RhoA activation during polarization and cytokinesis of the early Caenorhabditis elegans embryo is differentially dependent on NOP-1 and CYK-4

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ABSTRACT The GTPase RhoA is a central regulator of cellular contractility in a wide variety of biological processes. During these events, RhoA is activated by guanine nucleotide exchange factors (GEFs). These molecules are highly regulated to ensure that RhoA activation occurs at the proper time and place. During cytokinesis, RhoA is activated by the RhoGEF ECT-2. In human cells, ECT-2 activity requires its association with CYK-4, which is a component of the centralspindlin complex. In contrast, in early Caenorhabditis elegans embryos, not all ECT-2–dependent functions require CYK-4. In this study, we identify a novel protein, NOP-1, that functions in parallel with CYK-4 to promote RhoA activation. We use mutations in nop-1 and cyk-4 to dissect cytokinesis and cell polarization. NOP-1 makes a significant, albeit large-ly redundant, contribution to cytokinesis. In contrast, NOP-1 is required for the preponderance of RhoA activation during the establishment phase of polarization.

INTRODUCTION The small GTPase RhoA is a central regulator of cellular contractility. When activated by exchanging its bound GDP cofactor for GTP, this molecular switch activates formins and Rho kinase, which stimulates polymerization of unbranched actin filaments and promotes assembly and activation of minifilaments of nonmuscle myosin II, respectively (for a review, see Pike et al., 2005). RhoA and its effectors constitute a contractile module that impacts diverse biological processes. Two such processes, cell polarization and cytokinesis, have been extensively analyzed in the early Caenorhabditis elegans embryo (Munro and Bowerman, 2009). These processes share a number of molecular, genetic, and cytological similarities. Notably, cortical contractility, which is critical for both processes, is dependent on RhoA activation by the guanine nucleotide exchange factor (GEF) ECT-2 (Dechant and Glotzer, 2003; Zonies et al., 2010).

Although cytokinesis and cell polarization share requirements for ECT-2, RhoA, and their downstream effectors, the mechanisms that trigger ECT-2 activation during these processes appear to be molecularly distinct. During cytokinesis, but not polarization, ECT-2 activation involves extensive interactions with CYK-4, a component of the centralspindlin complex that contains a Rho-family GTPase-activating protein (GAP) domain. The N-terminus of ECT-2 contains tandem BRCA1 C-Terminal (BRCT) domains that bind to phosphorylated CYK-4 and contribute to its activation (Burkard et al., 2009; Wolfe et al., 2009). The BRCT domains also bind to and inhibit the catalytic C-terminus (Kim et al., 2005), suggesting that CYK-4 binding promotes relief from auto-inhibition. The integrity of the GAP domain of CYK-4—although
not necessarily its catalytic activity—is also required for full activation of RhoA during cytokinesis in a variety of experimental systems (Yamada et al., 2006; Canman et al., 2008; Loria et al., 2012). However, loss-of-function mutations in CYK-4 that cripple its ability to promote RhoA activation in cytokinesis appear to have little effect on RhoA activation during embryo polarization (Canman et al., 2008; Loria et al., 2012).

Polarization of the early C. elegans embryo involves an initial establishment phase that is followed by a mechanistically distinct maintenance phase (Cuenca et al., 2003). The establishment phase is initiated by fertilization, which activates global cortical contractility. The sperm-provided centrosome and its associated astral microtubules locally suppress contractility near the posterior pole (Cowan and Hyman, 2004; Tsai and Ahringer, 2007). The resulting nonuniform contractility induces anterior-directed cortical flow and concentration of numerous proteins, including myosin and the polarity factors PAR-3, PAR-6, and PKC-3, to the anterior of the embryo (Munro et al., 2004). This flow is accompanied by a transient, cytokinetic-like furrow called the pseudocleavage furrow. Concomitant with this flow, posterior PAR proteins, such as PAR-1 and PAR-2, are recruited to the posterior cortex and prevent the return of the anterior PARs to the posterior cortex (Cuenca et al., 2003; Munro et al., 2004). A second, microtubule-dependent pathway can direct cortical PAR-2 recruitment when cortical flows are eliminated (Motegi et al., 2011). During the maintenance phase, anteriorly localized CDC-42 promotes polarized activation of myosin via myotonic dystrophy kinase–related Cdc42-binding kinase and activation of ARP2/3-dependent actin polymerization (Gotta et al., 2001; Kay and Hunter, 2001; Kumfer et al., 2010). Symmetry-breaking polarization is robust; perturbations that result in clear molecular phenotypes may be tolerated, but combinations of these perturbations can block polarization (Zonies et al., 2010).

Cleavage furrow ingression in early C. elegans embryos is mediated by two distinct pathways that can be distinguished by their dependence on the central spindle (Dechant and Glotzer, 2003; Bringmann and Hyman, 2005). Central spindle–independent, also known as aster-directed, furrowings resemble the pseudocleavage furrow in that both invariably undergo furrow regression. Additionally, a number of perturbations have been described that preferentially impair central spindle–independent furrowing; these perturbations also prevent pseudocleavage furrow formation (Werner and Glotzer, 2008). One such case is that of embryos depleted of anillin (ANI-1). ANI-1 promotes the organization of cortical myosin into large foci and its polarization along the anterior–posterior axis (Maddox et al., 2005; Tse et al., 2011). Interestingly, whereas cleavage furrows ingress asymmetrically in control embryos, they ingress symmetrically in embryos depleted of either ANI-1 or the septins (Maddox et al., 2007). No other factors are known to regulate the asymmetry of furrow ingression.

A maternal effect mutation, nop-1(it142), has been described that abrogates cortical contractility during the establishment phase of polarization without dramatically affecting cytokinesis (Rose et al., 1995). In fact, homozygous nop-1 mutant animals are viable and fertile, although ∼20% of these embryos fail to hatch. nop-1 mutant embryos also fail to form an anterior actin cap, indicating a defect in organization or function of the cytoskeleton. However, the molecular nature of NOP-1 has not been established. We report the molecular identification of NOP-1 and show that this protein promotes RhoA-dependent processes during the establishment phase of polarization and during cytokinesis. NOP-1 is required for the majority of RhoA activation during polarization, whereas it acts in parallel to the central spindle–dependent pathway for RhoA activation during cytokinesis.

**RESULTS**

**Molecular identification of NOP-1**

nop-1 was originally mapped to a position on chromosome III between dpy-17 and unc-32 (Rose et al., 1995). Further mapping experiments revealed that the mutation is located in the 0.01 cM region between sma-4 and cyc-3 on chromosome III (Figure 1A). To identify nop-1, we depleted all the genes in this region by RNA interference (RNAi) and filmed early embryos using Nomarski microscopy. RNAi depletion of one gene, F25B5.2, blocked formation of the pseudocleavage furrow and phenocopied the nop-1(it142) mutation (Figure 1B, Figure 1 Movie S1). Furthermore, in nop-1(it142), we identified a premature stop codon near the end of the predicted open reading frame, at amino acid 696 (Figure 1C). Embryonic lethality occurs in similar fractions of nop-1(it142), F25B5.2(RNAi), and nop-1(it142);F25B5.2(RNAi) embryos (∼27%). These results, together with the rescue data below, indicate that nop-1 is encoded by the F25B5.2 gene and that nop-1(it142) is a strong loss-of-function mutation, perhaps even a null.

NOP-1 is a 757-residue, serine-rich (16.8%) protein lacking recognizable domains or motifs. Orthologues of NOP-1 can be readily identified in the related nematodes Caenorhabditis briggsae, Caenorhabditis japonicus, Caenorhabditis brenneri, and Caenorhabditis remanei (Supplemental Figure S1). However, despite extensive efforts, we were unable to identify orthologues in other species.

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**FIGURE 1:** Identification of NOP-1. (A) nop-1 maps to −1.43 on chromosome III, between sma-4 and cyc-3. (B) Images from a time-lapse sequence of the first division of wild-type, nop-1(it142), and nop-1(RNAi) embryos at the indicated cell cycle phases. Wild-type embryos form pseudocleavage furrows (black arrows), whereas pseudocleavage is absent in nop-1(it142) and nop-1(RNAi) embryos. Scale bar: 10 μm. (C) Schematic depicting the domain structure of NOP-1. The dotted line indicates the premature stop codon found in nop-1(it142).
Localization of NOP-1
To determine the subcellular localization of NOP-1 in C. elegans embryos, we inserted a green fluorescent protein (GFP)-tagged NOP-1 transgene controlled by the germ line–active PIE-1 regulatory elements into the C. elegans genome using MosSCI (Freijka-Jensen et al., 2008). GFP::NOP-1 does not have a strong dominant effect on embryonic viability or progression of cytokinesis, although the brood size is slightly reduced (Figure 2, A and B). GFP::NOP-1 partially rescued both the pseudocleavage defect and the lethality resulting from the nop-1(it142) mutation (Figure 2, A and B, Figure 2 Movie S1), confirming the gene identification. We imaged embryos expressing GFP::NOP-1 from pseudocleavage to completion of cytokinesis using time-lapse confocal microscopy. GFP::NOP-1 dramatically concentrates in interphase nuclei. In addition to a significant cytoplasmic pool, a distinct cortical localization can be detected, particularly at ingressing pseudocleavage and cleavage furrows (Figure 2C, Figure 2 Movie S2). Similar results were obtained when the localization of endogenous NOP-1 was assessed with an antibody raised against a peptide from the C-terminus of NOP-1 (Supplemental Figure S2).

NOP-1 promotes cortical recruitment of RhoA effectors
To determine how NOP-1 regulates pseudocleavage, we investigated the dynamic localization of mediators of cortical contractility. We recorded time-lapse images of control and nop-1(it142) embryos expressing NMY-2::GFP, GFP::ANI-1 and GFP::Utophin from the time of pronuclear migration to telophase using spinning-disk confocal microscopy (Figure 3). As reported previously, in control embryos, NMY-2::GFP forms large patches in the anterior half of embryos during pseudocleavage (Munro et al., 2004). These patches then dissociate, and smaller punctae accumulate in the anterior region and form an anterior cap of myosin during prometaphase. On anaphase onset, this anterior cap dissipates and NMY-2::GFP reaccumulates, forming large patches primarily at the equatorial region and near the anterior pole (Figure 3A, Figure 3 Movie S1). The pattern of myosin recruitment was dramatically different in nop-1(it142) embryos. Large NMY-2::GFP patches were absent from nop-1(it142) embryos during pseudocleavage (Figure 3A and Supplemental Figure S3). As the pronucelus migrates, small NMY-2::GFP foci are evenly distributed throughout the entire cortex, and cortical flow is greatly attenuated. The anterior cap of smaller myosin punctae forms during prometaphase, although its boundary extends more posteriorly than in control embryos (see following section). This cap then dissipates as in control embryos, and discrete, small NMY-2::GFP foci reaccumulate specifically in the equatorial region and induce furrow ingression (Figure 3A, Figure 3 Movie S1). Thus the large myosin patches that appear during pseudocleavage and anaphase in control embryos are NOP-1–dependent.

RhoA activation also triggers the cortical accumulation of the cytokinetic scaffold protein anillin, which largely colocalizes with myosin in control embryos (Maddox et al., 2005). In anillin-depleted embryos, myosin accumulates in the cortex, but the patches are dramatically smaller. In contrast, cortical anillin accumulates normally in myosin-depleted embryos (Maddox et al., 2005; Tse et al., 2011). Therefore cortical recruitment of anillin and myosin are partially independent markers for RhoA activation. We therefore investigated whether inactivation of NOP-1 also affects anillin. Anillin patches are absent from nop-1(it142) embryos; only small cortical foci are observed during pseudocleavage and in the equatorial region during cytokinesis (Figures 3B and S3, Figure 3 Movie S1). Quantitative analysis reveals that NOP-1 promotes cortical recruitment of both anillin and myosin, as well as the accumulation of equatorial cortical actin (Supplemental Figure S3).

Examination of fixed embryos indicates that nop-1(it142) embryos are defective in cortical actin accumulation (Rose et al., 1995). To gain more insight into this phenotype, we characterized the dynamic pattern of actin in live embryos using GFP::Utophin, a probe derived
from the F-actin–binding calponin homology domain of utrophin (Burkel et al., 2007). In control embryos, cortical filamentous actin and actin puncta form in the anterior half of embryos during pseudocleavage. Like myosin, actin filaments become enriched in the anterior half of the embryo and form an anterior cap during prometaphase. The cap then disperses and actin filaments reappear at the site of cleavage (Figure 3C, Supplemental Figure S3, and Figure 3 Movie S1). The overall dynamic behavior of actin filaments in nop-1(it142) embryos is similar to controls. However, the actin cap during pseudocleavage is attenuated. In addition, accumulation of actin filaments at the equatorial region during cytokinesis is slightly delayed in NOP-1–depleted embryos. Thus NOP-1 promotes the cortical recruitment of anillin and myosin during both pseudocleavage and cytokinesis and promotes the accumulation of actin at furrowing regions.
NOP-1 functions upstream of ECT-2

Contractile behavior during pseudocleavage and cytokinesis in the early embryo is dependent on the RhoGEF ECT-2 (Jenkins et al., 2006; Motegi and Sugimoto, 2006; Schonegg and Hyman, 2006; Werner et al., 2007). The primary RhoGAPs that antagonize ECT-2 are RGA-3/4, two highly related RhoGAPs (Schmutz et al., 2007; Schonegg et al., 2007). Depletion of RGA-3/4 leads to increased cortical contractility during polarization and cytokinesis (Figure 3D).

The observations shown thus far indicate that NOP-1 promotes the accumulation of active RhoA. In principle, NOP-1 could perform this function by inhibiting RGA-3/4. If NOP-1 functioned upstream of RGA-3/4, then co-inactivation of NOP-1 and RGA-3/4 should phenocopy RGA-3/4 depletion. We acquired time-lapse differential interference contrast (DIC) images of control and nop-1(it142) embryos depleted of RGA-3/4. Cortical contractions are enhanced and pseudocleavage persists until the pronuclei centrate in rga-3/4(RNAi) embryos (Figure 3D, Figure 3 Movie S2). In contrast, cortical ruffling and pseudocleavage are absent in nop-1(it142);rga-3/4(RNAi) embryos (Figure 3D, Figure 3 Movie S2), indicating that NOP-1 is epistatic to RGA-3/4 and does not inhibit it. As a second test for residual RhoA activity in early nop-1 embryos, we used another mutant that causes hypercontractility, rfl-1(or198ts). Depletion of NOP-1 in rfl-1(or198ts) embryos also causes a cessation of cortical contractility during pseudocleavage (Supplemental Figure S4).

Further evidence that NOP-1 contributes to cytokinetic furrowing arises from analysis of ect-2(ax751ts);nop-1(RNAi) embryos. In ect-2(ax751ts) embryos, cortical ruffling and pseudocleavage are absent (Figure 3D) yet cytokinesis proceeds to completion in most embryos at the semipermissive temperature (Zonies et al., 2010), resembling the phenotype of nop-1(it142);rga-3/4(RNAi) embryos (Figures 1B and 6A). Whereas both ect-2(ax751ts) and nop-1(RNAi) embryos are able to complete cytokinesis efficiently, doubly compromised embryos do not exhibit cytokinetic furrowing (Figure 3D, Figure 3 Movie S2). This genetic interaction is consistent with a model in which NOP-1 and ECT-2 act in a common pathway or in parallel to activate RhoA.

If NOP-1 and ECT-2 functioned in parallel, contractility would be predicted to persist in embryos in which either protein is strongly depleted. During pseudocleavage, this is not the case, contractility is absent in both single mutants (Figure 3D). Furthermore, during cytokinesis, strong depletion of ECT-2 is sufficient to abrogate cortical contractility. Indeed, cortical accumulation of anillin during anaphase is reduced to near background levels by strongly depleting ECT-2 (Supplemental Figure S5). Depletion of ECT-2 in nop-1(it142) embryos results in a marginally stronger effect, although it is difficult to exclude the possibility that this slight difference depends on residual ECT-2 activity (Supplemental Figure S5). Thus, during cytokinesis, NOP-1 requires ECT-2 to generate contractility, but not vice versa. These data suggest that NOP-1 acts upstream of ECT-2.

NOP-1 mediates aster-induced furrowing

Two independent pathways contribute to furrow formation in the early C. elegans embryo: central spindle–directed furrowing and aster-directed furrowing (Dechant and Glotzer, 2003; Brungmann and Hyman, 2005). The molecular mechanism of aster-induced furrowing during cytokinesis shares numerous similarities with pseudocleavage during polarity establishment. The genetic requirements of the two processes are similar and, in each case, a furrow forms at the border between an anterior region enriched in cortical myosin and a posterior region depleted of this motor protein (Werner et al., 2007; Afshar et al., 2010). We therefore asked whether NOP-1 regulates aster-directed furrowing during cytokinesis.

Depletion of the central spindle subunit ZEN-4 inactivates central spindle-induced furrowing; all the residual furrowing is aster-dependent (Werner et al., 2007). In wild-type and nop-1(it142) embryos, furrows ingress during anaphase and cytokinesis completes in most embryos (Figure 4A). In zen-4(RNAi) embryos, furrows ingress and then regress (Figure 4A). However, cleavage furrow formation is completely blocked in nop-1(it142);zen-4(RNAi) doubly compromised embryos (Figure 4A, Figure 4 Movie S1). To further determine whether NOP-1 is essential for the aster-induced pathway, we spatially separated the furrows that result from these two pathways in the same embryo by depleting the microtubule-associated protein ZYG-9 (Werner et al., 2007). Inactivation of ZYG-9 results in shorter microtubules and the spindle becomes mispositioned at the posterior pole. ZYG-9-depleted embryos form central spindle–dependent and aster-induced furrows in the posterior and the anterior regions of the zygote, respectively (Werner et al., 2007). The formation of anterior furrow is blocked in both nop-1(it142);zyg-9(RNAi) and nop-1(it142);zyg-9(b244ts) embryos (Figure 4B). Interestingly, the posterior furrow is perturbed in a subset of nop-1(it142);zyg-9(RNAi) embryos (40%) and nop-1(it142);zyg-9(b244ts) embryos (25%). NOP-1 is thus critical for aster-directed furrowing and makes a subtle, but detectable, contribution to central spindle–dependent furrowing.

Control of cytokinetic furrowing by independent RhoA activators

Recent work has demonstrated that CYK-4 is required for activating RhoA during central spindle–directed furrowing. An allele of cyk-4, or749ts, contains a substitution mutation in the GAP domain; this is a separation-of-function allele in which central spindle assembly is normal, but there is a dramatic reduction in the cortical accumulation of myosin, F-actin, and anillin and central spindle–directed furrowing is absent (Canman et al., 2008; Loria et al., 2012). The previous experiment suggests that NOP-1 also promotes RhoA activation, although it primarily contributes to aster-directed furrowing. These results suggest that NOP-1 and CYK-4 could be partially redundant activators of ECT-2 that are collectively responsible for ECT-2 activation. This model predicts that nop-1;cyk-4(or749ts) embryos would fail to recruit RhoA effectors upon anaphase onset and would fail to form cleavage furrows altogether. We imaged nop-1(RNAi);cyk-4(or749ts) embryos and found that the doubly compromised embryos indeed fail to form cleavage furrows upon anaphase onset (Figure 4A and Figure 4 Movie S1). In addition, cortical recruitment of myosin and anillin at the equatorial region upon anaphase onset are attenuated in cyk-4(or749ts);nop-1(RNAi) embryos (Figure 4, C and D); F-actin accumulation is also modestly reduced (Figure 4E).

NOP-1 promotes RhoA activation

To confirm that the cortical recruitment of RhoA effectors reliably reflects RhoA activity, we developed a RhoA biosensor. This sensor consists of GFP fused to the C-terminal portion of C. elegans anillin, which contains its conserved region (AH) and pleckstrin homology (PH) domain. It lacks the N-terminal myosin- and actin-binding domains but retains its RhoA-binding domain. This sensor is similar to one used in human cells that colocalizes with the cortical pool of active RhoA (Piekn and Glotzer, 2008). This truncated anillin does not dominantly interfere with anillin function; pseudocleavage occurs normally, and cytokinetic furrowing is asymmetric. This sensor was validated in C. elegans by assaying its cortical abundance in control, ECT-2-depleted and RGA-3/4-depleted embryos. In control embryos, GFP::AHPH is cortically recruited during pseudocleavage and
FIGURE 4: NOP-1 regulates aster-induced cleavage furrows. (A) Selected images from time-lapse sequences of wild-type, nop-1(it142), zen-4(RNAi), nop-1(it142); zen-4(RNAi), cyk-4(or749ts), and cyk-4(or749ts);nop-1(RNAi) embryos at the indicated cell cycle phases. (B) The number of ZYG-9–depleted embryos with anterior and posterior furrows in control and NOP-1–depleted embryos. (C–E) Selected confocal images from time-lapse sequences of cyk-4(or749ts) and cyk-4(or749ts);nop-1(RNAi) embryos expressing NMY-2::GFP (C), GFP::ANI-1 (D), and GFP::Utrophin (E) at the indicated times relative to anaphase onset. Insets (top, right) show the chromatin as detected by mCherry::HIS. Scale bars: 10 μm.

The RhoA biosensor was used to assess whether loss-of-function mutations in NOP-1 and/or CYK-4 result in reduced RhoA activation. During polarization, GFP::AHPH accumulation at the cell cortex is NOP-1–dependent (Figure 5A). During cytokinesis, cortical GFP::AHPH in the furrow region is reduced in both nop-1(it142) embryos and cyk-4(or749ts) embryos (Figure 5, B and C). However, in each case there is residual accumulation of GFP::AHPH. In nop-1(RNAi) embryos, GFP::AHPH accumulation is moderately reduced, whereas in cyk-4(or749ts) embryos, GFP::AHPH is dramatically reduced, although it remains detectable (Figure 5, B and C). In nop-1(RNAi);cyk-4(or749ts) embryos, no cortical GFP::AHPH was observed, confirming that NOP-1 and CYK-4 act in parallel to activate RhoA (Figure 5, B and C, and Supplemental Figure S6A).

cytokinesis, a pattern that resembles that of GFP:anillin (Figure 5, A and B). The cortical abundance of this reporter in the equatorial region declines when ECT-2 is depleted (Supplemental Figure S6, A–C) and increases when RGA-3/4 are depleted (Supplemental Figure S6, B and C). Cortical recruitment of the biosensor does not require downstream effectors of RhoA. Moderate depletion of NMY-2 causes a slight increase in sensor recruitment (Supplemental Figure S6, B and C), stronger depletion results in a more dramatic increase (Supplemental Figure S6A). ANI-1 depletion causes a stronger increase in biosensor recruitment (Supplemental Figure S6, B and C). These results suggest that RhoA may be coupled through feedback regulation to downstream processes. Alternatively, or in addition, there may be competition of the sensor with endogenous ANI-1. These results indicate that GFP::AHPH is a reliable indicator of RhoA activation.
NOP-1 can modulate asymmetric cleavage furrow ingression

Because the cortical accumulation of myosin, anillin, and actin are altered in NOP-1–depleted embryos, we investigated whether these molecular defects affect ingression of the cleavage furrow. Initiation of cleavage furrow ingression is slightly delayed in NOP-1–depleted embryos (Figure 6A), but furrowing completes with kinetics similar to those of control embryos (Figure 6, B and C). Given that both anillin and NOP-1 primarily promote aster-directed furrowing, we examined the behavior of embryos in which both of these factors are inactivated. Both individual perturbations abrogate pseudocleavage furrowing, but neither prevents cytokinesis. Anillin-depleted embryos have an additional phenotype in which furrow formation occurs symmetrically (Maddox et al., 2007; Figure 6, A and B). In contrast, cleavage furrows of nop-1(it142) embryos ingress asymmetrically, similar to control embryos (Figure 6, A and B). Asymmetric furrowing in nop-1(it142) embryos could result from central spindle–directed accumulation of anillin (compare Figures 3B and 4D). Surprisingly, we found that furrow formation is asymmetric in ani-1(RNAi);nop-1(it142) embryos (Figure 6, A and B). The vast majority of these embryos also exhibit hyperaccumulation of PH:GFP at the furrow tips (Figure 6A). These results reveal the existence of an ANI-1–dependent mechanism for generating asymmetrically ingressing cleavage furrows.

NOP-1 regulates polarity establishment

As polarity is established, cortical myosin flows away from the sperm microtubule-organizing center to the opposite pole, where it forms an anterior cap enriched in large myosin foci. This cortical flow of myosin promotes the concurrent flow of PAR-6/PAR-3/PKC-3 and its accumulation at the anterior pole (Munro et al., 2004). In NOP-1–depleted embryos, the cortical flow of myosin is weaker, and the assembly of large myosin patches in the anterior is abolished (Figure 7A). A significant subset of nop-1(it142) embryos lacked cortical flows altogether (11/56); these embryos exhibited a high incidence of symmetric or failed first division (7/11 and 4/11, respectively), and all of these embryos failed to hatch. These results suggest that nop-1 is a Par gene with incomplete penetrance. Consistent with reduced cortical flows, the anterior cap of myosin occupies ∼65% of egg length in nop-1(it142) embryos, as compared with ∼52% in control embryos (Figure 7B), indicating that loss of NOP-1 is sufficient to impair polarization. Although cortical flows persist in par-2(RNAi) embryos (Munro et al., 2004), microtubule-directed localization of PAR-2 can establish polarity in embryos defective in RhoA activation (Zonies et al., 2010; Motegi et al., 2011). Depletion of PAR-2 eliminated the residual polarization observed in nop-1(it142) embryos (Figure 7A), analogous to the previously demonstrated redundancy between PAR-2 and ECT-2 (Zonies et al., 2010).

To gain further insight into the mechanism by which NOP-1 promotes embryo polarization, we investigated whether NOP-1 regulates the dynamics of PAR protein accumulation. We imaged embryos coexpressing mCherry::PAR-6 and GFP::PAR-2 in control and NOP-1–depleted embryos. In control embryos, mCherry::PAR-6 initially occupies the entire cortex near the end of meiosis. As PAR-6 begins to recede from the posterior pole, GFP::PAR-2 is recruited to the cortex at the posterior, and the PAR-2–PAR-6 boundary shifts anteriorly until both domains occupy ∼50% of the length of the cortex.
Recruitment of GFP::PAR-2 occurs around 600 s before nuclear envelope breakdown (NEBD) in control embryos (Figure 7, C and D). GFP::PAR-2 recruitment is delayed by ∼150 s in nop-1(it142) embryos and nop-1(RNAi) embryos, occurring ∼450 s before NEBD (Figure 7, C and D). A similar delay in symmetry breaking has been reported in ect-2(ax751ts) embryos (Zonies et al., 2010). Indeed, both hypomorphic, temperature-sensitive ect-2(ax751ts) embryos and ect-2(RNAi) embryos recruit GFP::PAR-2 ∼400 s before NEBD (Figure 7, C and D). GFP::PAR-2 recruitment is further delayed to ∼200 s before NEBD in ect-2(ax751ts);ect-2(RNAi) embryos. Likewise, PAR-2 recruitment is significantly delayed in nop-1(it142);ect-2(RNAi) embryos and ect-2(ax751ts);nop-1(RNAi) embryos, as compared with the single mutants (Figure 7, C and D). Thus the residual ECT-2 activity in ect-2(RNAi) embryos and in ect-2(ax751ts) embryos requires NOP-1. Intriguingly, the strong inactivation of ECT-2 by depletion of the mutant protein delays polarization more strongly than mutational inactivation of NOP-1, raising the possibility of trace ECT-2 activity in the absence of NOP-1.

**DISCUSSION**

We have demonstrated that both polarization and cytokinesis involve a protein, NOP-1, that promotes the characteristic events downstream of RhoA activation. NOP-1 is critical for these events during polarization. During cytokinesis, NOP-1 makes a significant contribution to these processes, but the centralspindlin subunit CYK-4 plays the dominant role in promoting RhoA-dependent events. Due to redundant pathways for polarization and cytokinesis, the majority of nop-1 embryos successfully polarize and divide, despite significant molecular defects.

NOP-1 is an unusual protein. It is serine-rich and lacks obvious conserved domains and orthologues outside the *Caenorhabditis* genus. Interestingly, pseudocleavage is observed in embryos of many, but not all, species within the Rhabditidae family (Brauchle et al., 2009). The absence of molecular clues creates challenges for the elucidation of NOP-1's function. However, thorough phenotypic characterization indicates that NOP-1 is required for the cortical recruitment of myosin and anillin and the proper organization of F-actin during establishment phase of polarization and during central spindle–independent cytokinesis (Figure 8A). Cortical accumulation of these RhoA effectors are largely independent of each other. Furthermore, NOP-1 promotes cortical recruitment of a RhoA biosensor.
Therefore NOP-1 is not likely to mediate the biosynthesis or recruitment of one effector and indirectly inhibit recruitment of the other effectors. More likely, NOP-1 acts upstream of these components by promoting accumulation of active RhoA (Figure 8B). In principle, it could do so by inhibiting an inhibitor of RhoA. However, NOP-1 is required for cortical contractility in embryos depleted of a Rho family GAP, RGA-3/4, which acts antagonistically to ECT-2, formally ruling out the possibility that NOP-1 is an inhibitor of this RhoGAP. NOP-1 is also required for contractility in nrl-1 embryos, another condition that would favor accumulation of active RhoA. Thus we favor the hypothesis that NOP-1 promotes RhoA activation. The factors known to promote RhoA activation include the RhoGEF ECT-2 and the RhoGAP CYK-4. Whereas ECT-2 is required for all RhoA activity in the embryo, NOP-1 and CYK-4 promote separable aspects of RhoA function. As NOP-1 function in the embryo requires ECT-2, it must function upstream of ECT-2.

How might NOP-1 promote RhoA activation? There are indications of ECT-2 auto-inhibition. The N- and C-termini of ECT-2 interact, and mutations in the N-terminus can activate aspects of ECT-2 function (Kim et al., 2005). During cytokinesis in human cells, direct binding of CYK-4 to ECT-2 is required for ECT-2 activity, and this may relieve ECT-2's autoinhibition (Yüce et al., 2005; Burkard et al., 2009; Wolfe et al., 2009). Interestingly, the GAP domain of CYK-4 also contributes to full activation of ECT-2. Although CYK-4 and NOP-1 are not detectably related, a phosphorylated form of NOP-1 might interact with the tandem BRCT domains of ECT-2, as is the case for CYK-4 (Wolfe et al., 2009).

NOP-1 is required for RhoA-dependent contractility events during polarization. In contrast, cyk-4(or749ts) embryos, which contain a substitution mutation in the GAP domain, do not have a detectable phenotype during polarity establishment. Although sperm-derived CYK-4 was proposed to promote polarization in the early embryo through local RhoA-GAP activity at the embryonic posterior (Jenkins et al., 2006), this proposal is not consistent with observations in cyk-4(or749ts) embryos, nor has it garnered additional experimental support.

During cytokinesis, RhoA activation relies on both NOP-1 and CYK-4. In nop-1 embryos, the vast majority of RhoA-dependent processes require CYK-4 and vice versa. These results suggest that NOP-1 and CYK-4 are parallel activators of ECT-2 during cytokinesis, with CYK-4 playing a quantitatively dominant role. Indeed, cytokinesis can proceed to completion in nop-1 embryos, but furrow progression is incomplete in cyk-4 embryos (Jantsch-Plunger et al., 2000).

Furrow formation during cytokinesis involves both astral microtubules and the central spindle. CYK-4 is required for central spindle-directed furrowing and NOP-1 is primarily involved in aster-induced furrowing. Previous studies (Werner et al., 2007), have shown that central spindle-directed accumulation of RhoA effectors is highly localized. In contrast, central spindle-independent accumulation of RhoA effectors, shown here to require NOP-1, is largely dispersed (Figure 8C). Microtubules and anillin function together to focus these dispersed effectors (Tse et al., 2011).

Depletion of a number of proteins can prevent furrowing in central spindle-defective embryos. This list includes factors that promote spindle elongation (GOA-1 and GPA-16, GPR-1/2, LET-99), factors that promote myosin organization (ANI-1), and factors that promote myosin contractility (protein phosphatase 6, PPH-6, and its associated subunit SAPS-1). When spindle elongation is perturbed in central spindle-defective embryos, cortical myosin
accumulates, but it is not sufficiently polarized to allow formation of a ingressing furrow (Dechant and Glotzer, 2003; Bringmann et al., 2007; Werner et al., 2007). When ANI-1 is depleted, myosin accumulates on the cortex, but it is not organized into large foci, nor is the myosin highly polarized, again preventing furrow formation in central spindle–defective embryos (Tse et al., 2011). Depletion of PPH-6 or SAPS-1 also allows myosin accumulation (Afshar et al., 2010), although it is less contractile, perhaps due to a failure in Aurora A activation (Zeng et al., 2010). Thus, in each of these cases, there is significant cortical recruitment of RHO-1 effectors, creating the potential for aster-dependent contractility. NOP-1 is distinct from these factors in that its inactivation in central spindle–deficient embryos abrogates recruitment of cortical myosin II and anillin. Thus NOP-1 depletion is the most stringent means available to inhibit aster-directed furrowing without greatly compromising central spindle–directed furrowing.

In wild-type C. elegans embryos, the cleavage furrow ingresses asymmetrically. Asymmetric furrow ingression is observed in a variety of cell types, but its mechanistic basis is not understood. Asymmetric ingression requires anillin and septins, and it is associated with increased myosin recruitment at the rapidly ingressing side of the furrow (Maddox et al., 2007). The trigger for symmetry breaking is not known. One proposal involves a positive feedback loop in which contractile proteins recruit anillin, which promotes increased accumulation of contractile proteins (Maddox et al., 2007). Unexpectedly, we have found that embryos deficient in both NOP-1 and anillin have asymmetrically ingressing cleavage furrows. Thus there are anillin–independent mechanisms that lead to furrow asymmetry. From a genetic perspective, a corollary of these findings is that NOP-1 prevents asymmetric furrow formation in ANI-1–deficient embryos. Furrow formation in embryos deficient in both NOP-1 and anillin is entirely due to central spindle–dependent activation of RhoA. One explanation of these results could be that central spindle–directed furrowing is intrinsically asymmetric, perhaps due to the effect of spindle oscillations on the proximity of the central spindle to the cortex. When anillin is absent, NOP-1–dependent cortical accumulation of RhoA effectors may antagonize the asymmetric furrowing induced by the central spindle pathway. However, other explanations are also viable, and additional analysis of the mechanisms governing recruitment of contractile proteins is required to more deeply understand the mechanics of cleavage furrow ingress.

NOP-1 is not only an important, nonessential regulator of cytokinesis, it is also an important, nonessential regulator of cell polarization. The molecular function of NOP-1 in these two processes appears similar, namely, promoting RhoA activation. Polarization is known to involve RhoA and ECT-2. A mutation in ect-2, ax751ts, causes a phenotype similar to that seen with mutations in nop-1 (Zonies et al., 2010). This allele of ect-2 is hypomorphic; further depletion of ECT-2 by RNAi or inactivation of NOP-1 strengthens the phenotype, consistent with the model that NOP-1 is an activator of ECT-2. Indeed, both nop-1(it142) and ect-2(ax751ts) delay embryo polarity establishment and the combination of either mutation with depletion of PAR-2 abrogates embryo polarization (this paper and Zonies et al., 2010). These results demonstrate that NOP-1 functions in a RhoA-dependent pathway for embryo polarization that acts in parallel to a PAR-2–dependent pathway.

Collectively our results indicate that NOP-1 serves as a second upstream activator of the RhoA GEF, ECT-2, acting in parallel with CYK-4. These two activators are not simply redundant. During polarization, CYK-4 appears largely inactive and NOP-1 is required for the majority of RhoA-dependent events. During cytokinesis, both activators are active, but they appear to function at different sites, as determined by their differences in localization. During cytokinesis, CYK-4 appears largely inactive and NOP-1 is required for polarity establishment and the combination of either mutation with depletion of ECT-2 by RNAi or inactivation of NOP-1 strengthens the phenotype, consistent with the model that NOP-1 is an activator of RhoA. Opposite CYK-4::GFP localization, it is also an important, nonessential regulator of cell polarization.

MATERIALS AND METHODS

Strains

C. elegans strains (listed in Table S1) were maintained on nematode growth medium (NGM) plates using standard procedures. Unless otherwise specified, strains were provided by the Caenorhabditis
Genetics Center, which is funded by the National Institutes of Health (NIH) National Center for Research Resources.

RNAi

Generally, RNAi was administered by feeding nematodes with Escherichia coli expressing the appropriate double-stranded RNA (dsRNA; Timmons and Fire, 1998). HT115 bacterial cultures were grown in Luria broth with 100 μg/ml ampicillin overnight at 37°C. Cultures (200 μl) were seeded on NGM plates containing 100 μg/ml ampicillin and 1 mM isopropyl β-D-thiogalactopyranoside and incubated at room temperature (∼23°C) for 8 h. RNAi plasmids were obtained from the library produced by Kamath et al. (2003). Young L4 hermaphrodites were picked onto the plates for feeding at 25°C at least 24 h prior to dissection. The one exception to this was for Supplemental Figure S5, for which ECT-2 was depleted by injection of in vitro synthesized dsRNA. In this case, analysis was restricted to embryos that exhibited no furrowing in either the first or the second division.

Mapping of nop-1

nop-1 was previously mapped to the region of chromosome III between dpy-17 and unc-32 (Rose et al., 1995). Deficiency mapping placed nop-1 in the 0.06 cM region between the deficiencies sDf121 and sDf127. For further refinement of the mapping, lon-1(e185);nop-1(t42);unc-36(e251)/qC1 animals were crossed with unc-32(e189);cyk-3(t1525);sma-4(e729)/qC1 animals. We picked 111 recombinants that separated sma-4 and unc-32. Thirteen recombinants’ lines produced viable progeny, indicating that the recombination event took place between sma-4 and cyk-3. We performed time-lapse Nomarski imaging of at least 10 embryos from these 13 lines. Seven lines produced some embryos with pseudocleavage defects, positioning nop-1 approximately midway between sma-4 and cyk-3. We then used RNAi to the genes in this specific region to identify the gene, F25B5.2, whose depletion phenocopies nop-1(e42).

Transgenic lines

Transgenic lines expressing GFP::NOP-1, GFP::Utrophin, and GFP::AHPH were integrated into the Mos element ttTi5605 on chromosome II using the MosSCI method (Frøkjær-Jensen et al., 2008).

Microscopy

Gravid hermaphrodites were dissected in egg salt buffer on coverslips, mounted on 2.5% agarose pads, and sealed with Vaseline. For Nomarski imaging, embryos were observed with a Zeiss (Thornwood, NY) Axiosplan II with a 100x/1.3 Plan-Neofluar objective. Images were captured with a charged-couple device (CCD) camera (Imaging Source, Charlotte, NC) controlled by Gawk (http://gawk.sourceforge.net). Images were acquired every 5 s and processed with ImageJ (http://rsweb.nih.gov/ij). For confocal imaging, embryos were imaged with a 63x/1.4 numerical aperture oil-immersion lens on 1) a Zeiss Axiovert 200M equipped with a Yokogawa CSU-10 spinning-disk unit (McBain, Simi Valley, CA) and illuminated with 50-mW, 473-nm and 20-mW, 561-nm lasers (Cobolt, Solna, Sweden), or 2) a Zeiss Axiosmager M1 equipped with a Yokogawa CSU-X1 spinning-disk unit (Solamere, Salt Lake City, UT) and illuminated with 50-mW, 488-nm and 50-mW, 561-nm lasers (Coherent, Santa Clara, CA). Images were captured on a Cascade 1K EM-CCD camera or a Cascade 512B7T (Photometrics, Tucson, AZ) controlled by MetaMorph (Molecular Devices, Sunnyvale, CA). Image processing was performed with ImageJ. Z-stacks were projected with a maximum-intensity algorithm. Time-lapse projections were assembled into movies using MetaMorph and ImageJ.

Image quantification

For measurement of the kinetics of membrane ingestion, montages produced from embryos expressing the plasma membrane marker GFP::PH were opened in ImageJ, and the tips of the membrane ingestion were manually positioned. The coordinates were then imported into Excel for further analysis. The maximum length of myosin cap and the timing of GFP::PAR-2 recruitment were manually analyzed in ImageJ. To quantify the accumulation of GFP::AHPH at the furrow region, we measured the intensity of pixels that exceeded a threshold of 1.45 × background for each slice. To quantify the accumulation of GFP::NMY-2 and GFP::AN-1 in the entire cortex, we measured the intensity of pixels that exceeded a threshold of 1.45 × background for each slice. To quantify the changes of GFP::Utrophin, we measured the average intensity in a 30-pixel-wide region at 50–75% egg length.

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