RESEARCH ARTICLE



Aurora B functions at the apical surface after specialized cytokinesis during morphogenesis in *C. elegans*

Xiaofei Bai^{1,*,§§}, Michael Melesse^{1,‡,§§}, Christopher G. Sorensen Turpin^{1,§§}, Dillon E. Sloan^{2,1,§§}, Chin-Yi Chen³, Wen-Cheng Wang³, Po-Yi Lee³, James R. Simmons¹, Benjamin Nebenfuehr^{1,§}, Diana Mitchell^{2,¶}, Lindsey R. Klebanow^{1,**}, Nicholas Mattson^{1,‡‡}, Eric Betzig⁴, Bi-Chang Chen^{4,3}, Dhanya Cheerambathur⁵ and Joshua N. Bembenek^{2,1,¶¶}

ABSTRACT

Although cytokinesis has been intensely studied, the way it is executed during development is not well understood, despite a long-standing appreciation that various aspects of cytokinesis vary across cell and tissue types. To address this, we investigated cytokinesis during the invariant Caenorhabditis elegans embryonic divisions and found several parameters that are altered at different stages in a reproducible manner. During early divisions, furrow ingression asymmetry and midbody inheritance is consistent, suggesting specific regulation of these events. During morphogenesis, we found several unexpected alterations to cytokinesis, including apical midbody migration in polarizing epithelial cells of the gut, pharynx and sensory neurons. Aurora B kinase, which is essential for several aspects of cytokinesis, remains apically localized in each of these tissues after internalization of midbody ring components. Aurora B inactivation disrupts cytokinesis and causes defects in apical structures, even if inactivated post-mitotically. Therefore, we demonstrate that cytokinesis is implemented in a specialized way during epithelial polarization and that Aurora B has a role in the formation of the apical surface.

KEY WORDS: Apical surface, Aurora B kinase, Cytokinesis, Midbody, Morphogenesis

INTRODUCTION

Cytokinesis is the final step of cell division and is not only required to generate two daughter cells, but also regulates development and

^{¶¶}Author for correspondence (bembenek@umich.edu)

D.E.S., 0000-0001-6065-176X; D.M., 0000-0003-3788-2265; L.R.K., 0000-0002-0125-2619; N.M., 0000-0002-6950-9222; B.-C.C., 0000-0002-1876-7432; J.N.B., 0000-0002-0966-2268

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Received 5 June 2019; Accepted 26 November 2019

cellular organization in tissues (Canman et al., 2000; Chen et al., 2013: Herszterg et al., 2014: Li, 2007: Oegema and Hyman, 2006). Signals from the anaphase spindle trigger cleavage furrow ingression (Bringmann and Hyman, 2005; Eggert et al., 2006). After furrowing, cells remain connected at the midbody, a membrane channel containing microtubules, vesicles, a central spindle and contractile ring proteins (El Amine et al., 2013; Green et al., 2012; Hu et al., 2012; Schiel et al., 2013; Skop et al., 2004). The ESCRT machinery mediates the final abscission event (Carlton and Martin-Serrano, 2007; Guizetti et al., 2011; Schiel et al., 2011). Aurora B kinase, part of the chromosome passenger complex, promotes cytokinesis and regulates abscission timing (Carlton et al., 2012; Carmena et al., 2015, 2012; Mathieu et al., 2013; Norden et al., 2006; Steigemann et al., 2009). We sought to investigate the dynamics of cytokinesis during the completely described invariant Caenorhabditis elegans embryonic divisions (Sulston et al., 1983).

Cells normally follow the standard cytokinetic process, but several exceptions are known. Some cells do not complete cytokinesis and become polyploid, such as liver and intestinal cells (Amini et al., 2015: Fox and Duronio, 2013: Hedgecock and White, 1985; Lacroix and Maddox, 2012). Germ cells do not complete abscission and remain connected, allowing cytoplasmic exchange (Greenbaum et al., 2007; Haglund et al., 2011; Hime et al., 1996; Maddox et al., 2005), which also occurs in several other cell types (Daniel et al., 2018; McLean and Cooley, 2013; Zenker et al., 2017). The cleavage furrow can be repositioning during anaphase (Ou et al., 2010) or ingress asymmetrically to the apical surface as observed in epithelial tissues (Bourdages et al., 2014; Daniel et al., 2018; Founounou et al., 2013; Guillot and Lecuit, 2013; Herszterg et al., 2014; Higashi and Miller, 2017; Paolini et al., 2015; Wang et al., 2018). Therefore, cytokinesis is altered in different contexts, but more investigation is required to understand the functional purpose of these changes and how they are achieved.

The midbody can be released extracellularly after abscission (Chen et al., 2013; Crowell et al., 2014; König et al., 2017). The midbody remnant (MBR) can persist extracellularly or be engulfed depending on the cell type (Dubreuil et al., 2007; Ettinger et al., 2011; Salzmann et al., 2014). Once internalized, the MBR can elicit intracellular signaling (Peterman et al., 2019). In the early *C. elegans* embryo, MBRs are reproducibly phagocytosed by specific daughter cells (Fazeli et al., 2016; Ou et al., 2014; Singh and Pohl, 2014). This suggests that the MBR regulates cell fate, although the mechanism needs to be further elucidated.

Madin–Darby canine kidney (MDCK) cells can form a lumen, which begins with delivery of apical membrane proteins to the midbody (Li et al., 2014; Reinsch and Karsenti, 1994; Schlüter et al., 2009). Abscission timing and midbody positioning impact lumen formation (Lujan et al., 2016). Vesicle trafficking during

¹Department of Biochemistry, Cellular and Molecular Biology, University of Tennessee, Knoxville, TN 37996, USA. ²Department of Molecular, Cellular and Developmental Biology, University of Michigan, Ann Arbor, MI 48109, USA. ³Research Center for Applied Sciences, Academia Sinica, Taipei, Taiwan. ⁴Janelia Research Campus, HHMI, Ashburn, VA 20147, USA. ⁵Wellcome Centre for Cell Biology, University of Edinburgh, Edinburgh, EH9 3BF, UK. *Present Address: National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health, Bethesda, MD 20892, USA. [‡]Present Address: Oak Ridge National Labs, Oak Ridge, TN 37830, USA. [§]Present Address: Molecular, Cellular & Developmental Biology, University of Colorado, Boulder, Boulder, CO 80309, USA. [¶]Present Address: Biological Sciences, University of Idaho, Moscow, ID 83844, USA. **Present Address: United States Food and Drug Administration, Silver Springs, MD 20993, USA. ^{§§}These authors contributed equally to this work

cytokinesis promotes lumen formation in other systems (Wang et al., 2014b). Delayed abscission allows vesicles to deliver apical proteins to the membrane in mouse blastomeres (Zenker et al., 2017). The midbody becomes the apical process in chick neuronal progenitors (Wilcock et al., 2007), defines the site of polarization for dendrite extension in *Drosophila* neurons (Pollarolo et al., 2011) and regulates polarity in *Drosophila* neuroblasts (Loyer and Januschke, 2018). Epithelial cells establish new junctions at the midbody to maintain tissue integrity during division (Daniel et al., 2018; Higashi et al., 2016; Wang et al., 2018). The MBR is a polarizing cue during dorsoventral axis formation in the *C. elegans* embryo (Singh and Pohl, 2014; Waddle et al., 1994). The MBR can play a role in cilium formation (Bernabé-Rubio et al., 2016). Further effort is needed to understand how cytokinesis and the midbody regulate pattern formation in tissues.

Forming an apical surface is crucial for proper tissue architecture (Overeem et al., 2015). During morphogenesis in C. elegans, cells complete the embryonic divisions and organize into tissues (Chisholm and Hardin, 2005; Leung et al., 1999; Mango, 2007). In epithelial tissues, cells polarize to form apical and basal membranes, reorganize the cytoskeleton and form junctions mediated by adhesion proteins (Bryant and Mostov, 2008; Li and Gundersen, 2008; Niessen et al., 2011). The C. elegans intestine polarizes after the E8-E16 division. PAR and adherens junction proteins accumulate in the polar membrane, associate with centrosomes, and then move to the nascent apical midline (Achilleos et al., 2010; Feldman and Priess, 2012). The pharynx divides and polarizes slightly later than the intestine (Portereiko and Mango, 2001). The amphid sensory neurons share features of epithelia (Low et al., 2019) and initially organize into a multicellular rosette with a central apical domain that extends into a dendrite (Fan et al., 2019). Kinetochore proteins regulate microtubules during the initial stages of dendrite extension (Cheerambathur et al., 2019). Cytokinetic regulators have been implicated in epidermal and pharyngeal morphogenesis in roles independent of cell division (Hardin et al., 2008; Portereiko et al., 2004; Von Stetina et al., 2017). Cytokinesis in neuroblasts is important for overlying epidermal cells to migrate during enclosure (Fotopoulos et al., 2013). Therefore, mitotic regulators have mitotic and post-mitotic functions during morphogenesis, although this is not well understood.

We investigated the dynamics of cytokinesis in the invariant *C. elegans* embryonic divisions, including a detailed analysis of several tissues at the onset of morphogenesis. We investigated furrow symmetry, central spindle dynamics, abscission timing and the fate of several different proteins during midbody inheritance. Finally, we investigated mitotic and post-mitotic functions of Aurora B in cytokinesis and at the apical surface in several tissues.

RESULTS

Furrow asymmetry and midbody inheritance in the early embryo

The first mitotic division of P0 generates AB and P1 daughter cells (Fig. 1A). We examined different regulators of cytokinesis to label the central spindle, the cytokinetic furrow and the flank and ring sub-structures of the midbody (Green et al., 2012). To observe the midbody flank region, we imaged Aurora B kinase (AIR-2) (Bembenek et al., 2013), microtubules (Lee et al., 2015), and the membrane trafficking regulator RAB-11 (Sato et al., 2008) (Fig. 1, Fig. S1G-J, Movie 1). We imaged NMY-2 (Nance et al., 2003), which labels the contractile ring and MBR, and the central spindle and MBR (Fig. 1G-P, Movie 1). The first furrow is slightly

asymmetric (Maddox et al., 2007) and the midbody forms in a central position (Fig. 1B,C,G,H,L,M). AIR-2::GFP and tubulin localized to the central spindle and midbody (Fig. 1B,C, Fig. S1G, H, Movie 1). The MBR from the first mitotic division was always inherited by the P1 daughter cell (Fig. 1A,D,I,N,S). Abscission occured within 8 min of furrowing onset, indicated by loss of microtubules (Fig. 1X, Fig. S1H) (Green et al., 2013; König et al., 2017). AIR-2 was lost from the midbody flank but remained on the MBR with other ring components after internalization (Fig. 1D,E,I, J,N,O, Movie 1). Therefore, multiple proteins remain on the MBR after internalization, which may affect its function in P1.

In the second mitotic divisions, several reproducible alterations to cytokinesis were observed. In AB cytokinesis, an asymmetric furrow ingressed from the outer surface until it contacted EMS (Fig. 1D,E,I,J,N,O,S,T). The asymmetry parameter, which is the ratio of furrow ingression distance from each side (Maddox et al., 2007), was 1.7 in the first division, but 21.6 in AB and 16.1 in P1 (Fig. 1V). AIR-2 localized to the spindle midzone in AB, which is swept toward EMS during furrow ingression (Fig. 1E, Fig. S1I,J, Movie 1), and remained associated with the MBR after it was engulfed (Fig. 1D-F, Movie 1). NMY-2 and ZEN-4 also remained on the MBR (Fig. 1I,J,N,O, Movie 1). RAB-11 briefly accumulated at the midbody in the first two mitotic divisions and was not observed on the MBR (Fig. 1Q-U). Therefore, furrow ingression is highly asymmetric during the second division.

The MBR is reproducibly inherited in the early divisions. The AB MBR was invariably engulfed by EMS instead of either of the AB daughter cells (Fig. 1F,K,P,U, Movie 1). Further, the P0 midbody was always inherited by EMS. Abscission timing was relatively fast in both AB and P1 cell divisions as indicated by microtubule disassembly (Fig. 1X, Fig. S1I-J). Disruption of polarity by *par-3(RNAi)* caused random midbody inheritance as previously observed (Ou et al., 2014), and altered furrow symmetry (asymmetry parameter increased to 4.2 in P0, and decreased to 3.0 in AB and 4.4 in P1; Fig. S1K-O). Therefore, both midbody inheritance and furrow symmetry depend on polarity in the early embryo.

Aurora B remains on the apical surface after E8-E16 gut cytokinesis

We next analyzed cytokinesis during morphogenesis, which revealed several novel patterns. The intestine is derived from the E blastomere, which undergoes five embryonic divisions (Leung et al., 1999). The E8 to E16 division occurs around 280 min after the first cleavage and is followed by epithelial polarization (Leung et al., 1999). We observed a highly modified cytokinesis during polarization (Fig. 2A). E8 cells underwent symmetric furrowing over 4.7 min (n=15; 1.0 asymmetry parameter; Fig. 1V), to produce a centrally placed midbody (Fig. 2B,H, Fig. S2A,D,G,J, Movies 2-4). Using lattice light-sheet imaging, we observed E8 midbodies migrating to the nascent apical midline over 30 min (Fig. 2C,E, Movie 2). AIR-2::GFP localized on elongated spindle midzone microtubules during movement (Fig. 2E,F, Fig. S2A-C, Movies 2-4). The length of the spindle midzone microtubules relative to the cell was 0.47 (average 4.6 µm/9.8 µm) in the intestinal cell division, which is more than twice that of the early two cell divisions 0.17 (average 9.3 μ m/53.4 μ m) in P0 and 0.17 (average 7.7 μ m/44.3 μ m) in AB (Fig. 1W). The midzone microtubules persisted for over 25 min on average before they could no longer be distinguished at the apical midline, indicating a delay in abscission (Figs 1X and 2I). The ring markers ZEN-4 and NMY-2 internalized shortly after the midbody reached the apical midline $(553\pm140 \text{ s and } 545\pm179 \text{ s},$ respectively), indicating MBR internalization (Fig. 2G, Fig. S2D-I,O,

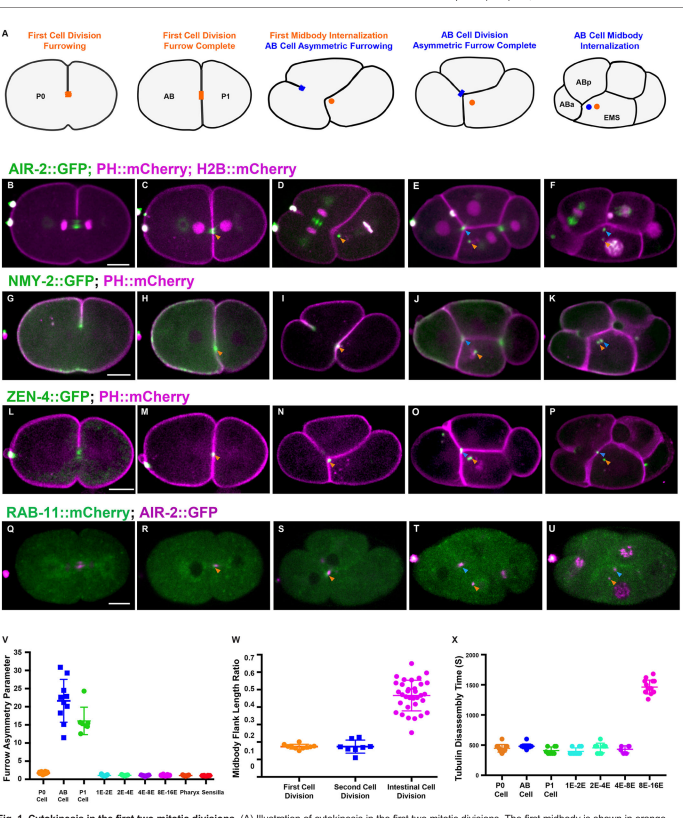


Fig. 1. Cytokinesis in the first two mitotic divisions. (A) Illustration of cytokinesis in the first two mitotic divisions. The first midbody is shown in orange, the AB midbody is blue. (B-F) Cytokinesis labeled with AIR-2::GFP (green; PH::mCherry and H2B::mCherry in magenta). AIR-2 localizes on the central spindle (B) and the midbody flank (C, orange arrowhead) and the MBR after internalization in AB (D, orange arrowhead). In the AB division, the asymmetric furrow pushes the midzone against EMS (E, blue arrowhead), which engulfs it (F, blue arrowhead). (G-K) NMY-2::GFP (green; PH::mCherry in magenta) localizes to the furrow (G) and midbody ring (H-K). (L-P) ZEN-4::GFP (green; PH::mCherry in magenta) appears on the central spindle (L) and the midbody (M-P). (Q-U) RAB-11:: mCherry (green) colocalized with AIR-2::GFP (magenta) at the midbody for a short time before internalization (R-U). Arrowheads in G-U are as described for B-F. (V) Furrow asymmetry parameter is shown for different divisions. (W) Ratio of midbody microtubule length to cell length in different divisions. (X) Quantification of microtubule persistence in different cell divisions. Error bars indicate s.d. Scale bars: 10 μm.

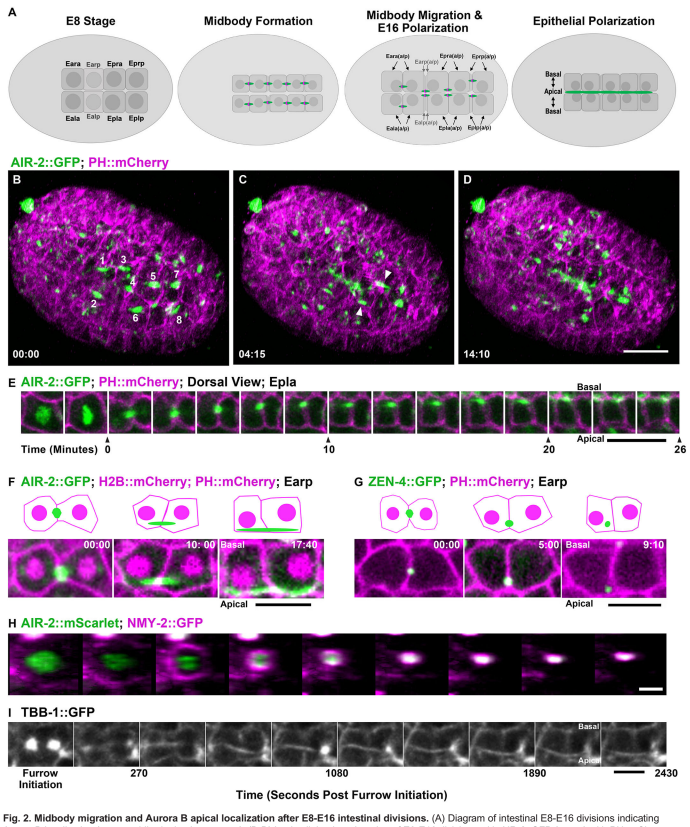


Fig. 2. Midbody migration and Aurora B apical localization after E8-E16 intestinal divisions. (A) Diagram of intestinal E8-E16 divisions indicating Aurora B localization (green; midbody ring in magenta). (B-D) Lattice light-sheet imaging of E8-E16 divisions with AIR-2::GFP (green) with PH::mCherry (magenta). AIR-2::GFP labels midbodies (labeled 1-8 in B) that migrate to the nascent apical surface (arrowheads, C) where it persists (D). Scale bar: 10 µm. (E) Montage of Epla division with AIR-2::GFP (green; PH::mCherry in magenta) showing midbody formation (t=0) and migration to the apical midline. Scale bar: 5 µm. (F,G) Comparison of AIR-2::GFP (F) and ZEN-4::GFP (G) localization to the apical midline. Scale bars: 5 µm. Time shown in minutes:seconds. In schematics, midbody is green, H2B::mCherry and PH::mCherry are magenta. (H) En face view of the E8-E16 contractile ring labeled with NMY-2::GFP (magenta; AIR-2::GFP (magenta; Scale bar: 5 µm. (I) Single z-plane imaging of midbody flank microtubules during Epra cell division. Scale bar: 5 µm.

Movie 3). Therefore, abscission occurs after the midbody migrates to the apical midline. To our knowledge, this is the first observation of apical migration of the midbody in *C. elegans*.

We observed another novel behavior of Aurora B after the E8-E16 division. AIR-2::GFP remained localized at the apical midline (Fig. 2D-F, Fig. S2C,O, Movies 2-4) after ZEN-4 and NMY-2 were internalized (Fig. 2G, Fig. S2D-I,O) colocalizing with the apical polarity marker PAR-6 (Schonegg et al., 2007) (Fig. S2N). Endogenously tagged AIR-2::GFP and immunostained endogenous AIR-2 could also be observed at the apical midline (Fig. S1A,D). The centrosome also migrates to the midline during E16 polarization (Feldman and Priess, 2012; Yang and Feldman, 2015). Centrosomes labeled with γ -tubulin::GFP (Redemann et al., 2010) moved to the apical surface at the same time as AIR-2::GFP localized in the spindle midzone (Fig. S2P). In contrast, the first three E cell divisions exhibited symmetric furrowing (Fig. 1V), rapid abscission timing (Fig. 1X) and no cortical localization of AIR-2::GFP after cytokinesis (Fig. S3). Therefore, after cytokinesis in the E8-E16 divisions, the MBR is internalized but Aurora B remains at the apical surface.

RAB-11 vesicle trafficking of apical proteins to the midbody establishes the apical membrane in other systems (Schlüter et al., 2009). In *C. elegans*, apical RAB-11 endosomes control trafficking in the intestine through adulthood (Sato et al., 2014). We imaged RAB-11::mCherry during the E8-E16 division and found that it colocalizes with AIR-2::GFP, migrates to the apical surface with the midbody and remains localized to the apical surface well after cytokinesis is complete (Fig. S2J-L). Therefore, apical RAB-11 localization is established during E8-E16 cytokinesis and intestinal epithelial polarization.

The anterior and posterior pairs of E16 cells (Ealaa, Earaa, Eplpp and Eprpp) undergo one last embryonic division to achieve the E20 intestine stage. In the four central E8 cells, which do not divide again (Ealp, Earp, Epla and Epra), the midbody migrated to the midline at E8-E16 as described above. However, the midbodies from Eala, Eara, Eplp and Eprp migrated toward the midline at E8-E16, but the AIR-2 signal diminished (Fig. S2M). The midbodies of the E16-E20 divisions also underwent apical migration after symmetrical furrowing (Movie 5). Therefore, apical midbody migration occurs both during and after epithelial polarization in the intestine.

Inactivation of Aurora B kinase disrupts E8-E16 cytokinesis and proper epithelial polarization

We next investigated the function of Aurora B and other cytokinetic regulators during the E8-E16 division. To bypass the essential function of cytokinetic regulators during the early divisions, we inactivated temperature-sensitive (ts) mutants after isolating twocell embryos and incubating them at the permissive temperature (15°C) until different stages before shifting them to the nonpermissive temperature (26°C) until they hatched (Fig. 3A). After epidermal enclosure, the embryo begins to elongate during the bean stage, followed by the comma, 1.5-fold, 2-fold and 3-fold stages, after which the larvae hatches from the egg. Two-cell air-2(or207) (Severson et al., 2000) embryos had only 53.6% (37/69) viability when kept at 15°C, indicating that this mutant is sick even at permissive temperature, whereas wild-type N2, zen-4(or153) (Severson et al., 2000) and spd-1(oj5) (O'Connell et al., 1998) embryos were 100% viable at 15°C (Table S1). Embryos shifted from 15°C to 26°C after 4.5 h (corresponding to late E4 to early E8 stages) or 6.5 h (corresponding to E8-E16 stage or early bean stage) showed significant lethality in both *air-2(or207)* and *zen-4(or153)*, but not *spd-1(oj5)* (Table S1). Mutant embryos shifted after most divisions had finished at the comma to 1.5-fold stage hatched at a rate similar to those at permissive temperature (Table S1). The *air-2(or207)* mutant has penetrant cytokinesis failure immediately after inactivation in early embryos (Severson et al., 2000) but this was not observed in in older embryos (37.8% cytokinesis failure in *n*=138 divisions; Fig. S5B,D; see Materials and Methods). Consistent with this, only mild lagging chromosome segregation defects were observed in older *air-2(or207)* mutants (Fig. S5F). A temperature-sensitive mutant of the INCENP homolog *icp-1(or663ts)* (Davies et al., 2014) also did not have penetrant cytokinesis failures at the E8-E16 division (9%, *n*=22 divisions). Therefore, these mutants allow us to bypass early divisions but do not yield rapid-onset, highly penetrant cell division phenotypes later in development.

Next, we tested whether AIR-2 was required for the specialized E8-E16 divisions and epithelial polarization. We first asked whether microtubule organization required AIR-2. air-2(or207) mutant embryos shifted at the E4-E8 stage had reduced spindle midzone microtubules relative to wild type (Fig. 3B-I, Movie 6). Inactivation of air-2(or207) caused cytokinesis failure in 27% of the observed E8 cells and a failure to normally polarize all nuclei at the midline (Fig. 3F-I, Fig. S5A-C). In air-2(or207) E8 cells that did not fail cytokinesis, weak spindle midzone microtubules moved to the apical surface where microtubules accumulated (Fig. 3F-I, Movie 6). In neighboring cells that failed cytokinesis, microtubule accumulation at the apical midline was diminished, which was most obvious when both left and right E8 divisions failed at the same time (Fig. 3H, Fig. S5A-C, Movie 6). Therefore, AIR-2 regulates central spindle microtubules and completion of cytokinesis during the E8-E16 divisions. Furthermore, nuclear polarization and apical microtubule accumulation are disrupted when cytokinesis fails after inactivation of Aurora B.

The adhesion complex accumulates at the apical surface during polarization to promote gut lumen formation after the E8-E16 divisions (Achilleos et al., 2010). We imaged the α -catenin HMP-1::GFP (Marston et al., 2016), part of the cadherin-catenin adhesion complex that links cell-cell contacts with actin. PAR and adhesion complexes localize to cortical foci (Achilleos et al., 2010) and move with centrosomes to the apical midline (Feldman and Priess, 2012). We also observed HMP-1::GFP at the furrow and midbody throughout apical migration (Fig. 4A-E, Movie 7). HMP-1::GFP localized to the furrow and membrane adjacent to the midbody in the first three E cell divisions (Fig. S4), indicating that this localization is not specific to the E8-E16 division. This dynamic adhesion localization during cytokinesis may be important to maintain proper cell contacts during the disruptive process of division in the early embryo. In air-2(or207) mutants, furrow HMP-1::GFP was reduced during E8 cytokinesis (Fig. 4F-J, Fig. S5D,E, Movie 7). In E8 air-2(or207) divisions that completed cytokinesis, HMP-1::GFP signal accumulated at the apical midline (Fig. 4I, Fig. S5D,E, Movie 7). However, in air-2(or207) E8 divisions that failed cytokinesis, accumulation of HMP-1::GFP was delayed especially when pairs of E8 daughters on opposite sides of the midline both failed (Fig. 4F-J, Fig. S5D,E, Movie 7). Therefore, Aurora B inactivation leads to reduced furrow localization of α -catenin and disrupts cytokinesis, delaying accumulation of α -catenin at the apical surface during polarization.

In order to understand the effect of cytokinesis failure on lumen formation, we performed staining of the polarized gut using apical markers. We shifted mutant E4-E8 embryos to 26°C and fixed after intestinal polarization. We evaluated the apical surface by staining for the Ezrin-Radixin-Moesin homolog ERM-1 (van Furden et al.,

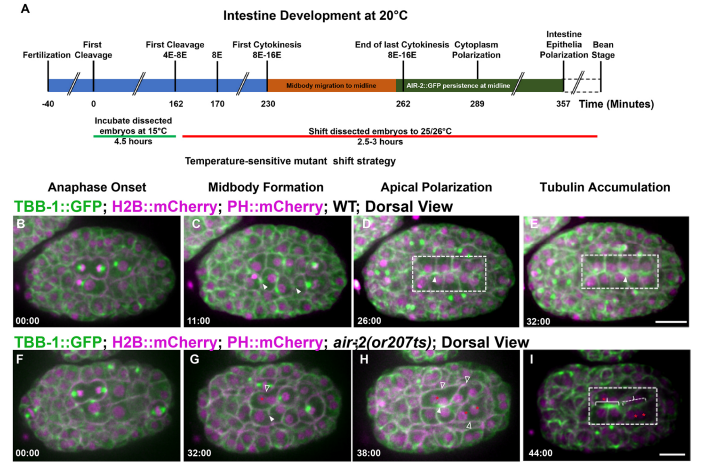


Fig. 3. Aurora B is required for E8-E16 cytokinesis and epithelial polarization. Microtubule dynamics during the E8-E16 divisions. (A) Time line of embryonic divisions and shifts for temperature-sensitive mutants. (B-E) In wild-type (WT) embryos, spindle midzone microtubules (TBB-1::GFP, green; H2B::mCherry and PH::mCherry shown in magenta) form in late anaphase (arrowheads, C) and migrate to the apical midline (D) and persist (E). (F-I) In *air-2(or207)* embryos, spindle midzone microtubules are diminished (G, filled arrowhead). When cytokinesis failures occur (G,H, unfilled arrowheads), nuclei fail to reach the apical midline (G-I, red asterisks) and apical microtubule accumulation is reduced (dashed bracket indicates failed divisions, solid bracket indicates successful divisions with apical accumulation). Dashed boxes indicate gut cells. Scale bars: 10 µm.

2004). We stained air-2(or207), zen-4(or153), spd-1(oj5) and *icp-1(or663)* and frequently observed deformed and binucleate cells in air-2(or207) and zen-4(or153) but not spd-1(oj5) mutants (Table 1, Fig. 5, Fig. S6). In all cases, ERM-1 was localized to the apical surface of the intestine and pharynx (Fig. 5A). However, ERM-1 staining was broadened, branched and/or discontinuous in air-2(or207) embryos (Fig. 5B-E, Table 1). Comma-stage air-2(or207) embryos still had disrupted ERM-1 staining, indicating that these defects are not resolved later in development (Fig. S6B,E). Furthermore, the intestine was highly mispositioned within the embryo (Fig. 5E) and the nuclei were disorganized (Fig. 5A-D, insets). Other apical markers also localized to the disorganized apical surface, including PAR-3, DLG-1 and IFB-2 (Fig. S6F-K). zen-4(or153) embryos had penetrant branched and discontinuous apical ERM-1 staining that was mispositioned at a lower rate (Table 1, Fig. S6C,E). spd-1(oj5) embryos displayed a significant but lower rate and severity of lumen defects (Table 1, Fig. S6D,E) despite having no lethality (Table S1) and minimal cytokinesis failures. We observed AIR-2::GFP dynamics in spd-1(oj5) E8-E16 divisions and found that AIR-2::GFP was lost from spindle midzone microtubules and instead accumulated at spindle poles, which moved to the apical surface in spd-1(oj5)embryos (Fig. S6E). Daughter cell pairs did not remain together in

spd-1(oj5) embryos, indicating that spindle midzone facilitates polarization (Fig. S6E). Therefore, AIR-2::GFP can still reach the apical surface through a compensatory mechanism when SPD-1 is inactivated. This compensatory mechanism is not perfect, leading to significant but reduced lumen defects. Finally, *icp-1(or663ts)* mutant embryos also had abnormal ERM-1 staining (Fig. S6L-M). Therefore, we conclude that Aurora B, the spindle midzone and other regulators of cytokinesis are required for normal apical lumen formation in the gut.

To inactivate AIR-2 more precisely, we tested whether tissuespecific depletion in the gut would cause lumen defects. We used a GFP-degradation system to deplete endogenously GFP-tagged Aurora B starting during the E8 stage (Wang et al., 2017). For comparison, we also depleted endogenously tagged NMY-2::GFP and ZEN-4::GFP (Dickinson et al., 2013; Lee et al., 2018). We quantified binucleate intestinal cells in L1 animals and observed 53.7% in AIR-2-depleted (n=23 animals), 19.7% in ZEN-4depleted (n=10), 13.6% in NMY-2-depleted (n=7) and 3% in control (n=10) embryos, indicating significant rates of cytokinesis failure. We stained embryos with ERM-1 after gut polarization to observe lumen defects. We measured four points along the length of the gut lumen and obtained an average width. In control embryos, the apical lumen was 1.15 ± 0.11 µm wide (n=10); AIR-2::GFP

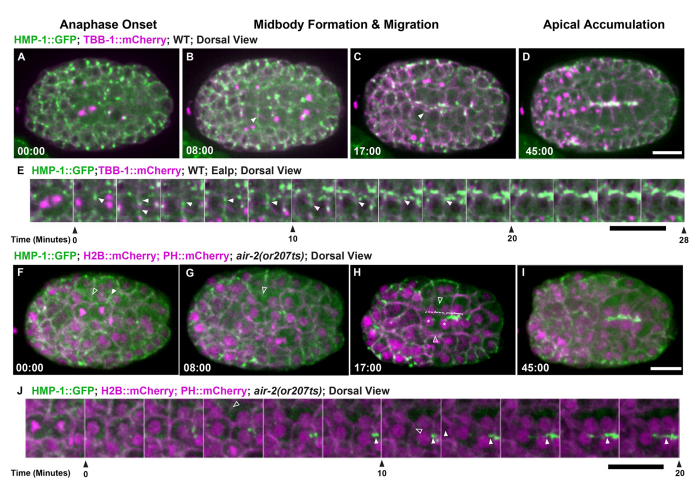


Fig. 4. Aurora B is required for adhesion dynamics during E8-E16 cytokinesis. Adhesion dynamics during E8-E16 division and polarization. (A-D) HMP-1:: GFP (green; microtubules in magenta) localizes to the furrow and midbody (B, arrowhead) during cytokinesis. HMP-1::GFP migrates with the midbody (C, arrowhead) to the apical surface where it accumulates after polarization (D). (E) Montage of HMP-1::GFP during E8-E16 division shows furrow and midbody (arrowheads) migration to the apical midline. (F-I) Aurora B mutants have reduced HMP-1::GFP on the furrow and midbody (F, unfilled arrowheads). When cells fail cytokinesis (G,H, unfilled arrowheads), HMP-1 accumulation is delayed (dashed bracket shows failed cytokinesis, solid bracket indicates successful E8 division with apical accumulation). Asterisks in H indicate nuclei. (I) HMP-1 signal eventually spreads along the midline. (J) Montage of HMP-1::GFP in Aurora B mutant E8-E16 cells that fail cytokinesis (unfilled arrowheads indicate furrow regression) and have delayed apical accumulation. Filled arrowheads indicate HMP-1 signal. Scale bars: 10 µm.

depletion caused lumens to be twice as wide on average $(2.53\pm0.33 \,\mu\text{m}, n=8, P<3.8\times10^{-7}; \text{Fig. 5F,G})$. The widest part of the lumen after AIR-2::GFP depletion was $2.94\pm0.41 \,\mu\text{m}$ ($P<3.33\times10^{-6}$), but only $1.27\pm0.03 \,\mu\text{m}$ in control embryos. Depletion of NMY-2::GFP ($1.05\pm0.17 \,\mu\text{m}, n=7$) or ZEN-4::GFP ($0.99\pm0.11 \,\mu\text{m}, n=4$) did not cause wide lumens, despite having cytokinesis failures (Fig. S6M,N). Therefore, tissue-specific depletion of Aurora B but not other cytokinetic regulators in the E8 gut leads to a highly consistent defect in the width of the lumen, consistent with a function for Aurora B at the apical surface.

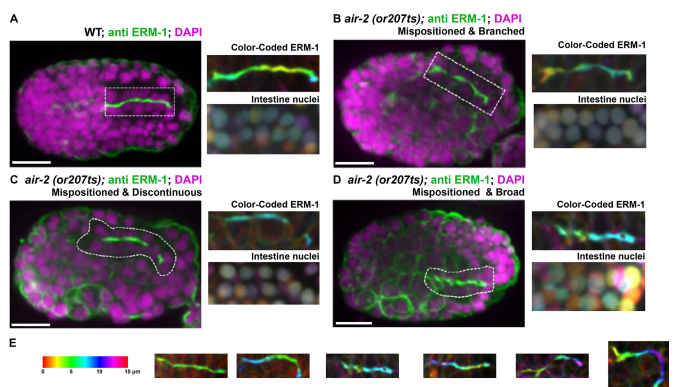
Table 1. Quantification of gut apical defects

	Mispositioned	Discontinuous	Broad	Branched
Wild type (N2)	0% (0/60)	0% (0/60)	3.3% (2/60)	20% (12/60)
air-2 (or207ts) zen-4 (or153ts) spd-1 (oj5ts)	60.9% (25/41) 27.3% (6/22) 0% (0/59)	43.9% (18/41) 27.3% (6/22) 28.8% (17/59)	12.1% (5/41) 81.8% (18/22) 10.2% (6/59)	65.9% (27/41) 81.8% (18/22) 40% (24/59)

Note that each embryo can exhibit more than one phenotype and can be included in more than one category.

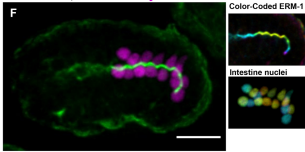
Apical midbody migration and AIR-2 apical localization in the pharynx

We also observed cytokinesis during the terminal divisions in the pharynx. The pharynx forms from more than 80 pharyngeal precursor cells (PPCs) and the final divisions occur 310-325 min after the first cleavage (Sulston et al., 1983). PPCs polarize and undergo apical constriction to form a lumen by 355 min (Rasmussen et al., 2013, 2012). We imaged from both dorsal and ventral aspects to observe PPC divisions and apical polarization (Fig. 6, Movie 8). We also used lattice light-sheet microscopy, which provides higher spatial resolution during the pharyngeal cell division (Movie 9). PPCs underwent symmetric furrowing that yielded a centrally placed midbody between daughter cells (Fig. 6A, F, Movie 8). PPC midbodies migrated from their central position between daughter cells toward the apical midline of the forming pharyngeal bulb (Fig. 6F,K,O, Movie 9). In PPC terminal divisions, AIR-2::GFP appeared on the spindle midzone, migrated with the midbody to the apical midline and persisted there (Fig. 6B-F, Movie 9). We confirmed apical localization using endogenously tagged AIR-2::GFP and immunofluorescence against endogenous AIR-2 (Fig. S1B,E). ZEN-4::GFP appeared on midbodies, migrated toward the apical surface, and rapidly disappeared (Fig. S7A-E,



	1-2 µm	3-4 µm	5-6 µm	7-8 µm	9-10 µm	11-12 µm
WT (N2)	25% (15/60)	40% (24/60)	31.7% (19/60)	3.3% (2/60)	0% (0/60)	0%(0/60)
air-2(or207ts)	0% (0/41)	22.0% (9/41)	12.2% (5/41)	34.1% (14/41)	14.6% (6/41)	17.1% (7/41)
zen-4 (or153ts)	0% (0/22)	27.3% (6/22)	40.9% (9/22)	22.7% (5/22)	4.5% (1/22)	0% (0/22)
spd-1 (oj5ts)	10.2 % (6/59)	55.9% (33/59)	27.1% (16/59)	1.7% (1/59)	1.7% (1/59)	0% (0/59)

Pelt-2::GFP degrader Anti ERM-1; H2B::mCherry



Pelt-2::GFP degrader; AIR-2::GFP Anti ERM-1; H2B::mCherry

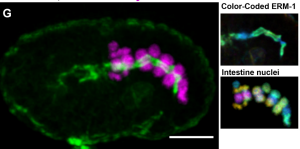


Fig. 5. Gut morphogenesis is disrupted in cytokinesis mutants. Apical surface staining after E8-E16 division and polarization. (A) ERM-1 apical staining (dashed rectangle) in wild-type (WT) bean-stage embryos. Maximum *z*-projected images of ERM-1 and nuclei color-coded according to *z*-depth (scale shown in F) show tissue organization. (B-D) In *air-2(or207)* embryos, apical surfaces are mispositioned (B-D), branched (B), contain gaps (C) or have broader staining (D). (E) Quantification of the defective apical *z*-plane distribution in different mutants (more colors indicate greater distortion in the *z*-plane). (F) ERM-1 staining and distribution of nuclei in a control embryo (GFP degrader only expressed) shows normal lumen width (1.15±0.11 μm, *n*=10) and nuclear distribution. (G) Endogenous AIR-2::GFP degradation in the intestine results in significantly broadened ERM-1 staining (2.53±0.33 μm, *n*=8) and disorganized nuclei. Scale bars: 10 μm.

Movie 8). NMY-2::GFP also labeled midbodies and moved to the apical surface, but remained there during apical constriction (Fig. 6G-K) (Rasmussen et al., 2012). RAB-11 and tubulin accumulated and remained localized to the apical surface after polarization (Fig. S7F,G). AIR-2 partially colocalized with PAR-6 and γ -tubulin::GFP (Toya et al., 2010) at the apical membrane (Fig. S7H,I). HMP-1::GFP localized to the furrow and midbody as it migrated to the

apical midline where it accumulated after polarization (Fig. 6L-O). Staining of embryos shifted slightly later in development revealed severe defects in pharyngeal formation in *air-2(or207)*, *zen-4(or153)* and *icp-1(or663ts)* mutant embryos (Fig. S6B-D,L). Therefore, similar patterns of symmetric furrowing, midbody migration and apical localization of AIR-2 are observed during epithelial polarization in the intestine and pharynx in *C. elegans*.

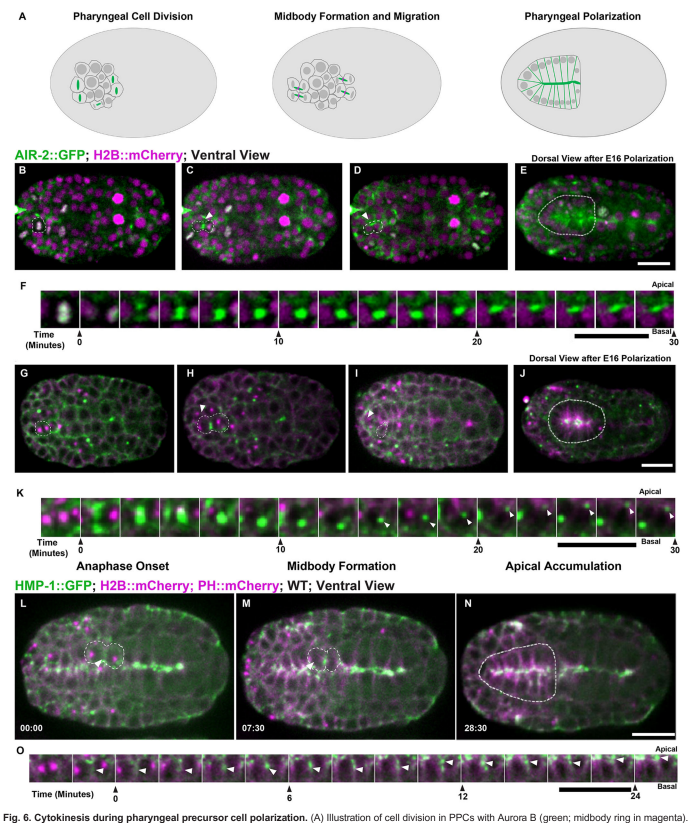


Fig. 6. Cytokinesis during pharyngeal precursor cell polarization. (A) Illustration of cell division in PPCs with Aurora B (green; midbody ring in magenta). (B-E) PPC division labeled with AIR-2::GFP (green; H2B::mCherry in magenta) from both ventral (B-D, dashed line highlights one cell, arrowhead indicates midbody) and dorsal (E, dashed line highlights pharynx) views. AIR-2::GFP localizes to chromosomes in metaphase (B), moves to the central spindle in anaphase (C), and appears on the midbody which moves toward the midline (D). AIR-2 persists at the pharyngeal apical surface for an extended time (E). (F) Montage showing AIR-2::GFP migrating toward the midline. (G-K) Imaging of NMY-2::GFP (green; TBB-1::mCherry in magenta), during midbody migration to the midline (I, K). NMY-2::GFP accumulates at the midline during apical constriction (J). (L-N) During PPC cytokinesis, α-catenin (HMP-1::GFP, green; tubulin in magenta) accumulates on the furrow, (arrowhead in L) and adjacent to the midbody (arrowhead, M) before accumulating at the midline (N). (O) Montage of HMP-1::GFP in PPC cell at the furrow, midbody and apical midline. Time shown in minutes:seconds. WT, wild type. Scale bars: 10 μm.

Apical midbody clustering and AIR-2 dendrite localization in sensilla neurons

Finally, we observed a unique form of cytokinesis in sensory neuron precursor divisions. The C. elegans amphid sensilla contain 12 neurons with dendrites that extend processes into the tip of the mouth. During morphogenesis, amphid neurons anchor at the tip of the animal and migrate back to extend dendrites (Heiman and Shaham, 2009). There are ten precursor cell divisions that occur between 280 and 400 min after the first cleavage to form the sensilla neurons (Sulston et al., 1983). Our observations show cytokinesis occurs in these sensilla precursor cells (SPCs) just before dendrite morphogenesis (Fig. 7A). SPCs underwent symmetrical furrowing with midbodies forming centrally between the daughter cells (Fig. 7B, Movies 10, 11). A group of at least six daughter cell pairs divided, initially forming multiple midbodies as observed with both confocal and lattice light-sheet imaging (Fig. 7C, Movies 10, 11). These midbodies migrated into a central cluster over a 60-min time window (Fig. 7D). AIR-2, RAB-11 and tubulin persisted in these clusters (Fig. 7D, Fig. S8A,B), whereas ZEN-4 rapidly disappeared, suggesting that abscission and MBR internalization had occurred (Fig. 7H, Movie 10). Cluster staining was observed with AIR-2 immunofluorescence or endogenous AIR-2::GFP (Fig. S1C,F). NMY-2::GFP migrated with the midbody to the cluster and persisted at the tip of the dendrites during extension (Fig. 7I, Movie 10). PAR-6 localized to this cluster, indicating that this site is the apical surface of these cells, which also accumulates γ -tubulin:: GFP (Fig. 7J, Fig. S7C,D). Recently, it was found that SPCs form a multicellular rosette with PAR-6 at the center, indicating that the midbody moves to the apical surface at the center of this rosette structure (Fan et al., 2019). HMP-1::GFP was observed at the furrow and midbody and migrated to the apical cluster and remained there during dendrite extension (Fig. 7E-G). Therefore, the midbody moves to the apical surface of the sensilla rosette where Aurora B and other midbody components accumulate.

After formation of the apical rosette, we observed that this region extends anteriorly toward the tip of the animal. AIR-2 and tubulin remained localized along the dendritic extension during elongation (Fig. 7K-M, Fig. S8D,E, Movie 12). As dendrites extended, other AIR-2 foci formed within the anterior region of the embryo and migrated toward the tip until six sensilla appeared (Fig. 7M, inset. Movies 12, 13). Although the individual cell divisions could not be easily discerned, these data suggest that other sensilla in the tip of the animal form through a similar process. Neuronal cell polarization may share mechanisms with epithelial morphogenesis (Low et al., 2019; McLachlan and Heiman, 2013), suggesting that modified cytokinesis may regulate epithelial polarization. Therefore, the midbody migrates from its original position at the end of furrowing to the apical surface in several tissues during morphogenesis. Interestingly, Aurora B remains localized at the apical surface well after completion of cytokinesis.

Aurora B has a post-mitotic function in dendrite formation

We tested whether Aurora B kinase and other cytokinesis components were required for sensilla formation. Cilia that form at the end of sensilla dendrites are exposed to the environment and can take up lipophilic dyes such as DiI (Hedgecock and White, 1985; Perkins et al., 1986). We inactivated cytokinesis mutants at different embryo stages and stained surviving L1 larvae with DiI. In wild type, amphid neurons were clearly labeled by DiI (Fig. 8A). In *air-2(or207)* mutants, we observed numerous defects in neuronal staining (Fig. 8B-E), including no DiI staining, indicating that none of the sensilla reached the environment (Table S2). *zen-4(or153)*

larvae showed severe DiI staining defects, which was dramatically reduced if embryos were shifted after the final divisions at the comma to 1.5-fold stage (Fig. S9A-C, Table S2). *spd-1(oj5)* animals had weak defects revealed by DiI staining but never showed a complete lack of staining (Fig. S9D, Table S2). Therefore, several cytokinetic regulators, including AIR-2, are required for dendrite formation.

Finally, we tested whether tissue-specific, post-mitotic depletion of AIR-2 would disrupt sensilla formation. We depleted endogenously tagged AIR-2::GFP from the sensory neurons by expressing the GFP degrader under the dyf-7 promoter (Fig. 8F), which is activated after the SPC divisions (Cheerambathur et al., 2019). AIR-2::GFP depletion in SPCs resulted in a wider distribution of cell bodies in L1 animals compared with control animals (Fig. 8G,H). This phenotype is also caused by the depletion of other kinetochore proteins, which also localize to the dendrite extension. Post-mitotic depletion of kinetochore proteins causes defects in the dendrite extension process, disrupts microtubule dynamics and also causes dispersion of the neuronal cell bodies (Cheerambathur et al., 2019). These data provide strong evidence that AIR-2 functions at the apical domain of the extending dendrite and has a role independent of its function during cytokinesis. Therefore, AIR-2 function is required for specialized cytokinesis during epithelial polarization and the subsequent formation of the apical surface during morphogenesis.

DISCUSSION

Our results reveal complex and reproducible patterns of cytokinesis during the invariant embryonic divisions in C. elegans. We observed reproducible alterations to furrow symmetry, central spindle length, abscission timing, midbody movement and MBR inheritance. We show that cells are completing cytokinesis when they polarize during morphogenesis. Aurora B inactivation disrupts cytokinesis and epithelial polarization. Many cells in the lineage divide and produce an apoptotic daughter cell rather than finish the divisions early, which may be due to the role of cytokinesis in polarization. Indeed, modified cytokinesis in the Q neuroblast generates a smaller daughter cell that undergoes apoptosis, which is prevented as cytokinesis parameters change (Ou et al., 2010). Cytokinesis is the transition into interphase and an ideal time to reorganize cellular architecture. Investigating the developmental plasticity of cytokinesis will be a fascinating question for future studies.

Asymmetric furrows drive efficient furrowing and help maintain proper cell contacts during cytokinesis (Founounou et al., 2013; Guillot and Lecuit, 2013; Maddox et al., 2007; Morais-de-Sá and Sunkel, 2013). Asymmetry of furrow ingression may contribute to the differential sensitivity to actin regulation in the four-cell embryo (Davies et al., 2018). The asymmetric furrow may facilitate MBR inheritance by EMS and it is worth noting that MS accumulates four MBRs, which could regulate its fate (Singh and Pohl, 2014). In polarized epithelial divisions, the furrow constricts to the apical surface to position the midbody (Herszterg et al., 2014). It is curious that cells furrow symmetrically before polarization, because asymmetric furrowing could apically position the midbody. Perhaps there is no cue to drive asymmetric furrowing to position the midbody at the apical midline before polarization. If polarization requires trafficking of apical components to the midbody, then the apical surface must form first and then be repositioned within the tissue to produce the final organization.

The apical migration of the midbody may represent a new phenomenon during cytokinesis. Midbody migration might occur in

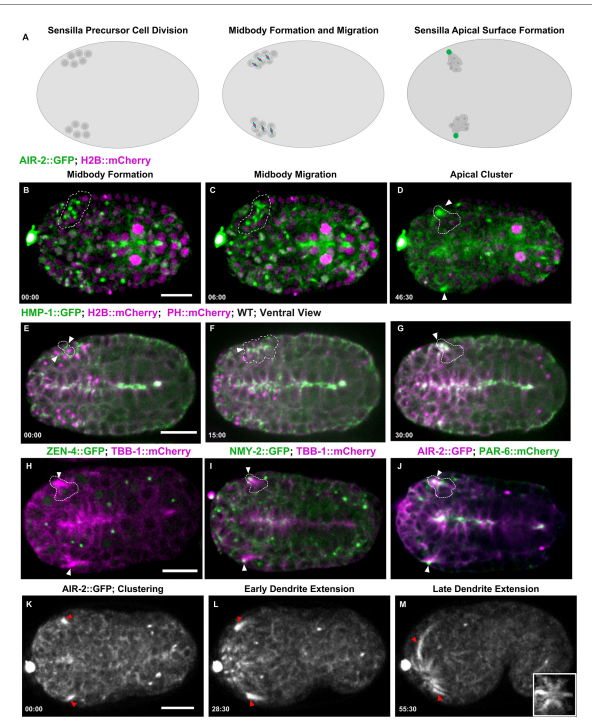


Fig. 7. Midbody components label dendrites of sensilla neurons. (A) Diagram of SPC divisions with Aurora B (green; midbody ring in magenta). (B-D) Cytokinesis in SPCs expressing AIR-2::GFP (green; H2B::Cherry in magenta) gives rise to multiple midbodies (dashed outline, B,C) that cluster together (arrowheads, D). (E-G) HMP-1::GFP accumulates at the furrow and midbody (arrowheads, E,F), accumulates at the apical cluster, and remains at the tip (G) during dendrite extension. (H) ZEN-4::GFP (green; microtubules in magenta) is internalized and degraded before the microtubule-rich cluster forms (arrowheads). (I) NMY-2::GFP (green; microtubules in magenta) remains at the tip of the dendrite as it extends (arrowheads). (J) PAR-6::mCherry (green) and AIR-2::GFP (magenta) colocalizes to the cluster (arrowheads), indicating that this is the apical surface. (K-M) AIR-2::GFP labels the dendrites during extension. Inset in M is a rotated maximum z-projection of sensilla after dendrite extension. Time shown in minutes:seconds. Scale bars: 10 μm.

other cells undergoing a mesenchyme-to-epithelial transition. Abscission is delayed during midbody migration indicating that daughter cells polarize while connected at the midbody, which might facilitate their reorganization. Indeed, SPD-1 inactivation causes disorganized polarization with AIR-2 reaching the apical surface through a compensatory mechanism. Midbody movements

are poorly understood and occur under normal or mutant conditions (Bernabé-Rubio et al., 2016; Herszterg et al., 2013; Morais-de-Sá and Sunkel, 2013; Singh and Pohl, 2014). Septate junction formation drives basal migration of the intercellular canal in *Drosophila* epithelia (Daniel et al., 2018; Wang et al., 2018). Midbody movements could also be regulated by global cortical

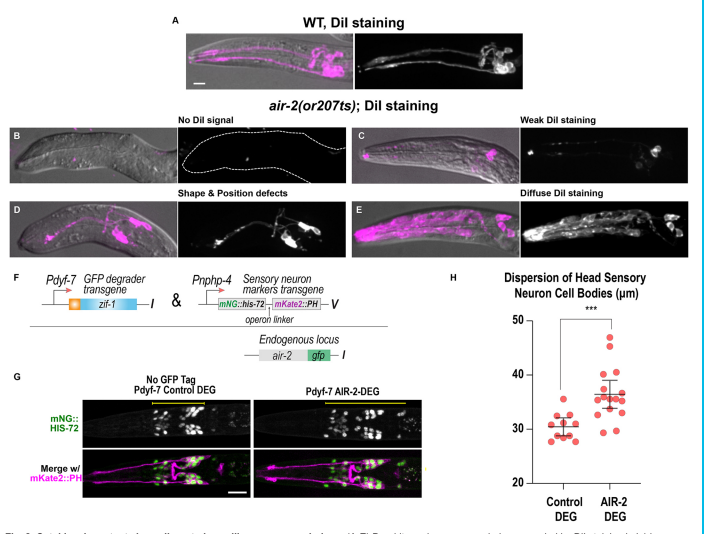


Fig. 8. Cytokinesis mutants have disrupted sensilla neuron morphology. (A-E) Dendrite and neuron morphology revealed by Dil staining in L1 larvae. (A) In wild type (WT), two dendrite bundles and amphid and phasmid neurons are labeled. (B-E) *air-2(or207ts)* mutants show no Dil signal (B), weak signal (C), dendrite shape and positioning defects (D) and diffuse staining throughout the head of the animal (E). Dashed line outlines indicate animal position in B. (F) Construct used for post-mitotic degradation of AIR-2::GFP in sensory neurons. (G) Sensory neuron nuclei (green) and plasma membranes (magenta) for the indicated conditions. Scale bars: 10 µm. (H) Quantification of sensory neuron cell body distribution, measured as indicated by the yellow lines in G. Error bars are the 95% confidence interval. ****P*<0.001 (two-tailed unpaired *t*-tests in GraphPad Prism; *n*=12 for control DEG and *n*=16 for AIR-2 DEG).

actin dynamics, which are regulated during cytokinesis (Jordan and Canman, 2012). In the future, it will be important to investigate how the midbody moves to the apical midline.

Cytokinesis in the intestinal lineage undergoes distinct regulatory phases during development. Post-embryonic intestinal divisions involve nuclear but not cytoplasmic divisions leading to the formation of binucleate cells that endoreduplicate to become polyploid (Hedgecock and White, 1985). Each division pattern might require gene expression programs that alter cytokinesis and produce unique proteins delivered to the midbody and apical surface. Transmembrane proteins localized to the tip of the dendrites in amphid sensilla could be delivered during cytokinesis (Heiman and Shaham, 2009). The initial secretory apparatus built during cytokinesis could contribute to the ability of the dendrite cilia to release exosomes (Wang et al., 2014a). Further investigation is required to define the contributions of the midbody to the apical surface in different tissues.

Aurora B regulates several aspects of the specialized cytokinesis that occurs during morphogenesis. Future work will be required to define how Aurora B moves from the midzone microtubules to the apical surface while the MBR is internalized. Aurora B regulates the cytoskeleton during cytokinesis to control cell shape and may have a similar role in lumen formation (Ferreira et al., 2013; Floyd et al., 2013; Goto et al., 2003; Kettenbach et al., 2011). Aurora B regulates abscission timing (Mathieu et al., 2013; Steigemann et al., 2009) and may delay abscission until after midbody migration. Aurora B delays abscission in mouse embryos to allow delivery of adhesion proteins to the midbody (Zenker et al., 2017). Aurora B regulates the central spindle (Bastos et al., 2013), and may contribute to the elongated midzone in E8-E16 gut cells. Aurora B is regulated by a cadherin in zebrafish embryos to organize spindle midzone microtubules (Chen et al., 2018), indicating reciprocal regulation between adhesion and cytokinesis. In the Xenopus embryo, altered spindle midzones correlate with changes to furrow ingression and midbody behavior (Kieserman et al., 2008). Although ZEN-4 is rapidly internalized with the midbody in the three tissues, it was previously implicated in morphogenesis of the epidermis and arcade cells independently of cytokinesis (Hardin et al., 2008; Portereiko et al., 2004; Von Stetina et al., 2017). Therefore, further study will be required to understand the role of Aurora B and the spindle midzone during specialized cytokinesis in morphogenesis.

Development (2020) 147, dev181099. doi:10.1242/dev.181099

In the sensilla, the centriole is required to form sensory cilia (Dammermann et al., 2009; Nechipurenko et al., 2017; Perkins et al., 1986). Central spindle proteins localize to cilia in *Xenopus* epithelial cells and regulate cilia morphology in the sensilla dendrites (Kieserman et al., 2008; Smith et al., 2011). Aurora B kinase regulates neuronal axon morphology and axonal outgrowth in zebrafish (Gwee et al., 2018). These results indicate a post-mitotic function for various cytokinetic regulators in cilia and cellular architecture. In the intestine, γ -tubulin but not centrioles remain at the apical surface, which forms microvilli rather than cilia (Feldman and Priess, 2012; Leung et al., 1999). Therefore, the midbody and centrosome may contribute different components to regulate the cytoskeletal architecture of the apical surface, which will be a major focus of future studies.

MATERIALS AND METHODS

C. elegans strains

C. elegans strains were maintained with standard protocols. *C. elegans* strains expressing midbody proteins driven by the *pie-1* promoter are listed in Table S3. All temperature-sensitive mutants were obtained from the *Caenorhabditis* Genetics Center.

CRISPR/Cas9 generation of GFP::AIR-2 and mScarlet::AIR-2 transgenic strains

We followed the CRISPR/Cas9 protocol generated by the Seydoux lab for C-terminus GFP and mScarlet tagging of the *C. elegans air-2* gene (Paix et al., 2015). The repair templates were amplified from pDD282 and pMS050, gifts from Bob Goldstein (Addgene plasmids #66823 and #91826, respectively). The primer sequences were as follows: AIR-2-mScarlet forward: GCAGCAAAAGATTGAAAAAGAAGCAAG TCTTCGAAATCACATGGTCTCCAAGGGAGAGG; AIR-2-mScarlet reverse: AGATGATTGAAAGAAGGACGGGAAAATCAGTAGTTGAT CACTTGTAGAGCTCGTCCATTCC; AIR-2-GFP forward: GCAGCAAA AGATTGAAAAAGAAGCAAGTCTTCGAAATCACGGAGCATCGGGA GCC; AIR-2-GFP reverse: AGATGATTGAAAGAAGGACGGGAAAA TCAGTAGTTGATCACTTGTAGAGCTCGTCCATTC. The guide RNA sequence is: UUGAAAAAGAAGCAAGUCUU.

Embryo preparation and imaging

For live imaging, young gravid hermaphrodites were dissected in M9 buffer containing polystyrene microspheres and sealed between two coverslips with petroleum jelly (Pohl and Bao, 2010). Live-cell imaging was performed on a spinning disk confocal system consisting of a Nikon Eclipse inverted microscope with a 60×1.40 NA objective, a CSU-22 spinning disk system, and a Photometrics EM-CCD camera from Visitech International. Images were acquired by Metamorph (Molecular Devices) and analyzed by ImageJ/Fiji Bio-Formats plugins (National Institutes of Health) (Linkert et al., 2010; Schindelin et al., 2012). Whole-embryo live imaging was performed on lattice light-sheet microscopes housed in the Eric Betzig lab, Bi-Chang Chen lab, or the Advanced Imaging Center at HHMI Janelia. The system is configured and operated as previously described (Chen et al., 2014). Briefly, embryos were dissected out and adhered to 5 mm round glass coverslips (Warner Instruments, CS-5R). Samples were illuminated by lattice light sheet using 488 nm or 560 nm diode lasers (MPB Communications) through an excitation objective (Special Optics, 0.65 NA, 3.74-mm WD). Fluorescent emission was collected by detection objective (Nikon, CFI Apo LWD 25XW, 1.1 NA) and detected by a sCMOS camera (Hamamatsu Orca Flash 4.0 v2). Acquired data were de-skewed as previously described (Chen et al., 2014) and deconvolved using an iterative Richardson-Lucy algorithm. Point-spread functions for deconvolution were experimentally measured using 200 nm TetraSpeck beads adhered to 5 mm glass coverslips (Invitrogen, T7280) for each excitation wavelength.

Immunostaining assay in C. elegans embryos

Apical marker staining was performed with the freeze-crack methanol protocol (Leung et al., 1999). Immunostaining with anti-AIR-2 antibodies

was performed as described (Schumacher et al., 1998). Primary antibodies and dilutions used were: anti-ERM-1 (1:200; AB_10584795, Developmental Studies Hybridoma Bank); P4A1/PAR-3 (1:200; P4A1, Developmental Studies Hybridoma Bank); DLG-1 (1:200; AB_2617529, Developmental Studies Hybridoma Bank); MH33 (1:150; MH33, Developmental Studies Hybridoma Bank); AIR-2 (1:50; gift from Jill Schumacher, University of Texas MD Anderson Cancer Center, TX, USA). Alexa 488- and 568-conjugated secondary antibodies (1:200-1:1000; A-11059 and A-11004, Invitrogen/Thermo Fisher Scientific) were used in the study. To stain temperature-sensitive mutants, two-cell-stage embryos were dissected from gravid worms, mounted in 10 μ l of M9 buffer, and kept cold on ice. The two-cell-stage embryos were incubated at 15°C for 4-7 h until specific stages, then shifted to the restrictive temperature (25°C) for 2-4 h and stained as described above.

Dil staining in C. elegans

DiI staining of wild type and temperature-sensitive mutants was performed as previously described (Tong and Burglin, 2010). Two-cell-stage embryos were incubated at 15°C for 6.5-7 h until they reached the polarized E16 stage, then shifted to the restrictive temperature (25° C) with 1:200 dilution of stock DiI dye solution containing 2 mg/ml DiI in dimethyl formamide for 18-24 h. Hatched larvae were transferred to M9 and washed twice in M9 before mounting in 25 mM levamisole on 2% agar pads for imaging.

Temperature-shift experiments

Temperature-sensitive mutants were maintained at 15°C. To perform temperature shifts on staged embryos, gravid adults were transferred to a dissection chamber (<4°C), which was precooled in an ice bucket with 20 µl of ice-cold M9 Buffer. Two-cell-stage embryos were quickly transferred (within a 5-10 min time window) via mouth pipette (Aspirator tube assemblies, Sigma-Aldrich) to Fisherbrand Hanging Drop Slides (12-560B, Fisher Scientific) on ice. The slide was placed into a humidified chamber and incubated at 15°C until the appropriate stages were reached, then shifted to 26°C. Incubation times were determined based on C. elegans embryonic lineage timing and adjusted according to DAPI staining to ensure each mutant was shifted at a similar stage of embryo development. To inactivate air-2(or207), mutant embryos were incubated for 5 h at 15°C and shifted to 26°C for 3 h to reach the bean stage or for 5 h at 26°C to reach the comma stage. This was the minimum amount of time required to shift embryos to the non-permissive temperature and observe significant cytokinesis defects by the E8-E16 division, indicating significant reduction of AIR-2 function. Most embryos reached the E4-E8 division at the time of the shift. By live imaging we found that there was little disruption of the E4-E8 division under these conditions as air-2(or207) embryos (n=4/5) have eight normal E8 cells. N2, spd-1(oj5) and zen-4(or153) embryos were incubated for 4.5 h at 15°C to reach the E4-E8 stage, followed by 3 h at 26°C to reach the bean stage and 5 h at 26°C to reach the comma stage. To shift embryos at the comma stage, air-2(or207) embryos were incubated for 12 h at 15°C and N2, spd-1(oj5) and zen-4(or153) embryos were incubated for 11-11.5 h at 15°C.

Acknowledgements

Lattice light-sheet microscopy was performed at the Advanced Imaging Center at HHMI Janelia Research Campus, a facility jointly supported by the Gordon and Betty Moore Foundation and the Howard Hughes Medical Institute. CGC and Wormbase provided *C. elegans* strains and information, funded by the NIH (P40 OD010440) and NHGRI (U41 HG002223). We thank Martha Soto, Julie Ahringer, Arshad Desai, Karen Oegema, Barth Grant, Tony Hyman and Asako Sugimoto for sharing strains and members of the Bembenek laboratory for support. We also thank Dr Max Heiman, Dr Zhirong Bao, Dr Amy Maddox, Dr Pablo Lara-Gonzalez and Sophia Hirsch for discussions, and Dr Don Fox, Dr John White, Dr John Heddleston, Dr Heidi Hehnley-Chang, Lindsay Rathbun and Erica Colicino for manuscript feedback.

Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: X.B., M.M., C.G.T., D.C., J.N.B.; Methodology: M.M., C.G.T., D.S., C.-Y.C., W.-C.W., P.-Y.L., E.B., B.-C.C., D.C., J.N.B.; Software: W.-C.W.,

P.-Y.L., B.-C.C.; Validation: X.B., M.M., C.G.T., J.R.S., D.C., J.N.B.; Formal analysis:
X.B., M.M., C.G.T., D.S., J.R.S., B.N., L.R.K., D.C., J.N.B.; Investigation: X.B., M.M.,
C.G.T., D.S., C.-Y.C., J.R.S., B.N., D.M., L.R.K., D.C., J.N.B.; Resources: D.S.,
D.M., L.R.K., N.M., E.B., B.-C.C., D.C., J.N.B.; Data curation: X.B., M.M., D.S.,
W.-C.W., P.-Y.L., J.R.S., B.N., D.M., L.R.K., D.C., J.N.B.; Writing - original draft: X.B.,
J.N.B.; Writing - review & editing: M.M., C.G.S.T., D.S., D.C., J.N.B.; Visualization:
X.B., M.M., C.G.S.T., D.S., C.-Y.C., W.-C.W., P.-Y.L., J.R.S., B.N., E.B., B.-C.C.,
D.C., J.N.B.; Supervision: E.B., B.-C.C., D.C., J.N.B.; Project administration:
E.B., B.-C.C., D.C., J.N.B.; Funding acquisition: E.B., B.-C.C., D.C., J.N.B.

Funding

Funding was provided by the Ministry of Science and Technology, Taiwan (105-2119-M-001-026-MY2 to B.-C.C.), the Howard Hughes Medical Institute (E.B.), the Wellcome Trust (208833 and 203149 to D.C.), and the National Institutes of Health (R01 GM114471 to J.N.B.). Deposited in PMC for immediate release.

Supplementary information

Supplementary information available online at http://dev.biologists.org/lookup/doi/10.1242/dev.181099.supplemental

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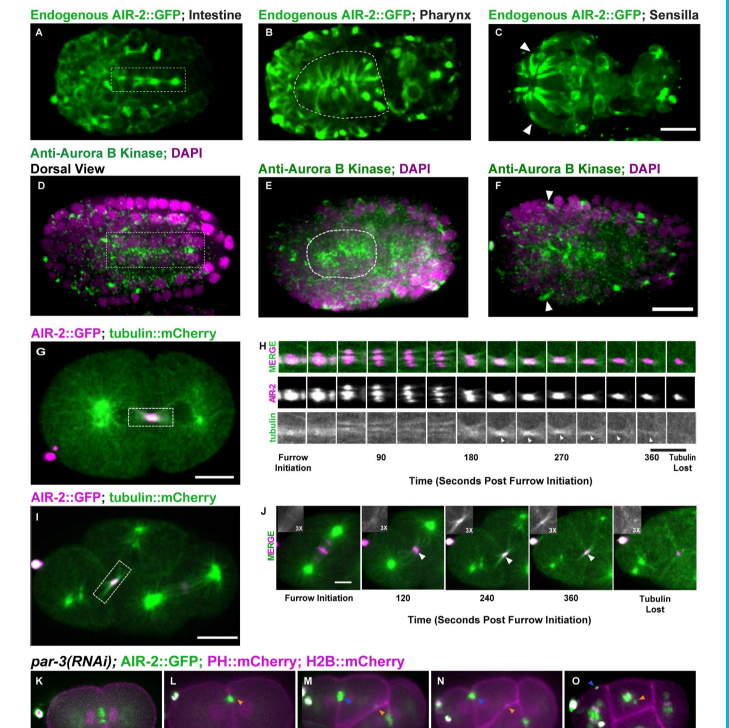
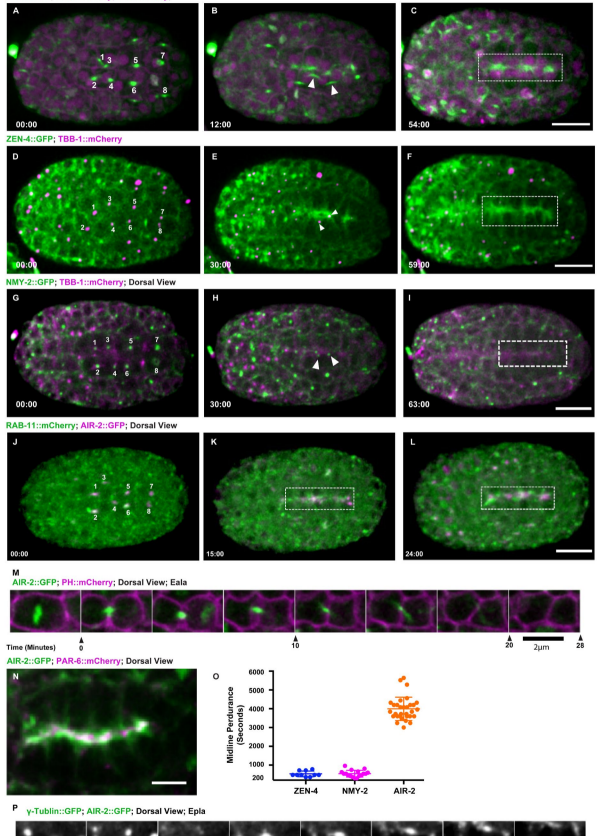


Figure S1. Aurora B kinase during embryo development.

Endogenously tagged AIR-2::GFP shows apical localization in E16 intestinal cells (dashed box in A), polarized pharyngeal cells (dashed shape in B), and elongating sensilla neurons (arrowheads, C). (D) AIR-

2 staining also shows apical surface of polarized E16 intestine (D, rectangle), pharynx (E, dotted circle) and at the apical cluster of the amphid sensilla (F, arrowheads). (G-H) AIR-2::GFP (magenta) and tubulin::mCherry (green) colocalize at the central spindle during cytokinesis in the first cell division. AIR-2::GFP persists at the midbody remnant after microtubules are lost, which indicates rapid abscission timing (H). (I-J) AIR-2::GFP (magenta) and tubulin::mCherry (green, inset) colocalize on the AB central spindle and midbody (arrowhead). Microtubules quickly disappear indicating abscission (J). (K-O) In *par-3(RNAi)* embryos, the P0 furrow becomes more asymmetric causing offset midbody positioning (L, orange arrowhead). The AB furrow is less asymmetric and ingresses in the opposite direction (M, blue arrowhead indicates misplaced AB midbody). (O) Midbody remnants are inherited randomly. Scale bar, 10 μm.



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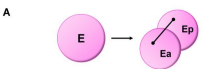
AIR-2::GFP; H2B::mCherry; PH::mCherry; Dorsal View

A0

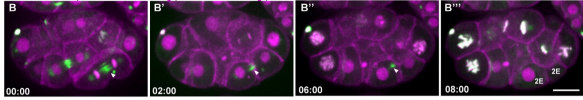
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Figure S2. Cytokinesis in the intestine epithelia

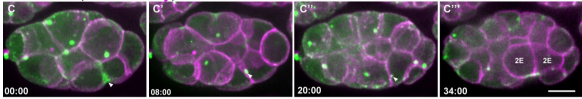
Cytokinesis in the E8-E16 division. (A-C) AIR-2::GFP (green, H2B::mCherry and PH::mCherry are magenta) localizes to midbodies (A, numbered 1-8) that migrate toward the apical midline (B). AIR-2:GFP remains at the apical surface after polarization (C). (D-F) ZEN-4::GFP (magenta, TBB-1::mCherry in green) appears on midbodies (D, numbered 1-8) that migrate to the midline (E) and are quickly removed (F, rectangle box). (G-I) NMY-2 (green, microtubules in magenta) localizes to midbody rings (labeled 1-8 in G) that move to the midline (arrowheads, H) but do not persist (rectangle box in I). (J-L) RAB-11::mCherry (green) and AIR-2::GFP (magenta) colocalize on midbodies (labeled as 1-8 in J) as they migrate to the midline (K) and persist well after polarization is complete (L, rectangle). (M) Midbody from Eala at the E8-E16 division migrates towards the midline but the AIR-2 signal diminishes (green, PH::mCherry in magenta). (N) PAR-6::mCherry (magenta) colocalizes with AIR-2::GFP (green) at the apical midline. (O) Quantification of midline perdurance of different midbody components (measured from the end of furrowing to internalization or loss of signal) shows that AIR-2 remains at the apical surface after the midbody remnant is internalized. Error bars represent the standard deviation. (P) Image series showing the simultaneous movement of AIR-2::GFP (red arrowhead) on the midbody and ytubulin::GFP (white arrowheads) on centrosomes to the apical surface. Time shown in minutes: seconds. Scale bar, 10 µm.

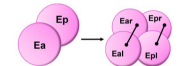


AIR-2::GFP; H2B::mCherry; PH::mCherry; 1E-2E



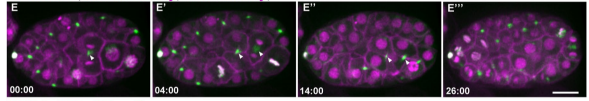
NMY-2::GFP; PH::mCherry; 1E-2E



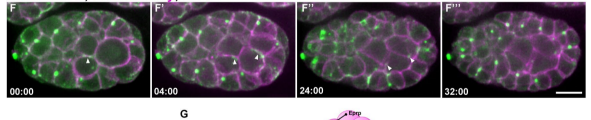


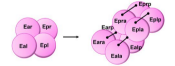
AIR-2::GFP; H2B::mCherry; PH::mCherry; 2E-4E

D

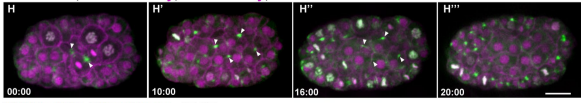


NMY-2::GFP; PH::mCherry; 2E-4E





AIR-2::GFP; H2B::mCherry; PH::mCherry; 4E-8E



NMY-2::GFP; PH::mCherry; 4E-8E

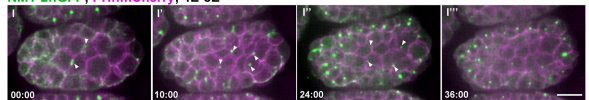
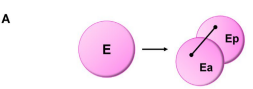
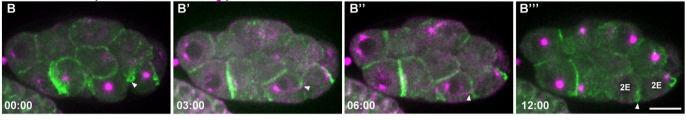


Figure S3. Midbody dynamics during divisions of the early E lineage

(A) Diagram of the E-E2 division. Dynamics of Aurora B (B) and NMY-2 (C) show formation of the midbody (arrowheads) and rapid internalization. A similar pattern is also observed during the E2-E4 divisions (D-F) and the E4-E8 divisions (G-I), demonstrating normal midbody dynamics in the first three E lineage embryonic divisions. Scale bar, 10 μm.

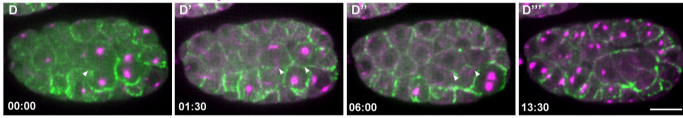


HMP-1::GFP; TBB-1::mCherry; 1E-2E



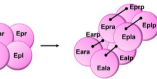
Ea Ep Ear Epr Eal Epi

HMP-1::GFP; TBB-1::mCherry; 2E-4E



Е

С



HMP-1::GFP; TBB-1::mCherry; 4E-8E

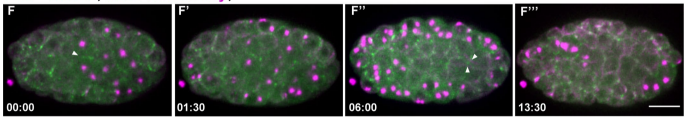
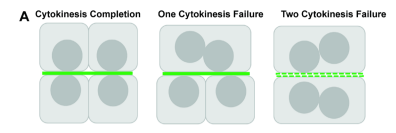


Figure S4. Localization of α-catenin during the E lineage cell divisions

(A) Diagram of the E-E2 division. HMP-1::GFP localizes to the furrow and membrane adjacent to the midbody during cytokinesis (arrowheads) and remains there after spindle midzone microtubules disappear.
(C) Diagram of the E2-E4 division. HMP-1::GFP can be observed in the cortex of the dividing cells (arrowhead, D) and localizes along the furrow membrane and adjacent to the midbody (arrowheads, D-D"). (E) Diagram of the E4-E8 divisions. (F) Arrowheads indicate accumulation of HMP-1::GFP near the midbody during cytokinesis. HMP-1::GFP does not accumulate at the midline until the end of the E8-E16 division. Scale bar, 10 μm.

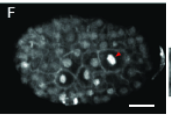


Defects in Microtubule Accumulation

В	Genotype	Total Embryo	Total 8E Cells	Cytokinesis Failure	Nuclei Polarization Defects	Chromosome Segregation Defects	Microtubule Accumulation Defects
	WT (N2)	8	34	0/34 (0%)	0/34 (0%)	0/34 (0%)	0/8 (0%)
	air-2(or207ts)	11	66	18/66 (27.3%)	17/66 (25.8%)	3/66 (4.5%)	5/11 (45.5%)
С	Genotype	Pair of 8E Cells	Pair Number	One Cytokinesis Failure	Microtubule Discontinuous	Two Cytokinesis Failure	Microtubule Discontinuous
	WT (N2)	Ea(r/l)a	8	0/8 (0%)	0/0 (0%)	0/8 (0%)	0/0 (0%)
		Ea(r/l)p	8	0/8 (0%)	0/0 (0%)	0/8 (0%)	0/0 (0%)
		Ep(r/l)a	8	0/8 (0%)	0/0 (0%)	0/8 (0%)	0/0 (0%)
	air-2(or207ts)	Ea(r/l)a	11	5/11 (45.5%)	1/5 (20%)	0/11 (0%)	0/0 (0%)
		Ea(r/l)p	11	6/11 (54.4%)	0/6 (0%)	0/11 (0%)	0/0 (0%)
		Ep(r/l)a	11	3/11 (27.3%)	1/3 (33.3%)	2/11 (18.2%)	1/2 (50.0%)

Defects in Adhesion Complex Accumulation

Genotype	e Total Embryo	Total 8E Cells	Cytokinesis Failure	Nuclei Polarization Defects	Chromosome Segregation Defects	HMP-1 Accumulation Defects
WT (N2)	5	30	0/30 (0%)	0/30 (0%)	0/30 (0%)	0/5 (0%)
air-2(or207	ts) 12	72	30/72 (41.7%)	25/72 (34.7%)	0/72 (0%)	6/12 (50.0%)
Genotype	Pair of 8E Cells	Pair Number	One Cytokinesis Failure	s HMP-1 Discontinuous	Two Cytokinesis Failure	HMP-1 Discontinuous
WT (N2)	Ea(r/l)a	5	0/5 (0%)	0/0 (0%)	0/5 (0%)	0/0 (0%)
	Ea(r/l)p	5	0/5 (0%)	0/0 (0%)	0/5 (0%)	0/0 (0%)
	Ep(r/l)a	5	0/5 (0%)	0/0 (0%)	0/5 (0%)	0/0 (0%)
air-2(or207ts	s) Ea(r/l)a	12	3/12 (45.5%)	0/3 (0%)	3/12 (0%)	3/3 (100%)
	Ea(r/l)p	12	6/12 (54.4%)	0/6 (0%)	1/12 (8.3%)	0/1 (0%)
	Ep(r/l)a	12	4/12 (33.3%)	0/4 (0%)	4/12 (33.3%)	3/4 (75.0%)



H2B::mCherry; PH::mCherry; air-2(or207ts)



Figure S5. Quantification of defects observed in *air-2(or207ts)* mutant embryos

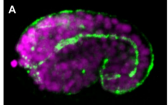
(A) Diagram of failure patterns of pairs of E8-E16 divisions on opposite sides of midline and observed consequence on apical microtubule accumulation (green). (B) Quantification of defects in *air-2(or207ts)* mutant E8-E16 divisions showing significant cytokinesis failures where most cells that failed also have nuclear polarization defects. Of the 11 embryos observed, five have apical microtubule accumulation defects. The low incidence of chromosome segregation defects are consistent with weak Aurora B inactivation. (C) Quantification of individual cell division failures and incidence of discontinuous microtubule accumulation at the apical midline after polarization. (D) Quantification of defects in *air-2(or207ts)* mutant E8-E16 divisions showing significant defects in HMP-1::GFP accumulation with cytokinesis failures. (E) Quantification of individual cell division failures and incidence of discontinuous adhesion accumulation at the apical midline after polarization. (F) An example of an intestinal cell in an Aurora B mutant embryo showing lagging chromosome segregation (red arrowheads).

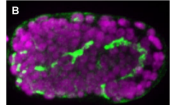
N2; Anti ERM-1; DAPI



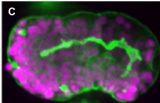
zen-4(or153); Anti ERM-1; DAPI Branched & Broad lumen

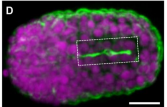




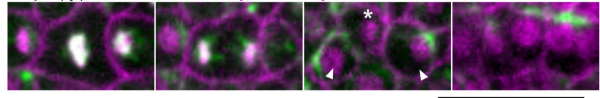


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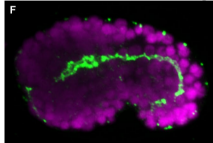




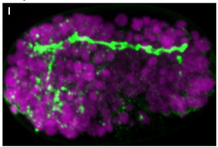
E spd-1(oj5); AIR-2::GFP; H2B::mCherry; PH::mCherry



WT; Anti PAR-3; DAPI; Comma stage WT; Anti DLG-1; DAPI; Comma stage WT; Anti IFB-2; DAPI; Comma stage

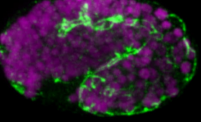


air-2(or207); Anti PAR-3; DAPI Mispositioned & Branched

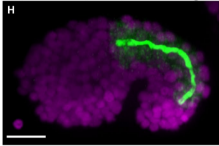


icp-1(ts); Anti ERM-1; DAPI

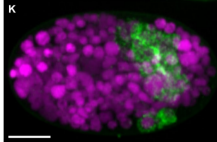




Pelt-2::GFP degrader; NMY-2::GFP; Anti ERM-1; pelt-2::histone::mKate2



air-2(or207); Anti IFB-2; DAPI Mispositioned & Discontinous



Pelt-2::GFP degrader; ZEN-4::GFP Anti ERM-1; pelt-2::histone::mKate2

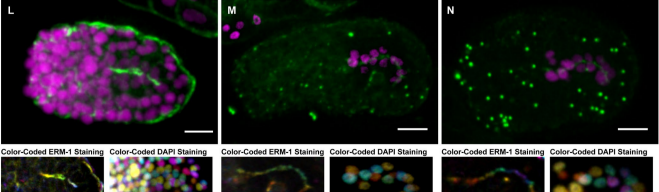
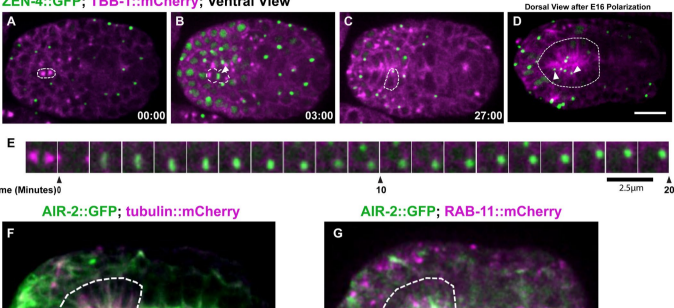


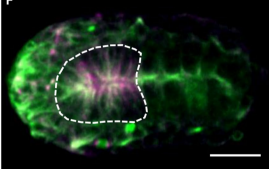
Figure S6. Cytokinesis mutants have disrupted intestinal and pharyngeal tubulogenesis

(A) ERM-1 stains the apical lumen of the gut and pharynx in wild-type (N2) embryos at the comma stage. Gut lumen defects are observed in (B) *air-2(or207)*, (C) *zen-4(or153)* and (D) *spd-1(oj5)* embryos. (E) In *spd-1(oj5)* embryos, AIR-2::GFP does not accumulate at the spindle midzone and instead appears on spindle poles (arrowheads). A neighboring cell comes between E16 sisters (asterisk), but AIR-2 eventually moves to the apical midline with the poles. (F) PAR-3, (G) DLG-1 and (H) IFB-2 localize to the lumen in N2 embryos. (I-K) In *air-2(or207)* mutants, apical surface markers localize to distorted and mispositioned lumens. (L) Lumen defects are observed in *icp-1(or663ts)* mutant embryos. (M-N) Tissue-specific degradation of endogenously tagged NMY-2::GFP (M) or ZEN-4::GFP (N) in the gut does not cause significant widening of the apical lumen. Scale bar, 10 μm.

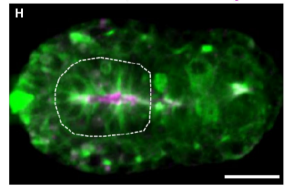
ZEN-4::GFP; TBB-1::mCherry; Ventral View



Time (Minutes)0



AIR-2::GFP; PAR-6::mCherry



α-TUBULIN::mCherry; v-TUBULIN::GFP: H2B::GFP

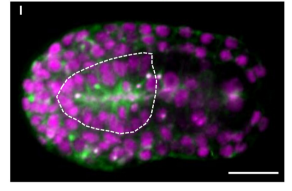


Figure S7. Cytokinesis in Pharynx Precursor Cells

(A-E) Centralspindlin ZEN-4::GFP dynamics in PPC divisions. Midbody remnants labeled with ZEN-4::GFP migrate to the midline (arrowheads, C) and are rapidly internalized and eventually degraded (arrowheads in D show internalized ZEN-4 labeled midbodies). AIR-2::GFP (F, G green) colocalized with Tubulin::mCherry (F, magenta) and partially colocalized with RAB-11::mCherry (G, magenta) at the apical surface of the polarized pharynx (dotted circle). AIR-2::GFP (H, green) partially co-localized with PAR-6::mCherry (H, magenta) at the apical surface of the pharynx (dotted circle). (I) The apical surface of the pharynx accumulates γ -tubulin::GFP (magenta in merge, microtubules in green). Scale bar, 10 μ m.

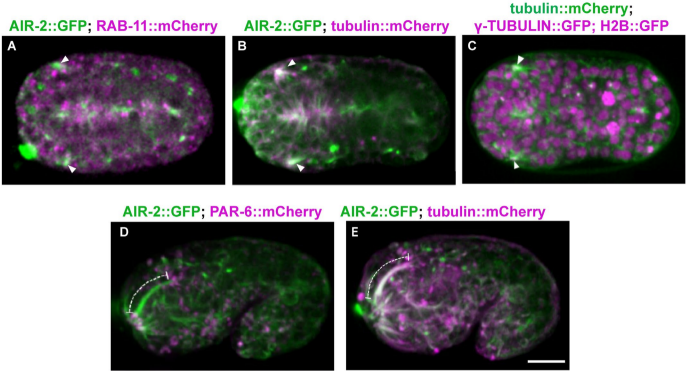


Figure S8. Midbody components label dendrites of sensilla neurons

Aurora B kinase (green) colocalizes with (A) RAB-11::mCherry (magenta) and (B) tubulin::mCherry (magenta) at the apical surface of sensilla (white arrowheads). (C) γ-tubulin::GFP (magenta) localizes at the apical cluster with microtubules (green). (D) AIR-2::GFP (green) localizes along the dendrite extension (dashed line) and PAR-6::mCherry is observed at the tip of the extension (arrowhead), indicating it is the apical domain. (E) AIR-2::GFP (green) and microtubules (tubulin::mCherry, magenta) colocalize along the length of the extended dendrite (dashed line). Scale bar, 10 μm.

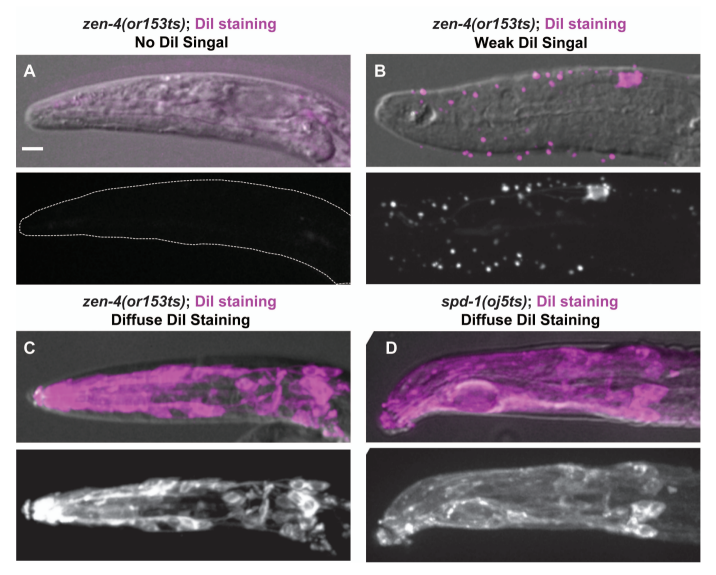
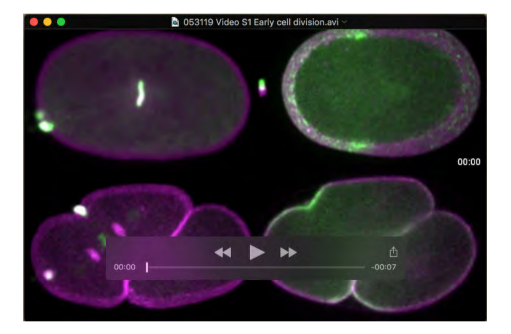


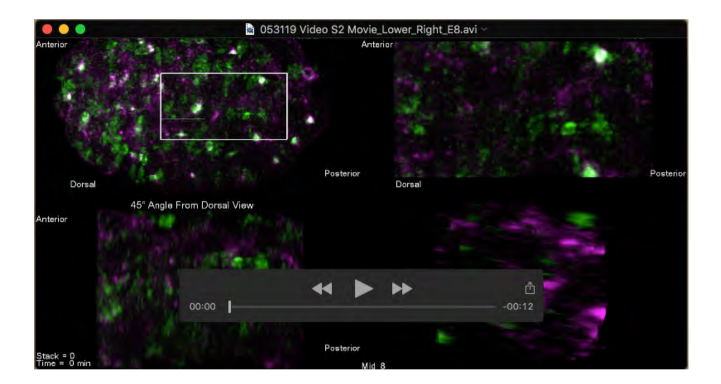
Figure S9. Cytokinesis mutants have disrupted sensilla neuron morphogenesis

Visualizing dendrite and neuron morphology by DiI staining in surviving larvae at non-permissive temperature. (A-C) Hatched *zen-4(or153ts)* mutant larvae display a variety of defects including No-DiI signal (A), weak (B) and Diffuse (C) DiI signal. *spd-1(oj5)* mutant embryo also show sensilla neuron morphogenesis defects including diffuse DiI staining pattern (D).



Movie 1. Cytokinesis in the first two mitotic divisions

Cytokinesis in the first (top row) or second (bottom row) division in embryos expressing AIR-2::GFP (green, left, with H2B::mCherry and PH::mCherry in magenta) or NMY-2::GFP (green, right, with PH::mCherry in magenta). White arrowheads indicate first midbody that is internalized by AB and red arrowheads indicate the AB midbody, which is engulfed by EMS. Images are maximum Z projections of 15 central planes spaced 1µm apart taken every 90 seconds. Playback rate is 2 frames/second.



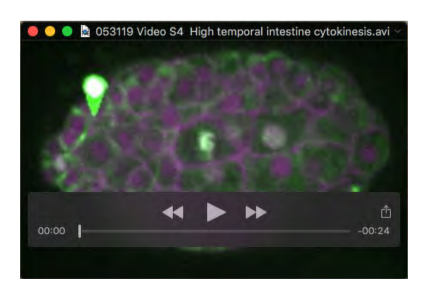
Movie 2. Lattice Light Sheet Imaging of Intestinal Cytokinesis

E8-E16 intestinal cell divisions in embryos expressing AIR-2::mScarlet (green) with NMY-2::GFP (magenta) imaged with lattice light sheet microscopy. A ventral view of the whole embryo (top left) is shown with a region of interests (white box) highlighted in the top right panel. Rotated views shown a dorsal view (bottom right) and a 45 degrees from a dorsal view (bottom left) and show that AIR-2::GFP accumulates to a narrow band around the emerging apical surface. Images are maximum Z projections of images acquired every 60 seconds.



Movie 3. Cytokinesis in the intestine epithelia

E8-E16 intestinal cell division in embryos expressing AIR-2::GFP (green, left, PH::mCherry in magenta), NMY-2::GFP (green, middle, tubulin::mCherry in magenta) or ZEN-4::GFP (green, right, tubulin::mCherry in magenta). The midbodies (indicated by arrowheads) form and move toward the apical surface. ZEN-4::GFP and NMY-2::GFP rapidly disappear, while AIR-2::GFP persists at the apical midline. Images are maximum Z projections of 10 planes spaced 1µm apart, taken every 60 seconds. Playback rate is 6 frames/second.



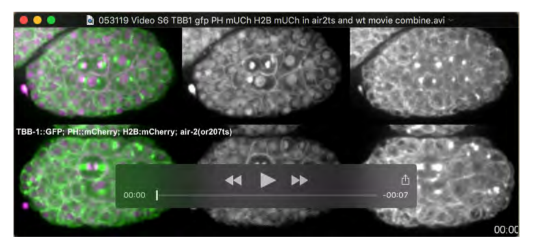
Movie 4. High temporal resolution of midbody dynamics in the intestine

Imaging Earp cell division with high temporal resolution in an embryo expressing AIR-2::GFP (green, PH::mCherry in magenta) shows the lengthening of the central spindle and midbody migration event. Single plane images were acquired every 10 seconds. Playback rate is 15 frames/second.



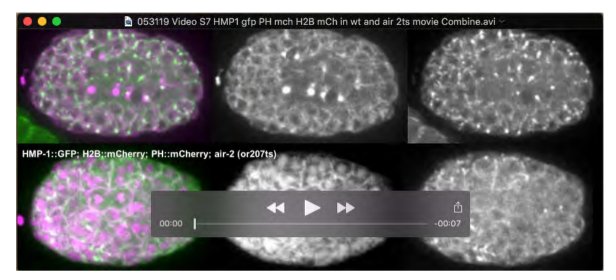
Movie 5. Cytokinesis in E16 to E20 cell division

E16-E20 cell division in embryos expressing AIR-2::GFP (green) with PH::mCherry and H2B::mCherry (magenta). Images are Maximum Z projections of 15 z planes 1 µm apart that were acquired every 90 seconds. Playback rate is 6 frames/second.



Movie 6. Microtubule dynamics during E8-E16 in Aurora B mutants.

WT (top row) and *air-2(or207)* (bottom row) mutant embryos expressing TBB-1::GFP (green in merge, right movie) and PH::mCherry and H2B::mCherry (middle, magenta in merge). Spindle midzone microtubules in WT embryos move to the midline where microtubules accumulate. In Aurora B mutant embryos, spindle midzone microtubules are diminished and cells that fail cytokinesis (right pair of gut cells) delay microtubule accumulation compared with the cells that do not fail. Images are maximum Z projections of 10-15 z planes 1 µm apart that were acquired every 60 seconds. Playback rate is 6 frames/second.



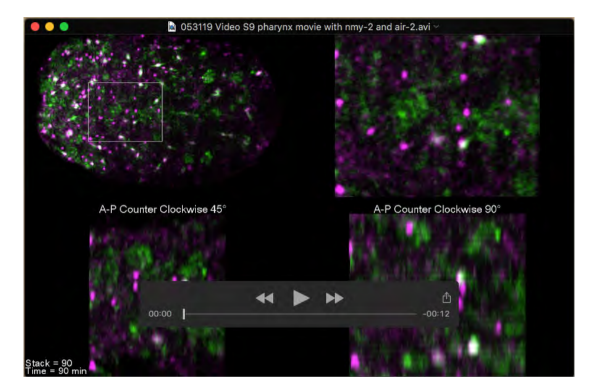
Movie 7. Aurora B regulates adhesion dynamics during E8-E16 epithelial polarization.

WT (top row) and *air-2(or207)* (bottom row) mutant embryos expressing HMP-1::GFP (green in merge, right movie) and TBB-1::mCherry (top middle, magenta in upper left merge) or PH::mCherry and H2B::mCherry (bottom middle, magenta in lower left merge). In WT, HMP-1::GFP accumulates at the furrow and midbody as it migrates to the apical surface where it accumulates during polarization. In Aurora B mutant embryos, HMP-1::GFP is reduced in the furrow and midbody and cells that fail cytokinesis (left pair of gut cells) delay adhesion accumulation at the midline. Images are maximum Z projections of 10-15 z planes 1 µm apart that were acquired every 60 seconds. Playback rate is 6 frames/second.



Movie 8. Cytokinesis in the pharynx from ventral views.

Cell division in pharyngeal precursor cells in embryos expressing AIR-2::GFP (green, left, H2B::mCherry in magenta), ZEN-4::GFP (green, middle left, TBB-1::mCherry in magenta), HMP-1::GFP (green, middle right, TBB-1::mCherry in magenta) or NMY-2::GFP (green, right, TBB-1::mCherry in magenta). Midbodies (white arrowheads) migrate toward pharyngeal midline after forming centrally between daughter cell pairs. Images are maximum Z projections of 10-15 z planes 1 µm apart that were acquired every 90 seconds. Playback rate is 6 frames/second.



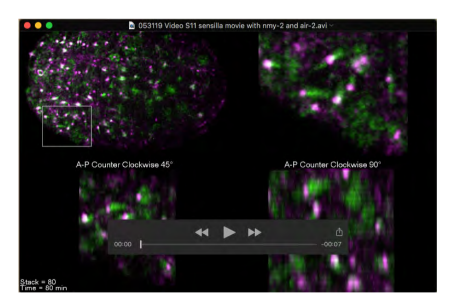
Movie 9. Lattice light sheet imaging of pharyngeal precursor cell divisions

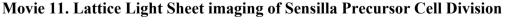
Dorsal view (top left) of lattice light sheet microscopy of pharyngeal precursor cell cytokinesis in embryos expression AIR-2::mScarlet (green) with NMY-2::GFP (magenta) shows migration of midbodies towards the apical midline. The pharyngeal precursor cells in the highlighted region (top right) are shown in zoomed in view of a 45-degree (bottom left) and 90-degree (bottom right) rotation along the anterior-posterior axis of the embryo. Images are max intensity projections of images acquired every 60 seconds.



Movie 10. Cytokinesis in the sensilla dendrite development

Cell division in sensilla precursor cell divisions in embryos expressing AIR-2::GFP (green, left, H2B::mCherry in magenta), ZEN-4::GFP (green, middle left, TBB-1::mCherry in magenta), NMY-2::GFP (green, middle right, TBB-1::mCherry in magenta), or HMP-1::GFP (green, right, TBB-1::mCherry in magenta). Midbodies (white arrowheads) migrate into an apical cluster. Images are maximum Z projections of 10 z planes 1 µm apart that were acquired every 90 seconds. Playback rate is 6 frames/second.



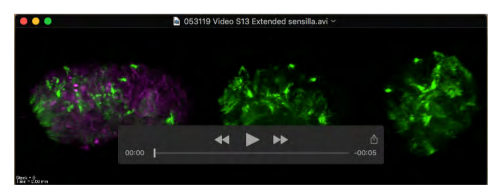


Dorsal view of embryos expressing midbody flank marker AIR-2::mScarlet (green) with NMY-2::GFP (magenta) during the sensilla precursor divisions (top left). The sensilla precursor cells are highlighted in the white box (top right) and zoomed in views of this region at a 45-degree (bottom left) and 90-degree (bottom right) rotation along the anterior-posterior axis of the embryo are also shown. Images are maximum intensity Z projections of images acquired every 60 seconds.



Movie 12. Dendrite Extension of sensilla neurons

Embryo expressing AIR-2::GFP from the apical cluster stage of amphid polarization through the process of dendrite extension. Images are maximum Z projections of 7 planes 1 µm apart viewed from the ventral aspect and acquired every 90 seconds. Playback rate is 6 frames/second.





Embryo expressing AIR-2::mScarlet (green) with NMY-2::GFP (magenta) during amphid neuron dendrite extension. A 45 degree (middle panel) and a 90 degree (right panel) rotated view of the mouth of the animal are also shown. Images are maximum Z projections of images acquired every 60 seconds.

Stage Before	Genotype	Hatch Rate % (Hatch Embryos/Total)
Shifting		
15 °C Forever	N2	100% (32/32)
	air-2(or207)	53.6% (37/69)
	zen-4(or153)	100% (28/28)
	spd-1(oj5)	100% (35/35)
E4-E8	N2	100% (26/26)
	air-2(or207)	6.3% (2/32)
	zen-4(or153)	0% (0/57)
	spd-1(oj5)	100% (48/48)
E8-E16	N2	100% (45/45)
	air-2(or207)	14.4% (13/90)
	zen-4(or153)	10.1% (10/99)
	spd-1(oj5)	100% (83/83)
Comma-1.5 Fold	N2	100% (36/36)
	air-2(or207)	33.7% (31/92)
	zen-4(or153)	85.7% (54/63)
	spd-1(oj5)	100% (27/27)

Table S1. Hatch rate of temperature sensitive mutants dissected at the two cell stage.

Table S2. Quantification of Dil Staining of TS Mutants

Stage Before	Genotype	No Dil Signal	Weak Dil	Shape & Position	Extended Dil Staining
Shifting	-	_	signal	Defect	
15 °C Forever	N2	0% (0/32)	0% (0/32)	0% (0/32)	0% (0/32)
	air-2(or207)	2.7% (1/37)	8.1% (3/37)	2.7% (1/37)	0% (0/37)
	zen-4(or153)	0% (0/18)	0% (0/18)	0% (0/18)	0% (0/18)
	spd-1(oj5)	0% (0/34)	0% (0/34)	0% (0/34)	0% (0/34)
E4-E8	N2	0% (0/10)	0% (0/10)	0% (0/10)	0% (0/10)
	air-2(or207)	50% (1/2)	0% (0/2)	50% (1/2)	0% (0/2)
	zen-4(or153)	0% (0/0)	0% (0/0)	0% (0/0)	0% (0/0)
	spd-1(oj5)	0% (0/44)	4.5% (2/44)	9.1% (4/44)	6.8% (3/44)
E8-E16	N2	0% (0/29)	0% (0/29)	0% (0/29)	0% (0/29)
	air-2(or207)	16.7% (2/12)	58.3% (7/12)	58.3% (7/12)	0% (0/12)
	zen-4(or153)	77.8% (7/9)	11.1% (1/9)	22.2% (2/9)	0% (0/9)
	spd-1(oj5)	0% (0/59)	0% (0/59)	8.5% (5/59)	5.1% (3/59)
Comma-1.5 Fold	N2	0% (0/21)	0% (0/21)	0% (0/21)	0% (0/21)
	air-2(or207)	6.9% (2/29)	10.3% (3/29)	0% (0/29)	13.8% (4/29)
	zen-4(or153)	0% (0/53)	0% (0/53)	0% (0/53)	22.6% (12/53)
	spd-1(oj5)	0% (0/18)	0% (0/18)	0% (0/18)	33.3% (6/18)

Table S3. Strains used in this study.

Strain	Genotype	Origin
AZ212	unc-119(ed3) iii; ruIs32[Ppi-1::GFP::His-58] iii	(Praitis et al., 2001)
DKC21	ltSi1016[pDC337; Pdyf-7::vhhGFP4::ZIF- 1::dyf-7_3'UTR; cb-unc-119(+)] air-2(lt58[air- 2::GFP::tev::loxP::3xFlag]) i;Pnphp- 4::mNeonGreen-his-72:tbb-2_3'UTR;;gpd-2/3 operon linker-mKate2-PH:unc-34_3'UTR] v	This Study
EKM48	unc-119(ed3) iii; ojIs51[Ppie-1::GFP::air-2; unc-119(+)]	(Bembenek et al., 2013)
EKM50	unc-119(ed3) iii; ojIs51[Ppie-1::GFP::air-2; unc-119(+)]; ltIs37[(pAA64) pie- 1p::mCherry::his-58 + unc-119(+)] iv; ltIs44[pie-1p::mCherry::PH(PLC1delta1) + unc-119(+)] v	(Bembenek et al., 2013)
EKM51	unc-119(ed3) iii; ojIs51[Ppie-1::GFP::air-2; unc-119(+)]; ltIs37[(pAA64) pie- 1p::mCherry::his-58 + unc-119(+)] iv	This Study
EKM52	unc-119(ed3) iii; ojIs51[Ppie-1::GFP::air-2; unc-119(+)];ltIs44[pie- 1p::mCherry::PH(PLC1delta1) + unc-119(+)] v	This Study
EU630	air-2(or207) i	(Severson et al., 2000)
EU716	zen-4(or153) iv	(Severson et al., 2000)
JA1559	unc-119(ed3) iii; weIs21[pJA138 (pie- 1::mCherry::tub::pie-1)]	(Lee et al., 2015)
JAB23	unc-119(ed3) iii; ojIs51[Ppie-1::GFP::air-2; unc-119(+)]; weIs21[pJA138 (pie- 1::mCherry::tub)]	This Study

JAB24	zen-4(or153ts) iv; xsEx6[zen-4::GFP; rol-6	This Study
	(su1006)]; unc-119(ed3) iii; weIs21[pJA138	
	(pie-1::mCherry::tub)]	
JAB32	unc-119(ed3) iii; ddIs26[pie-1p::mCherry::par-	This Study
	6; unc-119(+)] v; ojIs51[Ppie-1::GFP::air-2;	
	unc-119(+)]	
JAB34	unc-119(ed3) iii; zen-4(or153) iv; xsEx6[zen-	This Study
	4::GFP; rol-6 (su1006)]; ltIs44[pie-	
	<i>lp::mCherry::PH(PLC1delta1) + unc-119(+)]</i>	
	ν	
JAB36	unc-119(ed3) iii; ddIs26[pie-1p::mCherry::par-	This Study
	6; unc-119(+)] v; ojIs51 [Ppie-1::GFP::air-2 +	
	unc-119(+)]	
JAB38	unc-119(ed3) iii; air-2(or207); ltIs44[pie-	This Study
	<i>lp::mCherry::PH(PLC1delta1) + unc-119(+)]</i>	
	v; ojIs37[Ppie-1::H2B::mCherry; unc-119(+)]	
	iv; ojIs2[alpha-tubulin::GFP unc-119(+)]	
JAB39	unc-119(ed3) III; ojIs51[Ppie-1::GFP::air-2 +	This Study
	unc-119(+)]; ruIs32[Ppi-1::GFP::His-58; unc-	
	119(ed3) iii; ddIs6[tbg-1::GFP + unc-119(+)]	
	V	
JAB52	unc-119(ed3) iii; ddIs6[tbg-1::GFP + unc-	This Study
	119(+)] v; ruIs32[Ppi-1::GFP::His-58; unc-	
	119(ed3) iii; weIs21[pJA138 (pie-	
	1::mCherry::tub)]	
JAB60	unc-119(ed3) iii; ojIs51[Ppie-1::GFP::air-2;	This Study
	unc-119(+)];	
	11]	
JAB116	unc-119(ed3) iii; weIs21[pJA138 (Ppie-	This Study
	1::mCherry::tub)]; unc-119(+)]; zuIs45[nmy-	

	2::NMY-2::GFP; unc-119(+)] v	
JAB142	unc-119(ed3) iii; ojIs2[alpha-tubulin::GFP unc-119(+)]; ltIs37[(pAA64) pie- 1p::mCherry::his-58 + unc-119(+)] iv; ltIs44[pie-1p::mCherry::PH(PLC1delta1) + unc-119(+)] v	This Study
JAB194	air-2(erb80[air-2::mScarlet]) i	This Study
JAB200	<pre>hmp-1(cp20[hmp-1::gfp + LoxP unc-119(+) LoxP]) v; weIs21[pJA138 (pie- 1::mCherry::tub::pie-1)]</pre>	This Study
JAB205	air-2(erb-81[air-2::linker::GFP])	This Study
JAB207	unc-119(ed3) iii; air-2(or207) i; hmp- 1(cp20[hmp-1::gfp + LoxP unc-119(+) LoxP]) v; ltIs44[pie-1p::mCherry::PH(PLC1delta1) + unc-119(+)] v; ojIs37[Ppie-1::H2B::mCherry; unc-119(+)] iv	This Study
JAB210	air-2(erb80[air-2::mScarlet) i; zuIs45[nmy- 2::NMY-2::GFP; unc-119(+)] v	This Study
JAB223	unc-119(ed3) iii; air-2(erb-81[air- 2::linker::GFP]); ltSi910[pOD2044/pSW378; Pelt-2::vhhGFP4::ZIF-1::operon- linker::mCherry::histone::tbb-2_3'UTR; cb- unc-119(+)] ii	This Study
JAB224	unc-119(ed3) iii; zen-4(lt30[GFP::loxP::zen- 4]) iv; ltSi910[pOD2044/pSW378; Pelt- 2::vhhGFP4::ZIF-1::operon- linker::mCherry::histone::tbb-2_3'UTR; cb- unc-119(+)] ii	This Study
JAB225	unc-119(ed3) iii; nmy-2(cp13[nmy-2::GFP + LoxP]) i; ltSi910[pOD2044/pSW378; Pelt-	This Study

	2::vhhGFP4::ZIF-1::operon-	
	linker::mCherry::histone::tbb-2_3'UTR; cb-	
	unc-119(+)] ii	
JAB235	<i>spd-1(oj5) I; air-2(erb-81[air-2::linker::GFP]);</i> <i>ltIs37[(pAA64) pie-1p::mCherry::his-58 + unc- 119(+)] iv; ltIs44[pie- 1p::mCherry::PH(PLC1delta1) + unc-119(+)]</i> <i>v</i>	This Study
JCC401	icp-1(or663ts) i	(Davies et al., 2014)
JJ1473	unc119(ed3) iii; zuIs45[nmy-2p::nmy-2::GFP;	(Nance et al., 2003)
	unc-119(+)] v	
LP162	nmy-2(cp12[nmy-2::GFP + LoxP] i	(Dickinson et al., 2013)
LP169	unc-119(ed3) iii; hmp-1(cp21[hmp-1::GFP _	(Marston et al., 2016)
	LoxP unc-119(+) LoxP]) v	
MG170	zen-4(or153) iv; xsEx6[zen-4::GFP + rol-	(Kaitna et al., 2000)
	6(su1006)	
N2	Bristol (wild-type)	CGC
OD56	unc-119(ed3) iii; ltIs37[(pAA64) pie-	(McNally et al., 2006)
	<i>1p::mCherry::his-58 + unc-119(+)] iv</i>	
OD70	unc-119(ed3) iii; ltIs44[pie-	(Kachur et al., 2008)
	<i>lp::mCherry::PH(PLC1delta1) + unc-119(+)]</i>	
	v	
NWG002	unc-119(ed3) iii; ltIs44[pie-	(Redemann et al., 2010)
	<i>1p::mCherry::PH(PLC1delta1) + unc-119(+)]</i>	
	<i>v; zuIs45[nmy-2p::nmy-2::GFP; unc-119(+) v</i>	
OD2768	unc-119(ed3) iii; ltSi910[pOD2044/pSW378;	(Wang et al., 2017)
	Pelt-2::vhhGFP4::ZIF-1::operon-	
	linker::mCherry::histone::tbb-2_3'UTR; cb-	
	unc-119(+)] ii	
	1	l

OD2979	zen-4(lt30[GFP::loxP::zen-4]) iv	(Lee et al., 2018)
OD3025	unc-119(ed3) iii; ltSi1016[pDC337; Pdyf- 7::vhhGFP4::ZIF-1::dyf-7_3'UTR; cb-unc- 119(+)] i #2	(Cheerambathur et al., 2019)
OD3230	air-2(lt58[air-2::GFP::tev::loxP::3xFlag]) i	(Cheerambathur et al., 2019)
OD3262	<i>ltSi1016[pDC337; Pdyf-7::vhhGFP4::ZIF-</i> <i>1::dyf-7_3'UTR; cb-unc-119(+)] air-2(lt58[air-</i> <i>2::GFP::tev::loxP::3xFlag]) i</i>	(Cheerambathur et al., 2019)
OD3919	unc-119(ed3) iii; ltSi1174[oxTi365; pDC591; Pnphp-4::mNeonGreen-his-72: tbb- 2_3'UTR;;gpd-2/3 operon linker-mKate2- PH:unc-34_3'UTR] v	(Cheerambathur et al., 2019)
RT1196	unc-119(ed3) iii; pwIs476[Ppie- 1:mCherry:RAB-11.1, unc-119(+)]	Gift from Barth Grant
SA240	unc-119(ed3) iii; tjIs54[pie-1 promoter- gfp::tbb-2; pie-1 promoter-2xmCherry::tgb-1; unc-119 ⁺]	(Toya et al., 2010)
SA245	unc-119(ed3) iii; tjIs57[pie-1 promoter- mCherry::his-48; unc-119 ⁺]	(Toya et al., 2010)
SA250	unc-119(ed3) iii; tjIs54[pie-1 promoter- gfp::tbb-2; pie-1 promoter-2xmCherry::tgb-1; unc-119 ⁺]; tjIs57[pie-1 promoter-mCherry::his- 48; unc-119 ⁺]	(Toya et al., 2010)
TH27	unc-119(ed3) iii; ddIs6[tbg-1::GFP + unc- 119(+)] v	(Redemann et al., 2010)
TH110	unc-119(ed3) iii; ddIs26[pie-1p::mCherry::par- 6; unc-119(+)]	(Schonegg et al., 2007)
WH12	spd-1(oj5) i	(O'Connell et al., 1998)
WH210	unc-119(ed3) iii; ojIs2[alpha-tubulin::GFP unc-119(+)]	(Kemp et al., 2004)

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