples of interacting protein pairs where RNAi for both corresponding genes resulted in a defect in the same marker strain. In panels 2–3, arrowheads point to spreading of SAD-1::GFP into dendrites. In panels 4–6 arrowheads point to spreading of mCherry::RAB-3 into dendrites or cell body. In panels 7–9, arrowheads indicate defects that arise during the development of the PVD neuron. Arrow in panel 11 indicates failed cell division. Arrow in panel 12 points to larger than normal distance between cells and elongated connection between cells. In the yolk

trafficking panels, arrows point to accumulation of yolk in the body cavity. In the excretory canal panels, arrowheads point to defects in canal morphology. (**b**) Examples of phenotypes observed in the PVD neuron, the Excretory canal, and the intestine during phenotypic analysis of the bait proteins. In PVD neuron panels, an asterisk indicates the location of the cell body, and arrowheads indicate spreading of RAB-3 into dendrites. Images are representative of phenotypes observed in >5 animals. Scale bars are 10 μm.

Figure 6.

The PAR-6 – PAC-1 interaction is required for radial polarization (**a–c**) Delineation of the protein regions that mediate the interaction between PAR-6 and PAC-1. (**a,b**) Schematic representations of protein fragments (grey lines) tested for interaction by Y2H. Presence or absence of interaction is indicated to the left of each fragment. Yellow regions represent the minimal regions of interaction. (**c**) Example of Y2H analysis using the fragments indicated. Left: permissive plate containing histidine, right: selective plate lacking histidine. Representative image of 3 experiments. (**d**) Validation of the PAC-1 – PAR-6 interaction and interaction domains by co-purification. Western blots show co-purification of GFP::PAR-6 with biotinylated mCherry::PAC-1. Protein fragments tested are indicated. See legend of Fig. 2d for details. Co-purifications were performed at least twice. Unprocessed scans are shown in Supplementary Figure 5. (**e–g**) Subcellular localization of mCherry::PAC-1 (aa 1221 – 1604) and EGFP::PAR-6 (aa 133 – 155) in HeLa cells. Insets are magnified 2x. Localization patterns were observed in >30 cells. (**h,i**) PAR-6 is enriched at contact-free surfaces in wildtype embryos (n=58/58) and localizes symmetrically to cell surfaces in pac-¹ mutant embryos (n=62/62). (**j, j**′) PAR-6 is enriched at contact-free surfaces in pac-1; GFP::pac-1 embryos (n=50/50). (**j**′) GFP::PAC-1 localizes to sites of cell contact. (n=50/50) (**k**) PAR-6 localizes symmetrically to cell surfaces in pac-1; GFP::pac-1(PBD) embryos (n=61/61). (**k** $'$) GFP::PAC-1(PBD) localizes to sites of cell contact, like GFP::PAC-1. $(n=61/61)$ (**l**) Quantification of PAR-6 asymmetry in embryos of the indicated genotypes (see Methods for

details). Circles represent individual embryos, and the bar indicates mean value. 8 embryos were measured for each genotype. P-values were calculated using the Mann-Whitney U test. n.s. = not significantly different, $** = p < 0.01$. Samples were pooled from 3 independent experiments. (**j**) Expression levels of GFP::PAC-1 and GFP::PAC-1(PBD), measured as fluorescence intensity at the four-cell stage. Statistical analysis was performed using the Mann-Whitney U test, $n = 7$ GFP::PAC-1 embryos and $n = 9$ GFP::PAC-1(PBD) embryos. Box represents first and third quartiles, bars represent maximum and minimum values, and the line within the box is the mean fluorescence intensity. Samples were pooled from 2 independent experiments. All scale bars are 10 μm.