Epidermal PAR-6 and PKC-3 are essential for postembryonic development of *Caenorhabditis elegans* and control non-centrosomal microtubule organization

Victoria G. Castiglioni¹, Helena R. Pires¹, Rodrigo Rosas Bertolini¹, Amalia Riga¹, Jana Kerver¹, Mike Boxem¹

¹Corresponding author. Email: m.boxem@uu.nl (MB)

¹These authors contributed equally to this work.

¹Division of Developmental Biology, Institute of Biodynamics and Biocomplexity, Department of Biology, Faculty of Science, Utrecht University, Utrecht, The Netherlands

Short title: Essential roles for PAR-6 and PKC-3 in larval *C. elegans* epidermal epithelia

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Summary

Using inducible protein degradation, we show that PAR-6 and PKC-3/aPKC are essential for postembryonic development of *C. elegans* and control the organization of non-centrosomal microtubule bundles in the epidermis, likely through recruitment of NOCA-1/Ninein.

Abstract

The cortical polarity regulators PAR-6, PKC-3 and PAR-3 are essential for the polarization of a broad variety of cell types in multicellular animals, from the first asymmetric division of the *C. elegans* zygote to apical–basal polarization of epithelial cells. In *C. elegans*, the roles of the PAR proteins in embryonic development have been extensively studied, yet little is known about their functions during larval development. Using auxin-inducible protein depletion, we here show that PAR-6 and PKC-3, but not PAR-3, are essential for postembryonic development. We also demonstrate that PAR-6 and PKC-3 are required in the epidermal epithelium to support animal growth and molting, and the proper timing and pattern of seam cell divisions. Finally, we uncovered a novel role for PAR-6 in controlling the organization of non-centrosomal microtubule arrays in the epidermis. PAR-6 was required for the localization of the microtubule organizer NOCA-1/Ninein, and microtubule defects in a *noca-1* mutant are highly similar to those caused by epidermal PAR-6 depletion. As NOCA-1 physically interacts with PAR-6, we propose that PAR-6 promotes non-centrosomal microtubule organization through localization of NOCA-1/Ninein.
Introduction

Polarity is a near universal property of cells that is essential for establishing proper cellular architecture and function. Epithelial cells – one of the major polarized animal cell types – polarize along an apical–basal axis and establish molecularly and functionally distinct apical, basal, and lateral membrane domains. The boundary between apical and lateral domains is marked by the presence of cell–cell junctions that provide adhesion between cells and prevent unwanted paracellular passage of molecules. The polarization of epithelial cells is orchestrated by conserved cortical polarity regulators that establish opposing membrane domains through mutually antagonistic interactions. In metazoans, the partitioning defective (PAR) proteins Par3, Par6, and atypical protein kinase C (aPKC) play a central role in the establishment of epithelial cell polarity. These highly conserved polarity regulators are essential determinants of apical domain identity, and are required for the positioning, maturation, and maintenance of apical cell junctions (Achilleos et al., 2010; Franz and Riechmann, 2010; Georgiou et al., 2008; Harris and Peifer, 2004; Harris and Peifer, 2005; Harris and Tepass, 2008; Hutterer et al., 2004; Izumi et al., 1998; Joberty et al., 2000; Leibfried et al., 2008; Lin et al., 2000; Totong et al., 2007; Wodarz et al., 2000; Yamanaka et al., 2001).

Par6 and Par3 are both PDZ domain containing scaffold proteins that can interact with each other, with aPKC, and with numerous other proteins. Par6 and aPKC form a stable subcomplex by interacting through their PB1 domains (Hirano et al., 2005; Wilson et al., 2003). The association of Par6–aPKC with Par3 is dynamic. In C. elegans zygotes, PAR-6/PKC-3 shuttle between a kinase inactive complex with PAR-3 that promotes anterior segregation, and an active complex with the small GTPase CDC-42 (Aceto et al., 2006; Beers and Kemphues, 2006; Rodriguez et al., 2017; Wang et al., 2017). In epithelia, Par3 can promote the apical recruitment of Par6–aPKC (Franz and Riechmann, 2010; Harris and Peifer, 2005; Hutterer et al., 2004; Joberty et al., 2000; Lin et al., 2000; Wodarz et al., 2000). In mature epithelia, however, the bulk of Par3 segregates to the apical/lateral border, where it plays an essential role in the positioning and assembly of apical junctions (Achilleos et al., 2010; Georgiou et al., 2008; Harris and Peifer, 2004; Harris and Tepass, 2008; Izumi et al., 1998; Leibfried et al., 2008; Totong et al., 2007; Yamanaka et al., 2001). The release of Par3 in epithelia depends on phosphorylation of Par3 by aPKC, and involves handoff of Par6–aPKC to Cdc42 and the epithelial specific Crumbs polarity complex (Bilder et al., 2003; Harris and Peifer, 2005; Hong et al., 2003; Krahn et al., 2010; Morais-de-Sá et al., 2010; Nagai-Tamai et al., 2002; Nunes de Almeida et al., 2019; Walther and Pichaud, 2010).

In addition to interactions that mediate the subcellular localization of Par6–aPKC or Par3, both Par6 and Par3 can interact with effector proteins to connect cortical polarity with downstream pathways (McCaffrey and Macara, 2009). For example, Par3 modulates phospholipid levels by recruiting the lipid phosphatase PTEN to cell junctions (Feng et al., 2008; Martin-Belmonte et al., 2007; Pinal et al., 2006; von Stein et al., 2005), inhibits Rac activity by binding to and inactivating the RacGEF Tiam1 (Chen and Macara, 2005; Mertens et al., 2005), and mediates spindle positioning in Drosophila neuroblasts through recruitment of Insuteable (Schober et al., 1999; Wodarz et al., 1999). For Par6, fewer downstream targets have been described. In mammals, Par6 can recruit the E3 ubiquitin ligase Smurf1 to promote degradation of the small GTPase RhoA, causing dissolution of tight junctions.
Par6 can also bind to the nucleotide exchange factor ECT2 to regulate epithelial polarization and control actin assembly at metaphase in dividing epithelial cells (Liu et al., 2004; Liu et al., 2006; Rosa et al., 2015). As high-throughput studies have identified multiple candidate binding partners that have not yet been investigated (Boxem et al., 2008; Brajenovic et al., 2004; Giot et al., 2003; Grossmann et al., 2015; Hein et al., 2015; Huttlin et al., 2015; Lenfant et al., 2010; Luck et al., 2020; Waaijers et al., 2016), additional interactions important for the functioning of Par6 and for linking cortical polarity to other processes involved in epithelial polarization likely remain to be discovered.

Despite the conserved requirements for Par6–aPKC and Par3 in epithelial cells there are important context and cell-type dependent differences in the functioning of these polarity proteins (Pickett et al., 2019; St Johnston, 2018). For example, in Drosophila, Bazooka (Par3) is not required for junction positioning or polarization of cells in the follicular epithelium (Pickett et al., 2019; Shahab et al., 2015), and in the adult Drosophila midgut, the canonical Par, Crumbs, and Scribble polarity modules are all dispensable for apical–basal polarity (Chen et al., 2018). In *C. elegans*, requirements for PAR-3 and PAR-6 in embryonic epithelia also vary. PAR-6 appears to be required for apical junction formation in all epithelia, including the epidermis, intestine, foregut, and pharyngeal arcade cells (Montoyo-Rosario et al., 2020; Totong et al., 2007; Von Stetina and Mango, 2015; Von Stetina et al., 2017). However, while arcade cells show a complete lack of polarization upon PAR-6 loss, foregut, intestinal, and epidermal epithelial cells still establish an apical domain (Totong et al., 2007; Von Stetina and Mango, 2015). PAR-3 is required for apical junction formation in embryonic intestinal and pharyngeal epithelia, but not in epidermal epithelial cells (Achilleos et al., 2010).

Studies of PAR-6, PKC-3, and PAR-3 in *C. elegans* have largely focused on embryonic tissues. Here, we make use of targeted protein degradation to investigate the role of PAR-6, PKC-3, and PAR-3 in larval epithelia of *C. elegans*. Ubiquitous depletion of PAR-6 and PKC-3, but not PAR-3, resulted in a larval growth arrest, demonstrating that these proteins are required for larval development. Through tissue-specific depletion, we identified an essential role for PAR-6 and PKC-3 in the *C. elegans* epidermis. Depletion in this tissue caused growth arrest, a cessation of the molting cycle, and severe defects in the division pattern of the epidermal seam cells. We also observed defects in the maintenance of apical cell junctions, and a failure to exclude LGL-1 from the apical domain. Finally, we identified a novel role for PAR-6 in organizing non-centrosomal microtubule arrays in the epidermis. Epidermal depletion of PAR-6 led to defects in the localization of the microtubule organizer NOCA-1/Ninein, as well as of the γ-tubulin ring complex component GIP-1, and of the sole Patronin/CAMSAP/Nezha homolog PTRN-1. Microtubule defects in a *noca-1* mutant closely resembled those in PAR-6 depleted animals, including the loss of GIP-1 localization. As NOCA-1 physically interacts with PAR-6, we conclude that PAR-6 likely organizes non-centrosomal microtubule arrays through localization of NOCA-1.
Results

PAR-6 and PKC-3 are essential for larval development

To investigate the role of PAR-6, PKC-3, and PAR-3 in larval development, we made use of the auxin-inducible degradation (AID) system. The AID system enables targeted degradation of AID-degron tagged proteins through expression of the plant-derived auxin-dependent E3 ubiquitin ligase specificity factor TIR1 (Zhang et al., 2015) (Fig. 1A). Using CRISPR/Cas9, we inserted sequences encoding the AID-degron and the green fluorescent protein (GFP) into the endogenous par-6, pkc-3, and par-3 loci, such that all known isoforms of each protein are tagged (Fig. 1B). PAR-6 was tagged at the shared C-terminus, and PKC-3 at the N-terminus. The par-3 locus encodes two groups of splice variants that use two alternative start sites; hence we inserted the GFP–AID tag at both start sites. To examine if the presence of the GFP–AID tags interfered with protein function, we examined the growth rates of the tagged strains. Homozygous animals were viable and showed the same growth rates as wild-type, indicating that the proteins are still functional (Fig. 1C–E). We also observed localization of each protein at the apical membrane domain of epithelial tissues, including the pharynx, excretory canal, intestine and epidermis (Fig. 1F, G). This localization pattern matches previous observations in C. elegans larvae (Li et al., 2010a; Li et al., 2010b), and further indicates that the GFP–AID tag does not interfere with protein functioning.

To investigate the role of PAR-3, PAR-6 and PKC-3 in larval development we degraded each protein using a ubiquitously expressed TIR1 under the control of the eft-3 promoter (Zhang et al., 2015). We tested the efficiency of protein degradation by exposing synchronized L3 larvae to auxin and examining protein expression. Apical enrichment of PAR-3, PAR-6, and PKC-3 became indistinguishable from background fluorescence within one hour of exposure to 4 mM auxin in the pharynx, excretory canal, intestine, and epidermis (Fig. 1G). To examine if the depletion of PAR-6, PKC-3, or PAR-3 affected larval development, we degraded each protein by addition of auxin at hatching and measured animal growth rates. Ubiquitous degradation of PAR-3 did not cause a defect in larval growth, and animals developed into morphologically normal looking and fertile adults (Fig. 1E). This lack of a visible phenotype may indicate that the functions of PAR-3 are not essential for larval development. Alternatively, despite visual absence of GFP::AID::PAR-3, degradation may be incomplete, or animals may express unpredicted non-tagged protein isoforms. In contrast to PAR-3, depletion of PAR-6 or PKC-3 caused a striking growth arrest with animals not developing beyond L1 size (Fig. 1C, D). Thus, PAR-6 and PKC-3 are essential for early larval development, and we focused our further analysis on PAR-6 and PKC-3.

PAR-6 and PKC-3 are essential in the larval epidermis, but not in the intestine

We next wanted to determine which larval tissue or tissues are severely affected by the loss of PAR-6/ PKC-3 and contribute to the growth arrest. We focused on the two major epithelial organs: the intestine and the epidermis. The intestine is an epithelial tube formed in embryogenesis by 20 cells, which do not divide during larval development. PAR-6 and PKC-3 are highly enriched at the apical luminal domain (Fig. 2A). The epidermis consists of two cell types: hypodermal cells and seam cells.
Fig. 1. PAR-6 and PKC-3 are essential for larval development. (A) Overview of the AID system, which enables targeted degradation of AID-tagged proteins by the plant-derived E3 ubiquitin ligase specificity factor TIR1 upon addition of auxin. (B) Schematic representation of endogenous tagging of par-6, pkc-3, and par-3 loci with sequences encoding a green fluorescent protein (GFP) and auxin-inducible degradation degron (AID) tag. (C–E) Growth curves of N2, par-6::aid::gfp, gfp::aid::pkc-3, and gfp::aid::par-3 animals in absence (- aux) or presence (+ aux) of 4 mM auxin. Data show mean ± SD. n = 6, 7, 8, and 8 for N2 - aux; 6, 7, 9, and 9 for N2 + aux; 7, 6, 9, and 9 for PAR-6 - aux; 8, 6, 7, and 9 for PAR-6 + aux; 22, 11, 10, and 14 for PAR-6 - aux; 19, 14, 9, and 10 for PKC-3 - aux; 10, 10, 10, and 10 for PAR-3 - aux, and 10, 10, 10, and 10 for PAR-3 + aux. (F) Graphical representation of larval epithelial tissues in C. elegans. Green indicates localization of PAR-6, PKC-3, and PAR-3. (G) Distribution of GFP::AID-tagged PAR-6, PKC-3, and PAR-3 in different larval tissues in absence (- auxin) or presence (+ auxin) of 4 mM auxin. Images are maximum projections. Dashed lines in - auxin panels outline pharynx (left panel), intestinal lumen (middle panel) or seam cells (right panel). White arrows point to the excretory canals.
Fig. 2. PAR-6 and PKC-3 are essential in the epidermis to support larval growth. (A, B) Distribution of PAR-6::AID::GFP and GFP::AID::PKC-3 in the intestine (A) and epidermis (B) in absence (- auxin) or presence (+ auxin) of 1 mM auxin. Images are maximum projections of the luminal domain for the intestine, and the apical domain for the epidermis. Drawings are schematic representation of the area imaged, with the localization of PAR-6 and PKC-3 indicated in green shades. Greys indicate absence of PAR-6 and PKC-3. (C–F) Quantification of apical GFP fluorescence intensity at the intestinal lumen and the hyp7–seam cell junction (indicated by colored lines in A, B) in par-6::aid::gfp and gfp::aid::pkc-3 animals in the absence (- aux) or presence (+ aux) of 1 mM auxin. Solid lines and shading represent mean ± SD. For the intestine, n = 10 animals for PAR-6 - aux, PAR-6 + aux, PKC-3 - aux, and PKC-3 + aux. For the epidermis, n = 8 animals for PAR-6 - aux, 6 for PAR-6 + aux, 5 for PKC-3 + aux, and 5 for PKC-3 - aux. (G–J) Growth curves of N2, par-6::aid::gfp, and gfp::aid::pkc-3 animals in absence (- aux) or presence (+ aux) of 4 mM auxin. Solid lines and shading represent mean ± SD. In G and H, degradation was induced in the intestine, and in I and J in the epidermis. In the intestine, n = 13, 10, 13, 14, and 12 for N2 - aux; 7, 7, 7, 5, and 9 for PAR-6 - aux; 6, 6, 6, 5, and 7 for PAR-6 + aux; 8, 7, 8, 4, and 9 for PKC-3 - aux; and 8, 7, 8, 8, and 8 for PKC-3 + aux. In the epidermis, n = 6, 7, 8, and 8 for N2 - aux; 6, 5, 11, and 8 for PAR-6 - aux; 5, 10, 8, and 9 for PAR-6 + aux; 7, 7, 10, and 8 for PKC-3 - aux; and 8, 7, 12, and 13 for PKC-3 + aux.
The hypodermal syncytial cell hyp7 covers most of the body. Embedded within hyp7 are two lateral rows of epithelial seam cells, which contribute multiple nuclei to hyp7 through asymmetric divisions in each larval stage. PAR-6 and PKC-3 localize to the apical domain of the seam cells and hyp7, and are enriched at the seam/seam and seam/hyp7 junctions (Fig. 2B).

To enable tissue-specific depletion of PAR-6 and PKC-3, we generated single-copy integrant lines expressing TIR1 in the intestine and epidermal lineages, using the tissue-specific promoters Pelt-2 and Pwrt-2, respectively. In both tissues, protein depletion occurred within one hour of addition of 1 mM auxin (Fig. 2A–F). To determine the contribution of the intestine and epidermis to the larval growth defects we observed above, we measured the growth rate of animals depleted of PAR-6 or PKC-3 in each tissue. Depletion of either protein from the intestine did not result in a growth delay or in obvious defects in morphology of the intestine (Fig. 2G, H). Simultaneous depletion of PAR-6 and PKC-3 also did not result in a growth delay or visible abnormalities in the intestine (Fig. S1A). These data indicate that PAR-6 and PKC-3 are not essential for the functioning and homeostasis of the larval intestine, though we cannot exclude that very low protein levels that we were not able to detect by fluorescence microscopy are sufficient in this tissue.

A recent study in Drosophila indicated a potential redundancy between aPKC and the kinase Pak1 (Aguilar-Aragon et al., 2018). To determine if a similar redundancy was the cause of the absence of intestinal PKC-3 depletion phenotypes, we combined depletion of PKC-3 with inactivation of C. elegans pak-1 by RNAi. However, we again did not observe a delay in larval growth (Fig. S1B).

In contrast to the intestine, depletion of PAR-6 or PKC-3 from hatching in the epidermis caused an early larval growth arrest, as observed with ubiquitous degradation (Fig. 2I, J). Thus, PAR-6 and PKC-3 play an essential role in the functioning and/or development of epidermal larval epithelia. We did notice that animals with ubiquitous PAR-6 or PKC-3 depletion appeared more sick than epidermal depleted animals, indicating that the functions of PAR-6 and PKC-3 are not limited to the epidermis.

**Cell autonomous and non-autonomous roles for PAR-6 and PKC-3 in molting, seam cell divisions and seam cell morphology**

One of the major functions of the epidermal epithelium is the synthesis and apical secretion of cuticle components, and defects in this tissue can lead to molting defects (Chisholm and Xu, 2012; Lažetić and Fay, 2017). As molting defects can be accompanied by a growth arrest (Brooks et al., 2003; Lažetić and Fay, 2017; Russel et al., 2011; Yochem et al., 1999), we explored the possibility that PAR-6 depleted animals are molting defective. By Nomarski differential interference contrast (DIC) microscopy, we observed incompletely released cuticles 30 hours past exposure to auxin, indicative of molting defects (Fig. 3A). To examine molting progression in more detail, we used a transcriptional reporter expressing GFP from the mlt-10 promoter (Meli et al., 2010). mlt-10 expression cycles, increasing during molting and decreasing during the inter-molt. Upon epidermal degradation of PAR-6 from hatching, mlt-10 driven GFP levels did not cycle and remained low (Fig. 3B, C), indicating a perturbation of the molting cycle. Together, these results demonstrate that PAR-6 performs essential functions in the epidermis required to support remodeling of the C. elegans cuticle.
Fig. 3. PAR-6 and PKC-3 are required in the epidermis for molting and seam cell development. (A) DIC microscopy images of molting defects upon epidermal depletion of PAR-6 or PKC-3. Animals were grown in absence (-auxin) or presence (+auxin) of 1 mM auxin since hatching, and images were taken 30 h after hatching. Dotted lines outline detached but unreleased cuticle in the pharynx and in the tail. Defects are observed in ~50% of the animals. (B) Expression of the molting reporter *Pmllt-10::gfp::pest* in *par-6::aid::gfp* animals in the absence (- auxin) or presence (+ auxin) of 1 mM auxin at 22 h of post-embryonic development. (C) Quantification of *Pmllt-10::gfp::pest* expression from 11 h to 32 h of post-embryonic development (mean fluorescence intensity ± SD) in *par-6::aid::gfp* animals in absence (- aux) or presence (+ aux) of 1 mM auxin. Measurements were done every hour. Each data point is an average of 3–12 measurements, with an average of 8 measurements per data point. (D) Examples of seam cell division and morphology defects observed upon depletion of PAR-6::GFP::AID. Seam cells are visualized by expression of nuclear H2B::GFP and membrane-bound PH::GFP markers (Wildwater et al., 2011). Arrows indicate membrane protrusions and arrowhead indicates abnormal division plane orientation. (E) Seam cell division pattern in *par-6::aid::gfp* animals in absence (control, blue) or presence (+ auxin, magenta) of 1 mM auxin. For the control, *n* = 14, 75, 40, and 28 animals for the L1, L2, L3 and
We next examined the effects of PAR-6 depletion on the stereotypical division pattern of the seam cells. In every larval stage, an asymmetric cell division creates a new seam cell daughter and a cell that differentiates to form neurons or fuse with hyp7 (Chisholm and Hsiao, 2012) (Fig. 3E, blue shaded lineage tree). In the second larval stage, a symmetric division precedes the asymmetric division to double the number of seam cells. Depletion of PAR-6 directly after hatching did not disrupt the L1 asymmetric division, but the divisions that normally take place in the L2 stage were severely delayed (Fig. 3D, E). At the time when control animals were already undergoing the L3 divisions, L2-stage divisions had still not taken place in PAR-6 depleted animals. Eventually, a next round of divisions did take place, but we observed various deviations from the normal L2 division pattern, including division failures and abnormal differentiation and fusion with hyp7. We did not observe any L3 divisions (Fig. 3E). We also observed numerous morphological abnormalities such as membrane protrusions, blebs, and abnormal division plane orientation (Fig. 3D). Exposure of synchronized populations to auxin starting after the L1 or L2 divisions resulted in similar defects, indicating that the seam cell lineage requires the functioning of PAR-6 throughout development (Fig. 3E).

Expression of TIR1 under the wrt-2 promoter results in degradation of target proteins in both hyp7 and the seam cells. While both tissues contribute to cuticle production, severe molting defects can result from the loss of hyp7-specific proteins, such as the NIMA-related kinases NEKL-2,3 (Yochem et al., 2015). It is possible therefore the seam cell defects are a secondary consequence of the molting defect. To address this, we expressed an exogenous copy of par-6::mCherry lacking the degron sequence in hyp7 using the hypodermis-specific dpy-7 promoter (Gilleard et al., 1997). In combination with auxin-induced depletion of PAR-6::GFP::AID by wrt-2 driven TIR-1, this results in absence of PAR-6 only from the seam cells. Hypodermal specific expression of par-6::mCherry rescued the cuticle defects and seam cell division delay observed upon PAR-6 epidermal degradation, and partially rescued the growth arrest (Fig. S2A–D). However, seam cell morphology defects and the abnormal cell division plane were not restored (Fig. S2C). Thus, PAR-6 expression in the hypodermis is sufficient to support molting and larval growth. Furthermore, at least some of the seam cell defects we observed, most notably the timing defect, appear to be a secondary consequence of hypodermal or molting defects. The fact that the growth arrest and seam abnormalities were not fully rescued may indicate cell autonomous roles for PAR-6 in the seam, or alternatively that the Pdpy-7::par-6::mCherry rescue construct is not fully functional.
To determine the effects of a molting arrest on seam cell divisions via an alternative approach, we used a CRISPR-tagged NEKL-2::AID strain that arrests molting upon auxin addition (Joseph et al., 2020). NEKL-2 depletion caused defects in the morphology of seam cells, as well as reduced cell division, confirming that abnormalities in the hypodermis can affect the seam cells (Fig. S2E, F). However, in contrast to PAR-6 depletion, not all seam cell divisions were affected, indicating that a block in molting does not fully explain the seam cell division delay in PAR-6 depleted animals.

Taken together, our data shows that PAR-6 and PKC-3 have essential roles in the *C. elegans* epidermis to support cuticle homeostasis and molting. Restoring *par-6(+)* expression specifically in the hypodermis revealed the importance of PAR-6 for this cell type and indicate that the hypodermis and/or molting are essential for proper seam cell development. Finally, PAR-6 may perform cell autonomous functions in the seam cells that contribute to proper seam cell shape and cell division plane.

**PAR-6 and PKC-3 mediate apical LGL-1 exclusion and promote junction integrity in the larval epidermis**

As one of the major functions of the apical PAR complex is to mediate the exclusion of basolateral proteins from the apical domain, we next examined the effects of PKC-3 depletion on two key aPKC target genes: LGL-1/Lgl and PAR-1. Both proteins are direct aPKC targets in epithelia, and in the *C. elegans* zygote their anterior exclusion is mediated by PKC-3 (Beatty et al., 2010; Betschinger et al., 2003; Doerflinger et al., 2010; Hoege et al., 2010; Hurov et al., 2004; Motegi et al., 2011; Plant et al., 2003; Ramanujam et al., 2018; Yamanaka et al., 2003). For these experiments we made use of integrated LGL-1::GFP transgene (Waaijers et al., 2015) and an endogenously tagged PAR-1::GFP fusion.

Depletion of PKC-3 in the intestine did not result in apical invasion of LGL-1 (Fig. S3A). In contrast, degradation of PKC-3 in the epidermis resulted in clear apical LGL-1 localization in the seam cells within 6 hours of auxin addition (Fig. 4A, B). Degradation of PKC-3 in the epidermis did not result in apical PAR-1 localization (Fig. 4C, D). Instead, prolonged depletion of PKC-3 for 24h resulted in fragmentation of the normally contiguous PAR-1 signal at cell junctions, which may reflect an indirect effect of PKC-3 on junction organization (Fig. 4C). These results demonstrate that PKC-3 is necessary to maintain the basolateral localization of LGL-1 in the seam cells, but not the intestine. In contrast, the apical exclusion of PAR-1 is not solely mediated by aPKC, though the organization of PAR-1 at cell junctions does depend on PKC-3.

In embryonic epithelia, PAR-6 and PKC-3 are essential for proper junction formation, with loss of either protein resulting in fragmented cell junctions (Montoyo-Rosario et al., 2020; Totong et al., 2007). To investigate the requirement for PAR-6 and PKC-3 in junction integrity in larval epithelia, we assessed the localization of an endogenous GFP fusion of the junctional protein DLG-1 upon degradation of PAR-6 or PKC-3 from hatching. In control animals not exposed to auxin, DLG-1 displays the typical ladder-like intestinal junction pattern and forms a continuous apical belt around the seam cells (Figs S3B and 4E, F). Upon degradation of PAR-6 in the intestine, we did not observe
junctional defects (Fig. S3B). We also did not observe any changes to the DLG-1 localization pattern in the epidermis after 6 hours of PAR-6 or PKC-3 depletion (Fig. 4E). However, after 24 h of degradation, DLG-1 no longer localized in a uniform band around the seam cells but appeared fragmented, with aggregates of bright DLG-1 interspersed with areas lacking fluorescent signal (Fig. 4F). We also observed fluorescent aggregates in the hypodermis (Fig. 4F). Thus, as in the embryo, PAR-6 and PKC-3 are essential for junction integrity in the epidermis. The fact that junctional defects took 24 h to develop, compared to 6 h for LGL-1 mislocalization, points to an inherent stability of cell junctions.

Fig. 4. PKC-3 excludes LGL-1 from the apical cortex and, together with PAR-6, regulates junctions. (A, B) Distribution and quantification of LGL-1::GFP in the epidermis of lgl-1::gfp animals without auxin and in lgl-1::gfp; gfp::aid::pkc-3; Pwrt-2::tir-1::bfp animals in the presence of 4mM auxin. Images are maximum projections of the apical domain. Quantifications show mean apical GFP fluorescence intensity ± SD at the hyp7–seam cell junction, normalized to background intensity of each animal measured in the hypodermis. n = 7 animals for both conditions. (C, D) Distribution and quantification of PAR-1::GFP in the epidermis in par-1::gfp animals without auxin and in par-1::gfp; gfp::aid::pkc-3; Pwrt-2::tir-1::bfp animals in the presence of 4mM auxin. Images are maximum projections of the apical domain. Quantifications show mean apical GFP fluorescence intensity ± SD at the hyp7–seam cell junction, normalized to the background intensity of each animal measured in the hypodermis. n = 6 animals for both conditions. (E, F) Junction organization visualized by DLG-1::mCherry expression in par-6::aid::gfp or gfp::aid::pkc-3 animals in the absence (- auxin) or presence (+ auxin) of 1 mM auxin for 6 (E) or 24 (F) hours. (G) Graphical representation of junctional defects in the seam cells upon PAR-6 or PKC-3 degradation.
Finally, we investigated the localization dependencies between PAR-6, PKC-3 and PAR-3. Several studies demonstrated that PAR-6 and PKC-3 co-localize throughout development, and are mutually dependent on each other for their asymmetric localization (Bossinger et al., 2001; Leung et al., 1999; McMahon et al., 2001; Nance and Priess, 2002; Nance et al., 2003; Tabuse et al., 1998; Totong et al., 2007). Moreover, binding of PAR-6 to PKC-3 is required for apical localization of PAR-6, including in larval epithelia (Li et al., 2010a). Not surprisingly therefore, degradation of PAR-6 resulted in the loss of PKC-3 from the apical domain of the seam cells, and degradation of PKC-3 similarly disrupted PAR-6 localization (Fig. S4A, B). These disruptions occurred rapidly, within 1 h of auxin addition. Depletion of PAR-6 in the intestine also caused rapid loss of PKC-3 from the apical plasma membrane, demonstrating that, despite the lack of intestinal phenotypes, PAR-6 and PKC-3 do form an apical protein complex (Fig. S3C, D). When we followed the apical loss of PKC-3 in the intestine over time, we observed similar dynamics of PAR-6 depletion and PKC-3 loss (Video S1). Our results thus confirm the interdependency between PAR-6 and PKC-3. Surprisingly, the effects of PAR-6 and PKC-3 depletion on PAR-3 were distinct: the apical localization of PAR-3 in the seam cells depended on the presence of PKC-3, but not PAR-6 (Fig. S4C, D). Finally, we examined the effects of PAR-3 degradation on PAR-6 and PKC-3 in the seam cells. We observed partially reduced levels of PAR-6, but no effect on PKC-3 (Fig. S4E, F). Thus, PAR-6 and PKC-3 may not form an obligatory dimer under all conditions.

PAR-6 and PKC-3 control the organization of non-centrosomal microtubule arrays in the hypodermis

The loss of PAR-6 or PKC-3 affected several epidermal processes in which cytoskeletal elements play important roles, including molting, seam cell divisions, and maintaining proper seam cell morphology. The PAR proteins play essential roles in organizing the actomyosin cytoskeleton and microtubules in different settings, including asymmetric cell division, neuronal differentiation, and epithelial polarization (Goldstein and Macara, 2007; Rodriguez-Boulan and Macara, 2014; St Johnston, 2018). We therefore investigated if PAR-6 degradation affects the organization of actin or microtubules in the epidermis. To assess the organization of the actin cytoskeleton we used an epidermal transgene expressing the actin-binding-domain of \textit{vab-10} fused to \textit{mCherry} (Gally et al., 2009). We depleted PAR-6 from hatching and examined actin organization after 24 h in late L2 larvae. Consistent with previous observations (Costa et al., 1997), we observed prominent circumferential actin bundles in hyp7, strong enrichment of actin along the hyp7/seam junctions, and largely anterior/posteriorly organized actin within the seam cells of control animals at this stage (Fig. 5A). Upon PAR-6 depletion, actin organization appeared largely undisturbed in both the seam cells and hypodermis (Fig. 5A), and actin bundles in hyp7 remained perpendicular to the seam cells (Fig. 5B). These data indicate that PAR-6 does not play a major role in regulating the actin cytoskeleton in the larval epidermis.

We next inspected the organization of the microtubule cytoskeleton using an endogenously GFP tagged variant of the microtubule-binding protein MAPH-1.1 (Waaijers et al., 2016). We degraded PAR-6 in the epidermis from hatching and assessed the organization of epidermal microtubule arrays after 24 h. In control animals, we observed highly ordered circumferential microtubule bundles in the dorsal and ventral sections of hyp7 underlying the muscle quadrants, and a microtubule meshwork in
Fig. 5. PAR-6 and PKC-3 control microtubule organization in the *C. elegans* epidermis. (A) Actin organization visualized by the *Plin-26::ABDvab-10::mCherry* reporter in *par-6::aid::gfp* animals in absence (- auxin) or presence (+ auxin) of 1 mM auxin for 24 hours. (B) Quantification of actin bundle orientation. Angle is measured relative to the anterior (180°) – posterior (0°) axis. n = 100 bundles in 5 animals per condition. (C) Microtubule organization of the indicated genotypes visualized by *MAPH-1.1::GFP* in absence (- auxin) or presence (+ auxin) of 1 mM auxin for 24 hours. Images are maximum intensity projections. (D) Hypodermal microtubule bundle density. n = 13 animals for control, 15 for PAR-6 deg, 14 for PKC-3 deg and 13 for *noca-1(ok3692)*. Bars show mean ± SD (E) Microtubule growth visualized by maximum intensity projection of the plus end marker *EBP-2::GFP* in absence (- auxin) or presence (+ auxin) of 1 mM auxin for 24 hours. To match the age of animals in (C), we depleted PAR-6 for 1 h starting with 23 h old L2 animals. (F) *EBP-2* comet density. n = 12 animals for control and PAR-6 deg, 8 for *noca-1(ok3692)*, and 8 for PAR-6 deg + rescue. Bars show mean ± SD (G) Microtubule growth rate. n > 400 comets. Bars = mean ± SD (H) Quantification of microtubule growth orientation. Vertical axis: left/right orientation; horizontal axis: anterior/posterior orientation. n = 150 comets. Bars = mean ± SD. Tests of significance: Tukey’s multiple comparisons test for D, and Dunn’s multiple comparisons test for F and G. ns = not significant, * = P ≤ 0.05, ** = P ≤ 0.01, *** = P ≤ 0.001, **** = P ≤ 0.0001.
the lateral sections of hyp7 abutting the seam cells, as previously reported (Chuang et al., 2016; Costa et al., 1997; Taffoni et al., 2020; Wang et al., 2015) (Fig. 5C). In the seam cells the microtubule network was less well defined but also forms a meshwork (Fig. 5C). In PAR-6 depleted animals, we observed a significant reduction in the density of circumferential microtubule bundles in the hypodermis (Fig. 5C, D). Epidermal depletion of PKC-3 resulted in similar defects (Fig. 5C, D). To understand the cause of the reduced microtubule density, we investigated microtubule dynamics using an endogenous fusion of the microtubule plus-end tracking protein EBP-2 to GFP. In control animals, EB1 comets moved along trajectories consistent with the organization of microtubule bundles in the epidermis, and both comet density and growth rates match previous reports (Fig. 5 E–G) (Chuang et al., 2016; Taffoni et al., 2020; Wang et al., 2015). Already within 1 h of inducing depletion of PAR-6, we observed reduced microtubule dynamics (Fig. 5E–G). The density of growing MTs was reduced by 56% (Fig. 5F), and microtubule growth rate was reduced by 14% in hyp7 and by 16% in the seam cells (Fig. 5G). These results suggest that the reduced density of microtubule bundles upon depletion of PAR-6 is the result of reduced growth or nucleation of microtubules. We also observed a defect in the directionality of microtubule growth. While 54% of the comets in control animals travel perpendicular to the seam cells (70–110°), this number is reduced to 24% upon PAR-6 degradation (Fig. 5H), consistent with the defects in organization observed with GFP::MAPH-1.1.

PAR-6 controls microtubule organization through its interaction partner NOCA-1/Ninein and the γ-tubulin ring complex

Two large-scale protein–protein interaction mapping studies in C. elegans had identified the microtubule organizing protein NOCA-1 as an interaction partner of PAR-6 (Boxem et al., 2008; Lenfant et al., 2010). Affinity purification experiments showed that PAR-6 interacts with NOCA-1 through its PDZ domain (Lenfant et al., 2010), and we were able to confirm the PAR-6 PDZ interaction with NOCA-1 by yeast two-hybrid (Fig. S5). NOCA-1 functions together with γ-tubulin to assemble non-centrosomal microtubule arrays in multiple tissues, including the epidermis, and is thought to be a functional homolog of the vertebrate microtubule organizer Ninein (Green et al., 2011; Wang et al., 2015). NOCA-1 localizes to the apical cortex in seam cells, similar to the localization of PAR-6 (Figs 1C, 6A), but the mechanisms that mediate apical localization of NOCA-1 are currently not known. The physical interaction between PAR-6 and NOCA-1 prompted us to investigate if PAR-6 regulates non-centrosomal microtubule arrays through NOCA-1.

We first examined the effect of epidermal PAR-6 depletion on the localization of NOCA-1. To visualize NOCA-1 we made use of an existing transgenic line that expresses the epidermis specific NOCA-1d and e isoforms fused to GFP from their endogenous promoter (Wang et al., 2015). In untreated control animals, we observed punctate localization of NOCA-1 in the epidermis, mostly clustered at the seam/seam and seam/hyp7 junctions, as previously observed (Fig. 6A) (Wang et al., 2015). Addition of auxin to induce epidermal PAR-6 degradation led to a 61% reduction in junctional levels of NOCA-1 within 6 hours (Fig. 6A, B), demonstrating that PAR-6 promotes the apical localization of NOCA-1.
NOCA-1 was reported to work together with γ-tubulin and redundantly with Patronin/PTRN-1 in controlling circumferential microtubule bundle organization in the hypodermis (Wang et al., 2015). We therefore examined the effect of PAR-6 depletion on the localization of PTRN-1 and GIP-1, two core components of the γ-TuRC required to localize other γ-TuRC components to the apical non-centrosomal microtubule organizing center (ncMTOC) in the embryonic intestine (Sallee et al., 2018). To visualize PTRN-1 and GIP-1 we used endogenous PTRN-1::GFP and RFP::GIP-1 fusion proteins. GIP-1 localized in a punctate pattern at the seam/seam and seam/hyp7 junctions, similar to NOCA-1 (Fig. 6C) (Sallee et al., 2018; Wang et al., 2015). PTRN-1 also localized in a punctate pattern, but dispersed through the epidermis and lacking the junctional enrichment seen upon PAR-6 depletion. All images are maximum projections of the apical domain. Quantifications in B, D and H show mean apical GFP fluorescence intensity ± SD at the hyp7-seam cell junction (indicated by colored lines), normalized to background intensity of each animal measured in the hypodermis. Quantification in F shows mean PTRN-1::GFP puncta density ± SD. Tests of significance: unpaired t-test for F. *** = P ≤ 0.001.

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for NOCA-1 and GIP-1 (Fig. 6E) (Wang et al., 2015). Upon PAR-6 degradation, junctional GIP-1 levels were strongly reduced (Fig. 6C, D), similarly to NOCA-1. We also observed that PAR-6 depletion resulted in a decrease in the number of PTRN-1 puncta in the epidermis (Fig. 6E, F). As NOCA-1 is a direct interaction partner of PAR-6, we examined if the loss of GIP-1 is due to the loss of NOCA-1 localization. Indeed, in a noca-1(ok3692) deletion mutation GIP-1 levels were significantly reduced (Fig. 6G, H), suggesting that NOCA-1 acts upstream of GIP-1 in the C. elegans larval epidermis.

Finally, we examined if the failure to properly localize NOCA-1 could explain the microtubule defects we observed upon PAR-6 depletion. We determined microtubule bundle density, EB1 comet density, and microtubule growth rate in noca-1(ok3692) animals. In the noca-1(ok3692) deletion mutant, we observed a significant reduction in the density of circumferential microtubule bundles in the hypodermis (Fig. 5C, D). As reported in a previous study (Wang et al., 2015), we also observed reduced microtubule dynamics, with the density of growing MTs reduced by 65%, and microtubule growth rates reduced by 65% (Fig. 5E, G). These values are all very similar to those we observed upon PAR-6 depletion and are consistent with a model in which the microtubule defects caused by PAR-6 depletion are a result of the requirement of PAR-6 in localizing NOCA-1. The effects on PTRN-1 may be a secondary consequence of microtubule defects caused by NOCA-1 loss.

**Discussion**

Par6 and aPKC are essential for apical–basal polarization across animal species. Here, we used inducible protein degradation to identify an essential role for PAR-6 and PKC-3 in the larval epidermis of C. elegans, and a novel role for PAR-6 in regulating the assembly of microtubule bundles through its interaction partner NOCA-1/Ninein. Most studies of the apical PAR proteins in C. elegans have focused on embryonic tissues, and an essential role in postembryonic development has not been described. Depletion of par-6, pck-3, or par-3 by RNAi in larval stages caused defects in polarization of spermathecal cells and in ovulation, but not in larval development (Aono et al., 2004). Similar results were recently observed using a temperature sensitive pck-3 allele grown at non-permissive temperature (Montoyo-Rosario et al., 2020). More severe phenotypes were observed in hatching progeny (escapers) of par-6, pck-3, or par-3 RNAi-treated mothers, which showed partially penetrant defects in outgrowth of vulval precursor and seam cells, migrations of neuroblasts and axons, and the development of the somatic gonad (Welchman et al., 2007). The lack of a growth arrest phenotype in these studies presumably reflects incomplete gene inactivation.

**Auxin-inducible protein depletion of PAR proteins**

The auxin-inducible degradation approach allowed us to bypass embryonic requirements and examine the roles of PAR-6, PKC-3, and PAR-3 in specific epithelial tissues during larval development. Despite these advantages, one drawback of all protein degradation approaches is that it remains difficult to draw conclusions from negative results. Although we tagged all known PAR-3 protein isoforms and observed efficient protein depletion, ubiquitous depletion of PAR-3 did not cause obvious defects in larvae. While PAR-3 may indeed not be essential in larval tissues, it is also possible that very low levels of PAR-3 are sufficient for its functioning, or that unpredicted splicing events cause the
expression of non-degron tagged PAR-3 isoforms. One approach to counteract the latter possibility would be to replace the endogenous gene with a re-engineered copy that is unlikely to express alternative splice variants, e.g. by replacing natural introns with artificial ones and removing internal promoters. However, removing this level of regulation and expressing only one isoform may affect the functioning of par-3 and cause unintended side effects. We also did not detect phenotypes upon depletion of PAR-6 or PKC-3 in the larval intestine. Similar caveats as for PAR-3 depletion apply here, though PAR-6 depletion did lead to complete loss of PKC-3 from the apical domain, and epidermal depletion caused severe phenotypes. These observations make it less likely that the lack of a phenotype is due to the expression of unknown isoforms. PAR-6 and PKC-3 are likely to play non-essential or redundant roles in the intestine, as a previous study found that PAR-6 contributes to endosome positioning in this tissue (Winter et al., 2012).

Roles of PAR-6 and PKC-3 in junction formation and cell polarity

Depletion of PAR-6 and PKC-3 in the epidermis resulted in a fragmented appearance of the hyp7/seam and seam/seam junctions, similar to previous observations in embryonic epithelia (Montoyo-Rosario et al., 2020; Totong et al., 2007; Von Stetina and Mango, 2015; Von Stetina et al., 2017). Our data therefore further demonstrate a general requirement for PAR-6 and PKC-3 in junction formation or maintenance. We also examined if PKC-3 functions to exclude the basolateral polarity proteins PAR-1 and LGL-1 from the apical domain. In the epidermis, PKC-3 depletion caused a rapid invasion of LGL-1 in the apical domain of the seam cells, while PAR-1 remained junctional and basal. Thus, as in the one cell embryo, PKC-3 functions to exclude LGL-1 in the seam cells. A recent study found that LGL-1 can suppress sterility of a temperature sensitive \textit{pkc-3} allele, further indicating that the interaction between LGL-1 and PKC-3 is functionally relevant (Montoyo-Rosario et al., 2020).

In contrast to the epidermis, LGL-1 localization in the intestine remained unchanged upon PKC-3 depletion, and we observed no obvious abnormalities in the intestine upon PAR-6 or PKC-3 depletion. Thus, while PAR-6 and PKC-3 are essential for development of the embryonic intestine (Totong et al., 2007), they do not appear to be essential in the larval intestine. Other cellular systems, such as polarized protein trafficking, may suffice to maintain cell polarity in the absence of the apical PAR proteins (Shafaq-Zadah et al., 2012; Zhang et al., 2011; Zhang et al., 2012). An analogous situation exists in the Drosophila midgut, where integrins, but not the apical PAR proteins, are essential for polarization (Chen et al., 2018). The lack of LGL-1 mislocalization also points to the existence of possible redundancies in polarization of cortical polarity regulators, which may be uncovered through enhancer screens in PAR-6 or PKC-3 depleted backgrounds.

In embryonic epithelia, the requirements of the apical PAR proteins also vary between tissues. Intestinal and epidermal cells depleted of PAR-6 or PKC-3 using the ZF1 system still show apicobasal polarization, as evidenced by apical localization of junctional and cytoskeletal proteins (Montoyo-Rosario et al., 2020; Totong et al., 2007). However, in the arcade cells of the pharynx, most PAR-6 depleted animals show no apical enrichment of junctional or apical cytoskeletal markers (Von Stetina and Mango, 2015). These data further highlight that the requirements for PAR-6 and PKC-3 can vary between tissues.
Interdependencies between the PAR proteins

PAR-6 and PKC-3 were mutually dependent in both the epidermis and intestine. This result was not surprising, as Par6 and aPKC act as a dimer and have been shown to be mutually dependent in other *C. elegans* tissues (Hung and Kemphues, 1999; Li et al., 2010a; Nance et al., 2003; Tabuse et al., 1998; Totong et al., 2007). We also found that depletion of PAR-3 from the epidermis resulted in a partial loss of PAR-6. This is consistent with findings that multiple mechanisms can promote cortical recruitment of Par6–aPKC, including binding of Par6 to Par3 or Cdc42, and binding of aPKC to phospholipids (Dong et al., 2020; Hong et al., 2003; Hutterer et al., 2004; Joberty et al., 2000; Nagai-Tamai et al., 2002; Nunes de Almeida et al., 2019; Rodriguez et al., 2017; Wang et al., 2017; Wodarz et al., 2000). Conversely, the apical localization of PAR-3 was dependent upon PKC-3. While Par3 often localizes independently of Par6/aPKC, a similar dependency has been observed in Drosophila and in the *C. elegans* zygote (Hutterer et al., 2004; Petronczki and Knoblich, 2001; Tabuse et al., 1998; Watts et al., 1996).

Two results were more surprising in light of the interdependency between PAR-6 and PKC-3. First, whereas depletion of PKC-3 causes a loss of apical PAR-3, depletion of PAR-6 did not have a similar effect. Second, PAR-3 depletion resulted in a partial reduction of apical PAR-6 levels, but not of PKC-3. An independence in PAR-6 and PKC-3 localization has been observed previously in the one cell embryo, where reduction of the chaperone protein CDC-37 by RNAi resulted in the localization of PAR-6 to the cortex independently of PKC-3 (Beers and Kemphues, 2006). Thus, under certain conditions, PAR-6 and PKC-3 may not act as an obligate dimer.

A novel role for PAR-6 in epidermal microtubule organization

Epidermal specific depletion uncovered a novel role for PAR-6 in organizing non-centrosomal microtubule bundles. In epithelial cells, apical non-centrosomal microtubule organizing centers (ncMTOCs) assemble apical–basal microtubule arrays. ncMTOCs contain proteins and complexes involved in microtubule anchoring, microtubule stabilization, and microtubule nucleation — such as the γ-tubulin ring complex (γ-TuRC) (Sanchez and Feldman, 2017). How apical ncMTOCs are organized is not well understood, but several studies indicate an important role for apical PAR proteins in this process. In the cellularizing Drosophila embryo, aPKC is required for the transition from centrosome emanated asters to non-centrosome associated apical–basal bundles (Harris and Peifer, 2007). In the developing embryonic intestine of *C. elegans*, PAR-3 is needed for the redistribution of γ-tubulin and other microtubule regulators from the centrosomes to the apical domain of the cell (Feldman and Priess, 2012). A role for Par6 in regulating microtubule organizing centers may not be limited to epithelial ncMTOCs. For example, in several mammalian cultured cell lines Par6 is a component of centrosomes and regulates centrosomal protein composition (Dormoy et al., 2013; Kodani et al., 2010).

Epidermal depletion of PAR-6 resulted in reduced numbers of circumferential microtubule bundles, as well as a reduced microtubule growth rate and EB1 comet density. Moreover, depletion of PAR-6 led to a loss of apical NOCA-1 enrichment at seam–seam and seam–hyp7 junctions. The effects of PAR-6 depletion on microtubule organization and dynamics are very similar to those we observed in a *noca-1*
mutant. While other models are possible, these data are consistent with PAR-6 acting through NOCA-1 to control microtubule organization in the epidermis. The reduced microtubule growth rate and EB1 comet density we observed in \textit{noca-1} mutant animals have been reported previously (Wang et al., 2015). However, no defects in circumferential microtubule bundle density were observed in that study, despite using the same \textit{noca-1(\textit{ok3692})} allele. The observed difference may be a result of a difference in exact experimental procedure or the precise genetic background used. For example, whereas we used the microtubule-binding protein GFP::MAPH-1.1 to label microtubules, the study by Wang \textit{et al.} used a GFP::\(\beta\)-tubulin fusion.

We also found that, in the epidermis, the localization of GIP-1 is dependent on NOCA-1. The relationship between NOCA-1 and \(\gamma\)-TuRC components has been examined previously in two different tissues (Sallee \textit{et al.}, 2018; Wang \textit{et al.}, 2015). In the germline, NOCA-1 co-localizes with \(\gamma\)-tubulin to non-centrosomal microtubule arrays but is not required for the localization of \(\gamma\)-tubulin (Wang \textit{et al.}, 2015). In fact, in this tissue the localization of a short NOCA-1 protein lacking isoform-specific N-terminal extensions is dependent for its localization on \(\gamma\)-tubulin. The longer NOCA-1h isoform, however, localizes independently of \(\gamma\)-tubulin, indicating the presence of multiple NOCA-1 localization signals (Wang \textit{et al.}, 2015). In the embryonic intestine, the localization of NOCA-1 was not altered by the depletion of GIP-1 (Sallee \textit{et al.}, 2018). However, microtubule organization in the intestine is regulated differently from the epidermis, as apical microtubule organization was largely normal even in \textit{ptrn-1} mutant animals depleted of intestinal NOCA-1 and GIP-1 (Sallee \textit{et al.}, 2018). Thus, differential effects of \(\gamma\)-TuRC component loss may reflect differences in the mechanisms of microtubule regulation. Whether PAR-6 plays a role in ncMTOC assembly and microtubule organization in tissues other than the epidermis remains to be investigated.

In addition to the effects on NOCA-1 and GIP-1, PAR-6 depletion resulted in a reduced number of PTRN-1 puncta in the epidermis. PTRN-1 is a member of the Patronin/CAMSAP/Nezha family of minus end-associated proteins, which stabilize and protect uncapped microtubule minus ends (Atherton \textit{et al.}, 2019; Goodwin and Vale, 2010; Hendershott and Vale, 2014; Jiang \textit{et al.}, 2014). NOCA-1 was previously shown to act in parallel with PTRN-1 in organizing circumferential microtubule arrays in the \textit{C. elegans} epidermis (Wang \textit{et al.}, 2015). The mechanistic details of the relationship between NOCA-1 and PTRN-1 have not been resolved, but their distinct localization patterns suggest that they act on distinct pools of microtubules. Our data does not reveal why PAR-6 depletion results in a reduced number of PTRN-1 foci, but a likely possibility is that this is a secondary consequence of the microtubule defects caused by the loss of NOCA-1 localization.

**Mechanisms of larval growth arrest**

The depletion of PAR-6 or PKC-3 in the epidermis led to a rapid growth arrest and cessation of the molting cycle. What causes this dramatic effect on animal development? The junctional defects we observed are unlikely to be the primary consequence, as effects on cell junctions appeared only after 24 h of exposure to auxin. The effects on LGL-1 were more rapid but are also not likely to explain the growth arrest, as \textit{lgl-1} mutants are viable (Beatty \textit{et al.}, 2010; Hoege \textit{et al.}, 2010). The effects on the microtubule cytoskeleton are likely to contribute to the growth arrest or molting defects. However,
noca-1 mutants displayed similar microtubule defects as PAR-6 depletion yet develop to adulthood. Interestingly, noca-1; ptrn-1 double mutant animals do grow slowly and frequently die before reaching adulthood (Wang et al., 2015). Thus, the combined defects in NOCA-1 and PTRN-1 localization we observed upon PAR-6 depletion may partially explain the growth defects. The roles of PTRN-1 may not be limited to microtubule regulation, as a recent study demonstrated that PTRN-1 stimulates actin polymerization during endocytic recycling in the intestine (Gong et al., 2018). A final possibility is that the molting defect is the primary cause of the developmental arrest, as failure to molt can cause a growth arrest (Brooks et al., 2003; Lažetić and Fay, 2017; Russel et al., 2011; Yochem et al., 1999).

One way in which PAR-6 and PKC-3 could affect molting is by affecting intracellular trafficking. Molting requires the coordinated activity of the endocytic and exocytic machineries to mediate apical secretion of new cuticle components, recycling of old cuticle components, and uptake of sterols from the environment (Lažetić and Fay, 2017). Accordingly, mutations in multiple regulators of intracellular trafficking cause molting defects (Lažetić and Fay, 2017). Several links between cortical polarity regulators and the polarized trafficking machinery have been uncovered (Rodriguez-Boulan and Macara, 2014). In C. elegans, par-3, par-6, and pck-3 were all found to be required for endocytic trafficking in oocytes, and RNAi for par-3 and par-6 causes scattering of multiple endosome types in the intestine (Balklava et al., 2007; Winter et al., 2012). It is possible, therefore, that PAR-6 and PKC-3 regulate vesicle trafficking in molting as well. Such regulation may be indirect, through regulation of cytoskeletal components, or through more direct mechanisms remaining to be uncovered.

In summary, our data supports that PAR-6 and PKC-3 have multiple roles in the epidermis that support larval development and molting. We have also uncovered an important role for PAR-6 in regulating the microtubule cytoskeleton, while additional mechanisms through which PAR-6 and PKC-3 control growth and/or molting likely remain to be discovered.

**Materials and Methods**

*C. elegans* strains

All *C. elegans* strains used in this study are derived from the N2 Bristol strain, and are listed in Table 1. All strains were maintained at 20 °C on Nematode Growth Medium (NGM) plates seeded with *Escherichiae coli* OP50 bacteria under standard conditions (Brenner, 1974).

**CRISPR/Cas9 genome engineering**

All gene editing was done by homology-directed repair of CRISPR/Cas9-induced DNA double-strand breaks, using plasmid-based expression of Cas9 and sgRNAs. All edits were made in an N2 background, with the exception of 2x(egfp::aid)::par-3, for which egfp::aid::par-3 was used as the starting background. All fusions were repaired using a plasmid-based template with 190–600 bp homology arms and containing a self-excising cassette (SEC) for selection (Dickinson et al., 2015). The homology arms included mutations of the sgRNA recognition sites to prevent re-cutting after repair. The par-6::aid::egfp, par-6::mCherry, dlg-1::mCherry and ebp-2::egfp vectors were cloned using Gibson assembly and vector pJJR82 (Addgene #75027) (Gibson et al., 2009; Ramalho et al., 2020) as the backbone. The 2x(egfp::aid)::par-3, *Pwrt-2::tir-1::bfp* and *Pelt::tir-1::bfp* vectors were cloned using SapTrap assembly into vector pMLS257 (Schwartz and Jorgensen, 2016), and the egfp::aid::pck-3 and mCherry::pck-3 vectors were cloned using SapTrap assembly into vector pDD379 (Dickinson et al., 2018). The sgRNAs were expressed from a plasmid under control of a U6 promoter. To generate sgRNA vectors, antisense oligonucleotide pairs were annealed and ligated into BbsI-linearized pJJR50 (Addgene #75026) (Waaijers et al., 2016), with the exception of the pck-3 fusions, in which the sgRNA was incorporated into assembly
vector pDD379 using SapTrap assembly. The targeted sequences can be found in Table 2. Injection mixes were prepared in MilliQ H$_2$O and contained 50 ng/ml Pef3::cas9 (Addgene ID #46168) (Friedland et al., 2013), 50–100 ng/µl U6::sgRNA, 50 ng/µl of repair template, with the exception of the pkc-3 fusions, in which the sgRNA-repair-template vector was used at a concentration of 65 ng/µl. All mixes also contained 2.5 ng/µl of the co-injection pharyngeal marker Pmyo-2::GFP or Pmyo-2::tdTomato to aid in visual selection of transgenic strains. Young adult hermaphrodites were injected in the germline using an inverted micro-injection setup (Eppendorf FemtoJet 4x mounted on a Zeiss Axio Observer A.1 equipped with an Eppendorf Transfomer 4r). Candidate edited progeny were selected on plates containing 250 ng/µl of hygromycin (Dickinson et al., 2015), and correct genome editing was confirmed by Sanger sequencing (Macrogen Europe) of PCR amplicons encompassing the edited genomic region. From correctly edited strains, the hygromycin selection cassette was excised by a heat shock of L1 larvae at 34 °C for 1 h in a water bath. Correct excision was confirmed by Sanger sequencing. Sequence files of the final gene fusions in Genbank format are in Supplementary File 1.

*C. elegans* synchronization

In order to obtain synchronized worm populations, plates with eggs were carefully washed with M9 (0.22 M KH$_2$PO$_4$, 0.42 M Na$_2$HPO$_4$, 0.85 M NaCl, 0.001 M MgSO$_4$) buffer in order to remove larvae and adults but leave the eggs behind. Plates were washed again using M9 buffer after an hour, to collect larvae hatched within that time span.

**Auxin Inducible Degradation**

Auxin treatment was performed by placing synchronized populations of worms on NGP plates seeded with *E. coli* OP50 and containing 1 or 4 mM auxin. To prepare plates, auxin (Alfa Aesar A10556) was diluted into the autoclaved NGM agar medium after cooling to 60 °C prior to plate pouring. Plates were kept for a maximum of 2 weeks in the dark at 4 °C.

**C. elegans growth curves**

To measure growth curves, L1 animals synchronized as described above were placed on NGM plates seeded with *E. coli* OP50 and either lacking auxin or containing 4 mM auxin. Images were taken in 24 h intervals up to 96 h, using a Zeiss Axio Zoom.V16 equipped with a PlanNeoFluar Z 1x/0.25 objective and Axiocam 506 color camera, driven by Zen Pro software. Animal length was quantified in ImageJ(FIJI) software by drawing a spline along the center line of the animal (Rueden et al., 2017; Schindelin et al., 2012).

**Feeding RNAi**

pak-1 RNAi clones were obtained from the genome-wide Ahringer and Vidal RNAi feeding libraries (Kamath and Ahringer, 2003; Rual et al., 2004). Ahringer clone supplied through Source BioScience. An HT115 clone expressing the L4440 vector lacking an insert was used as a control. For feeding RNAi experiments, bacteria were cultured overnight in 10 mL of Lysogeny Broth (LB) supplemented with 100 µg/ml ampicillin (Amp) and 2.5 µg/ml tetracyclin (Tet) at 37°C in an incubator rotating at 200 rpm. Cultures were incubated for 120 min in the presence of 1 mM Isopropyl β-D-1-thiogalactopyranoside (IPTG) to induce production of dsRNA. Cultures were pelleted by centrifugation at 4000 g for 15 min and concentrated 5x. NGM agar plates supplemented with 50 µg/ml Amp, 12.5 µg/mL Tet and 1 mM IPTG were seeded with 250 µl of bacterial suspension. 8-10 L4 hermaphrodites per strain were transferred to NGM-RNAi plates, and two plates were analyzed per condition. The F1 generation was synchronized and its growth curved determined as described above.

**Molting assay**

Synchronized L1 animals were placed on NGM plates seeded with *E. coli* OP50 and either lacking auxin or containing 1 mM auxin. Fluorescence images were taken in 1 h intervals from 11 h to 32 h of development, using a Zeiss Axio Zoom. V16 equipped with a PlanNeoFluar Z 1x/0.25 objective and Axiocam 506 color camera, driven by Zen Pro software. Expression levels of the *Pmlt-10::gfp::pest* reporter were quantified in ImageJ(FIJI) software (see image analysis).

**Seam cell lineage analysis**

Synchronized animals were placed on NGM plates seeded with *E. coli* OP50 and either lacking auxin or containing 1 mM auxin. Animals were placed on plates immediately after hatching (L1 degradation), at 7 h of development (L2 degradation) or at 19 h of development (L3 degradation). Every hour, from placing the worms on plates until seam cell fusion on control plates lacking auxin, a sample of animals were imaged by spinning disc confocal microscopy. The number of seam cell and hyp7 nuclei were counted manually.
Microscopy
Live imaging of *C. elegans* larvae was done by mounting larvae on 5% agarose pads in a 10 mM Tetramisole solution in M9 buffer to induce paralysis. DIC imaging was performed with an upright Zeiss AxioImager Z2 microscope using a 63× 1.4 NA objective and a Zeiss AxioCam 503 monochrome camera, driven by Zeiss Zen software. Spinning disk confocal imaging was performed using a Nikon Ti-U microscope driven by MetaMorph Microscopy Automation & Image Analysis Software (Molecular Devices) and equipped with a Yokogawa CSU-X1-M1 confocal head and an Andor iXon DU-885 camera, using 60× or 100× 1.4 NA objectives. All stacks along the z-axis were obtained at 0.25 μm intervals, and all images were analyzed and processed using ImageJ (FIJI) and Adobe Photoshop. For quantifications, the same laser power and exposure times were used within experiments.

Quantitative image analysis
All image analysis was done in using ImageJ (FIJI). For intensity profile measurements of spinning disk microscopy data, background values were subtracted from the intensity measurements. Mean background intensity was quantified on a circular region in an area not containing any animals, except in quantifications in Fig. 4A, 4C and 6A, where background intensity was quantified on a circular region in an area with no fluorescence inside the worm.

For the intensity profiles in the epidermis, except those of RFP::GIP-1, a 10 px-wide line was drawn in the apical focal plane, from the hyp7 cytoplasm to the seam cell cytoplasm. For the intensity profiles of RFP::GIP-1, 10 20-px wide lines were drawn per cell in the apical focal plane, from the hyp7 cytoplasm to the seam cell cytoplasm, and then averaged to obtain a single intensity profile per cell. For the intensity profiles in the intestine, 8 50 px-wide lines were drawn from the lumen to the cytoplasm of the intestinal cells and averaged to obtain a single value per worm. The intensity profiles were manually aligned at the apical peak value.

To quantify the fluorescence intensity for the molting assay, whole worm fluorescence was quantified. An ROI of each whole worm was created by drawing a freehand line around the worm using the transmitted light channel. The corresponding fluorescence of the ROI was measured in the GFP channel.

Microtubule bundles were counted manually. A 5-px-wide freehand line was drawn along the worm at the dorsal/ventral region, and the intensity profile was obtained. The number of fluorescent peaks was counted, and the microtubule bundle density was calculated by dividing the number of peaks by the measured distance.

EBP-2::GFP comet counting was done manually. Comet density was calculated by dividing the number of EBP-2::GFP comets by the surface of the area analyzed. Comets in both the lateral and dorsal/ventral hypodermis were measured.

PTRN-1::GFP puncta counting was done manually. Puncta density was calculated by dividing the number of PTRN-1::GFP puncta by the surface of the area analyzed. Puncta in both the seam cells and the hypodermis were measured.

Microtubule growth rate was calculated in an automated manner using the ImageJ plug-in ‘TrackMate’ (Tinevez et al., 2017). The following parameters were chosen: estimated blob diameter = 0.700 um; threshold = 200,000; simple LAP tracker; linking max distance = 1.5 um; gap-closing max distance = 1.5 um; gap-closing max frame gap = 3; duration of track = 10. The mean speed of the comets was averaged to obtain the average microtubule growth rate. Comets in both the seam cells and the hypodermis were measured.

To determine the directionality of the actin bundles and microtubule growth, images or movies were rotated to orient the seam cells horizontally. Lines were drawn along the microtubule or actin bundles, and the angle of these lines was calculated relative to the horizontal axis. Movies of EBP-2 were used to calculate the directionality of microtubule growth, where the direction of growth of individual comets was annotated manually. Maximum projections of EBP-2 movies were used to calculate the directionality of microtubule growth. Rose plots were generated using MatLAB.

Seam lineage analysis
To generate the seam cell lineage the number of seam cells and hyp7 nuclei were analyzed at 1 h intervals, using the marker *oulIs10[scmp::NLS::tdTomato; dpy-7p::2xNLS::YFP; wrt-2p::GFP::PH]* that marks the seam nuclei in red and the hypodermal nuclei in green. Animals were classified according to showing a wild-type seam cell division pattern, having developmental defects such as delayed or arrested seam cell divisions, or having inappropriate seam cell differentiation. Control animals were classified at each larval stage. PAR-6 depleted animals were classified after they had undergone the delayed L2-stage divisions. From the total number of worms analyzed, the percentages of worms in each category were calculated.

PAR-6::mCherry transgenic array
The *Pdpy-7::par-6::mCherry* vector used for PAR-6 hypodermal rescue was cloned into the pBSK(+) vector
using Gibson assembly. The promoter of dpy7, which is expressed in hyp7 but not in the seam cells (Gilleard et al., 1997; Myers and Greenwald, 2005), was amplified from C. elegans genomic DNA using primers 5'-TGAATACGACTATAAGGGCAATTTGcGcTcGtCtgggtc and 5'-tctgaagaataaagttaaagat. A fragment of 5.3 kb containing the entire genomic sequence of par-6 and a fragment of 402 bp of the par-6 3' UTR were amplified from C. elegans N2 genomic DNA using primers 5'-tttaagaatattctttacattttgttccagaATGTGTCCTACAACGGCTCTTA and 5'-GGCCTGTTGTCCTCCTCCCTTGGAATGTCCCTCCTCCACTGTCCGAAT, and 5'-CACTCCACCGGGAATTGAGCAGGCTCTACTTGAAAaacttctgtgageca and 5'-TAAGGGAAACAAAAAGGCTGAGGCTCACGGAaaaattaattttac, respectively. mCherry was amplified from pJJR83 (Addgene #75028) using primers 5'-TCAAGGGAGGGAGGACCTCGTCCATTCTC. Correct amplification and assembly was confirmed by Sanger sequencing. The plasmid generated can be found in Supplementary File 1. To generate transgenic lines young adult hermaphrodites were injected in the germline with 30 ng/µl of Pdpy-7::par-6::mCherry. mCherry fluorescence was used to select stable transgenic lines.

Yeast two-hybrid analysis
Sequences encoding the PAR-6 PDZ domain and full-length NOCA-1d were PCR amplified from a mixed-stage cDNA library using primers par-6_F: 5'-ggaggcgcgccATGATTGTGCCAGAAGCTCATCG, par-6_R: 5'-ggaggcgcgccTCAGGCGTTCGGTGTTCCTTGTT, noca-1d_F: 5'-ggaggcgcgccATGAATATTTGTTGTTGG and noca-1d_R: 5'-ggaggcgcgccATGTGACTCGTCCATTCC. PCR products were digested with AscI and NotI, and cloned into Gal4-DB vector pMB28 and Gal4-AD vector pMB29, respectively (Koorman et al., 2016). The resulting plasmids were transformed into Saccharomyces cerevisiae strains Y8930 (MATα) and Y8800 (MATa) (Yu et al., 2008) using the Te/LiAc transformation method (Schiestl and Gietz, 1989). DB::PAR-6/AD::NOCA-1 diploid yeast was generated by mating, and plated on synthetic defined (SD) medium plates lacking leucine, tryptophan, and histidine containing 2 mM 3-Amino-1,2,4-triazole (3-AT); and lacking leucine, tryptophan, and adenine to assess the presence of an interaction, and on an SD plate lacking leucine and histidine containing 1 mg/ml cycloheximide to test for self-activation by the DB::PAR-6 plasmid in the absence of the AD::NOCA-1 plasmid. Controls of known reporter activation strength and behavior on cycloheximide were also added to all plates.

Statistical analysis
All statistical analyses were performed using GraphPad Prism 8. For population comparisons, a D’Agostino & Pearson test of normality was first performed to determine if the data was sampled from a Gaussian distribution. For data drawn from a Gaussian distribution, comparisons between two populations were done using an unpaired t test, with Welch’s correction if the SDs of the populations differ significantly, and comparisons between >2 populations were done using a one-way ANOVA, or a Welch’s ANOVA if the SDs of the populations differ significantly. For data not drawn from a Gaussian distribution, a non-parametric test was used (Mann-Whitney for 2 populations and Kruskal-Wallis for >2 populations). ANOVA and non-parametric tests were followed up with multiple comparison tests of significance (Dunnett’s, Tukey’s, Dunnett’s T3 or Dunn’s). Tests of significance used and sample sizes are indicated in the figure legends. No statistical method was used to pre-determine sample sizes. No samples or animals were excluded from analysis. The experiments were not randomized, and the investigators were not blinded to allocation during experiments and outcome assessment.

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Author contributions
Conceptualization: V.G.G., H.R.P., M.B.; Methodology: V.G.G., H.R.P.; Formal analysis: V.G.G., H.R.P.; Investigation: R.R.B., A.R., J.K.; Writing - original draft: V.G.G., H.R.P., M.B.; Visualization: V.G.G., H.R.P.; Supervision: M.B.; Project administration: M.B.; Funding acquisition: M.B.

Supplementary materials
This manuscript is accompanied by three supplementary material files: Supplementary Figures.pdf, Supplemental File 1 - DNA files.zip, and Video S1.m4v.

Tables

Table 1 - List of strains used

| Strain | Genotype |
|--------|----------|
| BOX289 | par-6(mib30[par-6::aid::egfp-loxp]) I; ieSi57[egt-3p::TIR1::mRuby::unc-54 3'UTR + Cbr-unc-119(+)]] II |
| BOX570 | pck-3(mib78[egfp-loxp::aid::pck-3]) II; ieSi57[egt-3p::TIR1::mRuby::unc-54 3'UTR + Cbr-unc-119(+)]] II |
| BOX292 | ieSi57[egt-3p::TIR1::mRuby::unc-54 3'UTR + Cbr-unc-119(+)]] II; par-3(mib68[egfp-loxp::aid::pck-3]) II |
| BOX409 | par-6(mib30[par-6::aid::egfp-loxp]) I; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II |
| BOX607 | pck-3(mib78[egfp-loxp::aid::pck-3]) II; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II |
| BOX444 | pck-3(mib78[egfp-loxp::aid::pck-3]) II; mibs48[Pelt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II |
| BOX285 | par-6(mib30[par-6::aid::egfp-loxp]) I; mibs48[Pelt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II |
| BOX506 | par-6(mib30[par-6::aid::egfp-loxp]) I; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II; dlg-1(mib23[dlg-1::mCherry-LoxP]) X; mgIs49[mlt-10::gfp-pest] |
| BOX412 | par-6(mib30[par-6::aid::egfp-loxp]) I; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II; hels63[Pwrt-2::GFP::PH, Pwrt-2::GFP::H2B, Plin-48::mCherry]V |
| BOX490 | ouIs10[Pscm::NLS::tdTomato(pAW584)+Pwrt2::GFP::PH(pAW561)+Pdpy-7::2xNLS::YFP(pAW516)] I; par-6(mib30[par-6::aid::egfp-loxp]) I; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II |
| BOX041 | mibs23 [lgl-1::GFP-2TEV-Avi 10ng + Pmyo-3::mCherry 10ng + lambda DNA 60ng] V |
| BOX553 | pck-3(mib78[egfp-loxp::aid::pck-3]) II; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II; mibs23 [lgl-1::GFP-2TEV-Avi 10ng + Pmyo-3::mCherry 10ng + lambda DNA 60ng] V |
| BOX554 | pck-3(mib78[egfp-loxp::aid::pck-3]) II; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II; par-1(t324[par-1::gfp::par-1 exon 11a]) |
| BOX493 | pck-3(mib78[egfp-loxp::aid::pck-3]) II; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II |
| BOX402 | par-6(mib30[par-6::aid::egfp-loxp]) I; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II |
| BOX494 | mcIs40 [Plin-26::ABDvab-10::mCherry + Pmyo-2::GFP]; par-6(mib30[par-6::aid::egfp-loxp]) I; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] X; hels63[Pwrt-2::GFP::PH, Pwrt-2::GFP::H2B, Plin-48::mCherry]V |
| BOX483 | maph-1.1(mib12[egfp::maph-1.1]) I; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] X; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] X |
| BOX505 | maph-1.1(mib12[egfp::maph-1.1]) I; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] X; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] X |
| BOX592 | maph-1.1(mib12[egfp::maph-1.1]) I; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] X; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] X |
| BOX487 | par-6(mib25[par-6::mCherry-LoxP]) I; ebp-2[he293[ebp-2::egfp]] II; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II |
| BOX580 | ebp-2[he293[ebp-2::egfp]] II; mibs49[Pwrt-2::TIR-1::tagBFP2-Lox511::tbb-2-3'UTR, IV:5014740-5014802 (ctxTi10882 site)] II |
Table 2 - List of sgRNA target sites

| Target  | Sequence                        |
|---------|---------------------------------|
| par-6   | gacgcaaatgacagtgatagTGG         |
| pkc-3   | tgggtctccgacatcattagAGG         |
| par-3 sgRNA 1 | tttcagatcgatcatcatgtCGG   |
| par-3 sgRNA 2 | cacatgcataacggtcgtggTGG      |
| dlg-1   | gcacgctcattagtaaatgTGG         |
| mos IV  | agctcaatcgtgtacttgcgTGG         |
| ebp-2 sgRNA 1 | gcagggcaaattcggcagattgTGG    |
| ebp-2 sgRNA 2 | taccgggataggatgaacgattgTGG  |
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