## 1 Material aging causes centrosome weakening and disassembly 2 during mitotic exit

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### 14 **ABSTRACT**

Centrosomes must resist microtubule-mediated forces for mitotic chromosome 15 16 segregation. During mitotic exit, however, centrosomes are deformed and fractured by 17 those same forces, which is a key step in centrosome disassembly. How the functional 18 material properties of centrosomes change throughout the cell cycle, and how they are 19 molecularly tuned remain unknown. Here, we used optically-induced flow perturbations 20 to determine the molecular basis of centrosome strength and ductility in C. 21 *elegans* embryos. We found that both properties declined sharply at anaphase onset. 22 long before natural disassembly. This mechanical transition required PP2A phosphatase 23 and correlated with inactivation of PLK-1 (Polo Kinase) and SPD-2 (Cep192). In vitro, 24 PLK-1 and SPD-2 directly protected centrosome scaffolds from force-induced 25 disassembly. Our results suggest that, prior to anaphase, PLK-1 and SPD-2 confer 26 strength and ductility to the centrosome scaffold so that it can resist microtubule-pulling 27 forces. In anaphase, centrosomes lose PLK-1 and SPD-2 and transition to a weak, brittle 28 state that enables force-mediated centrosome disassembly.

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#### 30 INTRODUCTION

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Centrosomes nucleate and anchor microtubules that comprise the mitotic spindle, which segregates chromosomes during somatic cell division. Centrosomes are micron-scale, membrane-less organelles containing a structured centriole pair surrounded by an amorphous protein mass called pericentriolar material (PCM). PCM carries out most of the functions of a centrosome, including directing cell polarity, cell migration, and chromosomal segregation (Conduit et al., 2015; Woodruff et al., 2014)

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39 For chromosome segregation, centrosomes must bear microtubule-dependent loads that 40 create tensile stresses. Motor proteins anchored at the plasma membrane attach to and 41 walk along astral microtubules extending from centrosomes. These spatially-fixed motors 42 thus generate cortically-directed pulling forces on centrosomes, and the balance of those 43 forces determines the ultimate position of the mitotic spindle (Colombo et al., 2003; 44 Gonczy et al., 2001; Grill et al., 2001; McNally, 2013; Nguyen-Ngoc et al., 2007). During 45 this time, centrosomes maintain a compact, spherical shape. However, once 46 chromosome segregation is complete and the cell exits mitosis, centrosomes are 47 deformed and fractured by the same microtubule-mediated forces, which is a pronounced event during centrosome disassembly (Megraw et al., 2002; Severson and Bowerman, 48 49 2003). How the cell regulates the structural and material integrity of centrosomes is 50 unclear. One possibility is that an increase in cortical forces during mitotic exit induces 51 centrosome disassembly. In C. elegans embryos, the magnitude of microtubule-mediated 52 pulling forces does increase during the metaphase-anaphase transition. Yet, the same 53 increase in pulling forces also occurs in metaphase-arrested embryos without leading to 54 centrosome deformation or fracture (Labbe et al., 2004). Furthermore, artificially 55 increasing pulling forces via csnk-1 RNAi does not cause premature centrosome 56 disassembly (Magescas et al., 2019; Panbianco et al., 2008). These studies suggest that 57 induction of centrosome deformation and fracture during mitotic exit cannot be sufficiently 58 explained by increased microtubule-mediated forces. An alternative hypothesis is that 59 centrosome mechanical properties significantly change to permit force-driven fracture 60 and dispersal during mitotic exit.

61 PCM provides most of the mass and microtubule nucleation capacity of a centrosome, 62 and it is widely believed to be responsible for bearing microtubule-mediated forces. PCM 63 is dynamic and expands in size and complexity as cells prepare for mitosis. Selfassembly of coiled-coil proteins, such as Cdk5Rap2 (vertebrates), Centrosomin (D. 64 melanogaster) and SPD-5 (C. elegans), creates the underlying structural scaffold of PCM 65 which then recruits "client" proteins that nucleate and regulate microtubules (Conduit et 66 67 al., 2010; Conduit et al., 2014a; Fong et al., 2008; Hamill et al., 2002; Woodruff et al., 68 2017; Woodruff et al., 2015). Formation of such micron-scale scaffolds requires additional regulatory clients like Polo Kinase, Aurora A Kinase, and SPD-2/Cep192 (Conduit et al., 69 70 2014a; Conduit et al., 2014b; Gomez-Ferreria et al., 2007; Hamill et al., 2002; Hannak et 71 al., 2001; Haren et al., 2009; Lee and Rhee, 2011; Pelletier et al., 2004; Zhu et al., 2008). 72 PCM disassembles at the end of each cell cycle, but the mechanism is not well 73 understood. While this process involves microtubule-mediated PCM fracture and reversal 74 of Polo Kinase phosphorylation (Enos et al., 2018; Magescas et al., 2019; Pimenta-75 Margues et al., 2016), it remains unclear how PCM fracture is initiated, which key 76 molecular targets are de-phosphorylated, if these activities are linked, and how dynamic 77 material changes might contribute to the disassembly process.

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79 A material's load-bearing capacity is determined by its ability to resist permanent 80 deformation and fracture upon stress. In materials science, these properties are 81 described as "strength" and "ductility", respectively. Strength is achieved through high 82 affinity bonding and serves to maintain the material's shape but can sometimes sacrifice 83 flexibility. Ductility is achieved through breakage and reformation of sacrificial bonds or 84 localized neighbor exchange, which dissipates energy over time but sacrifices the 85 material's shape. For example, glass requires large forces to deform, but it cannot deform 86 much before shattering; thus, glass has high strength and low ductility. On the other hand, 87 chewing gum is easily deformed, and it will stretch to great lengths before breaking; thus, 88 gum has low strength and high ductility. Materials with the highest load-bearing capacity 89 are those that combine strength and ductility, such as rubbers, polyampholyte gels, and 90 high-entropy alloys (George et al., 2019; Sun et al., 2013). Over time, these properties

91 can change via chemical or physical modifications, which is referred to as "material92 aging".

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94 For the centrosome, it remains unexplored how molecular-level interactions between 95 PCM proteins generate meso-level mechanical properties like strength and ductility and how these properties change with time. The non-covalent interactions between PCM 96 97 scaffold proteins, as well as cross-linking of scaffold molecules by PCM clients, could all 98 contribute. In general, characterizing the mechanical properties of living soft matter—such 99 as cells, organelles, and protein assemblies—has been challenging due to their size (sub-100 micrometer scale) and low abundance (sub-milligram scale). Techniques like atomic 101 force microscopy and optical trapping can be useful in this respect, but they are limited to 102 easily accessible samples, like the outer membrane of cultured cells and reconstituted 103 protein complexes. As a complementary method to actively probe mechanical properties 104 in cells with limited accessibility, we previously used focused light-induced cytoplasmic 105 streaming (FLUCS)(Mittasch et al., 2018). Specifically, we showed how FLUCS can 106 reveal robust power-law-rheology signatures within the cell cytoplasm, to distinguish between fluid and gel-like states. Fluids undergo unconstrained motion proportional to 107 108 stimulus time (i.e., they flow), while solids and gels undergo only limited deformations, 109 which stall after small amounts of time due to their intrinsic elastic constraints. As FLUCS 110 functions via thermoviscous flows, which develop independent of the absolute viscosity 111 of the fluid (Weinert et al., 2008), FLUCS is particularly suited to distinguish between 112 highly viscous phases and elastic phases. Such a distinction could not be achieved by 113 passive microrheology, as these two states would exhibit the same fingerprint of reduced 114 motion.

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The *C. elegans* embryo is an ideal system to dissect the molecular determinants of PCM load-bearing capacity. First, *C. elegans* has a limited core set of proteins needed for rapid PCM assembly and disassembly, most of which are conserved across eukaryotes: PLK-1 (Polo Kinase homolog), SPD-2 (Cep192 homolog), SPD-5 (functional homolog of Centrosomin and Cdk5Rap2), and LET-92<sup>SUR-6</sup> (PP2A<sup>B55 $\alpha$ </sup> phosphatase homolog) (Decker et al., 2011; Enos et al., 2018; Hamill et al., 2002; Kemp et al., 2004; Magescas

122 et al., 2019; Pelletier et al., 2004; Schlaitz et al., 2007). Second, it is possible to 123 reconstitute PCM assembly and microtubule nucleation *in vitro* using purified *C. elegans* 124 proteins (Woodruff et al., 2017; Woodruff et al., 2015). These experiments previously 125 revealed that PCM forms via self-assembly of SPD-5 into spherical, micron-scale scaffolds that recruit PCM client proteins. SPD-2 and PLK-1 enhance SPD-5 self-126 assembly, while PP2A<sup>SUR-6</sup> removes PLK-1-derived phosphates and promotes PCM 127 128 disassembly. However, these experiments did not reveal the mechanical properties of the 129 SPD-5 scaffold nor how they are tuned in a cell-cycle-dependent manner.

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131 Here, we ask 1) how centrosomes undergo dynamic structural changes to withstand high 132 tensile stresses in mitosis but not during mitotic exit. 2) which mechanical properties are 133 associated with these distinct functional states, and 3) which molecular players and logics 134 regulate the transient existence of centrosomes. To answer these questions, we 135 combined genetics and pharmacological intervention with FLUCS to study the mechanical 136 properties of PCM in *C. elegans* embryos. Our results revealed that PCM transitions from a strong, ductile state in metaphase to a weak, brittle state in anaphase. This mechanical 137 transition is promoted by PP2A<sup>SUR-6</sup> and opposed by PLK-1 and SPD-2. Our data suggest 138 139 that mitotic PCM is a composite of a stable SPD-5 scaffold and proteins that dynamically 140 reinforce the scaffold, such as PLK-1 and SPD-2. During spindle assembly, accumulation 141 of PLK-1 and SPD-2 render the PCM tough enough to resist microtubule-pulling forces. 142 During mitotic exit, departure of PLK-1 and SPD-2 weakens the PCM scaffold to allow 143 microtubule-mediated fracture and disassembly. Thus, PCM undergoes cell-cycle-144 regulated material aging that functions to promote its disassembly.

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#### 147 **RESULTS**

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## 149 FLUCS reveals weakening of the PCM scaffold at anaphase onset in *C. elegans*

150 embryos

To study the molecular determinants of PCM load-bearing capacity, we studied *C. elegans* 1-cell embryos, where growth and disassembly of the PCM scaffold is easily visualized using fluorescently-labeled SPD-5 (mMaple::SPD-5)(Figure 1A). During spindle assembly, the *C. elegans* PCM scaffold is subject to microtubule-mediated pulling forces, but it maintains its spherical shape and structural integrity. However, during telophase, those same pulling forces deform and fracture the PCM scaffold (Figure 1A)(Enos et al., 2018; Severson and Bowerman, 2003). We hypothesized that PCM undergoes an intrinsic mechanical transition from a strong, tough state in metaphase to a weak state in telophase.

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161 We probed PCM mechanical properties using FLUCS-generated flows complemented with fluorescent imaging. Specifically, we equipped a spinning disk confocal microscope 162 163 with a laser control unit (wavelength = 1455 nm) that creates precise, sub-millisecond 164 thermal manipulations (Figure 1B). Unidirectional scans with this laser at 1500 Hz creates 165 travelling temperature fields that are sufficient to induce flows in a viscous medium, 166 including embryonic cytoplasm (Mittasch et al., 2018)(see Methods). A Peltier-cooled 167 stage insert dissipates excess heat to keep the sample within its physiological 168 temperature range.

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170 We applied FLUCS to *C. elegans* embryos expressing mMaple::SPD-5 and mCherry-171 labeled histones (mCherry::H2B). For each experiment, we induced flows crossing the 172 cytoplasm and continuing through the middle of a centrosome, which should apply stress 173 to the PCM scaffold orthogonal to microtubule-derived tensile stresses (Figure 1C). 174 Based on previous experiments (Mittasch et al., 2018), we predicted that cytoplasmic 175 shear flows should weaken the PCM by the relative displacement of scaffold proteins and 176 create a virtual "notch" in the flow path. We used three different amplitudes for the 177 scanning infrared laser (25, 32, 40 mW) to generate cytoplasmic flow velocities ranging 178 from 5-20 µm/min (Figure 1D; Movie S1); these flows scaled guadratically with laser 179 power ( $R^2$ = 0.97), as one would expect for a predominantly viscous medium (Figure S1A). 180 Simultaneously, we cooled embryos to 17°C, such that the embryo cytoplasm never 181 exceeded 23°C during laser scanning (C. elegans embryos develop properly at any 182 temperature between 16°C and 25°C)(Begasse et al., 2015).

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As shown in Figure 1E, FLUCS deformed and fractured mature PCM in anaphase 184 185 embryos. We also detected a slight increase in cytoplasmic fluorescence surrounding the 186 PCM during FLUCS, indicating that flows can dislodge SPD-5 from the PCM, as predicted 187 (Figure S1B). FLUCS-induced PCM deformation differed starkly between cell cycle 188 stages. FLUCS was not able to visibly deform PCM during metaphase or prior, even 189 though flows were strong enough to detach the centrosome from the spindle and move it 190 toward the cell cortex or out of the focal plane (Figure 1D and S1C and Movies S2, S3; 191 see Metaphase and Prometaphase). On the contrary, FLUCS deformed and eventually 192 fractured PCM during anaphase and telophase (Figure 1E,F; Movie S4,S5); in these 193 experiments, the untreated centrosome remained intact.

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195 We then quantified 1) the deformation rate, defined by the rate of PCM length change 196 orthogonal to the flow direction, and 2) the fracture probability, defined by the chance that 197 PCM segments detach completely after FLUCS (Figure 1G; see methods). The PCM 198 deformation rate and fracture probability increased with increasing flow velocity and 199 progression through mitosis (Figure 1 H,I). To pinpoint when PCM becomes susceptible 200 to FLUCS-induced deformation, we continuously applied FLUCS to centrosomes starting 201 in metaphase continuing into anaphase. PCM remained spherical and intact during 202 metaphase, but then fractured immediately after anaphase onset, as marked by 203 chromosome segregation (Figure S1D; Movie S6). Our results suggest that PCM 204 resistance to deformation and fracture is high during metaphase, then declines at 205 anaphase onset, ~150s prior to full PCM disassembly in telophase. We refer to this 206 change in PCM mechanical properties hereon as the "PCM weakening transition".

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Since the generation of FLUCS is accompanied by local temperature gradients, we tested if temperature alone affects PCM structure. Bidirectional scanning at 10 kHz creates local temperature gradients without flow, and these conditions did not cause significant PCM deformation or fracture (Figure 1H,I and S1E). Thus, centrosome perturbation during FLUCS is primarily due to the flows and not temperature gradients *per se*. Furthermore, embryos developed properly after cessation of FLUCS (Figure S1F)(Mittasch et al.,

214 2018). We conclude that this established method can be used to study organelles inside215 a living cell.

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### 217 **PCM** weakens in anaphase independent of cortical force generation

218 We next wondered if our FLUCS results could be explained by cell-cycle-dependent 219 changes in PCM mechanical properties or changes in cortical force generation. During 220 the metaphase-anaphase transition in early C. elegans embryos, cortical pulling forces 221 increase to induce transverse oscillations and posterior positioning of the mitotic spindle 222 (Pecreaux et al., 2006). Depleting the proteins GPR-1 and GPR-2 (gpr-1/2(RNAi)) 223 significantly reduces cortical microtubule-pulling forces and prevents spindle 224 displacement, spindle oscillation, and PCM deformation and fracture (Colombo et al., 225 2003; Enos et al., 2018; Grill et al., 2003; Magescas et al., 2019; Pecreaux et al., 2006). 226 Thus, we performed FLUCS in *gpr-1/2(RNAi*) embryos, where we expect only residual 227 cortical forces that remain relatively constant throughout mitosis.

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229 In both wild-type and gpr-1/2(RNAi) embryos, FLUCS deformed PCM in anaphase and telophase, but not in metaphase. However, in gpr-1/2(RNAi) embryos, FLUCS-induced 230 231 PCM deformation rates were slower (Figure 2 A,B; Figure S2A), and FLUCS caused 232 fracturing only in telophase (Figure 2C). These results suggest that PCM mechanical 233 properties change during mitotic exit, but that cortical pulling forces are required to disperse and fracture the pre-weakened PCM scaffold. To further test this idea, we 234 235 applied 10 µg/ml nocodazole to depolymerize microtubules and performed high-flow 236 FLUCS. Under these conditions, FLUCS did not visibly affect PCM during metaphase, 237 but did dislodge SPD-5 protein from the PCM during telophase (Figure 2D); we did not observe stretching or clean fracture of the PCM during any cell cycle stage. These results 238 239 indicate that 1) the interactions between SPD-5 scaffold molecules weaken independent 240 of microtubule-pulling forces during mitotic exit, and 2) microtubule-pulling forces are 241 required for stretching and fracture of the PCM during mitotic exit. To test if microtubule-242 mediated pulling forces could be sufficient to deform PCM already in metaphase, we 243 depleted a negative regulator of GPR-1/2, casein kinase 1 gamma (*csnk-1(RNAi*)), which 244 is reported to increase cortical pulling forces ~1.5-fold (Panbianco et al., 2008). We did

not observe premature deformation or fracture of PCM under these conditions, even
though the spindle rocked violently in metaphase (Figure S2B). We conclude that an
intrinsic mechanical change in the SPD-5 scaffold is the main driver of PCM weakening
during anaphase entry.

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250 We next examined the viscoelastic properties of PCM during anaphase by measuring the 251 time-dependent deformation of PCM during FLUCS and relaxation after FLUCS was 252 turned off. We used *gpr-1/2(RNAi*) embryos to allow residual pulling forces but prevent spindle oscillations, which could complicate our analysis. We observed that continuous 253 254 application of medium and high-flow FLUCS in *gpr-1/2(RNAi)* embryos caused time-255 dependent strain of the PCM scaffold (Figure 2E). Thus, anaphase PCM is ductile and 256 can experience micron-scale structural rearrangements without complete fracture during 257 stress. Such behavior is seen in viscous materials. When we turned off FLUCS, PCM 258 remained in its strained, elongated state, indicating the absence of a dominant elastic 259 element strong enough to return the PCM to its original shape (Figure 2 E,F).

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Overall, our FLUCS experiments suggest that the PCM structurally weakens after metaphase. This weakening transition would presumably facilitate PCM disassembly by enabling microtubule-dependent pulling forces to fracture and disperse the PCM scaffold in telophase.

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### **PCM undergoes stepwise compositional changes following anaphase onset**

267 We next investigated the molecular mechanism underlying the PCM weakening transition, 268 in particular, identifying the specific players that determine the dynamic regulation of PCM 269 strength and ductility. PCM is a heterogeneous assembly of proteins required for its 270 assembly and function (Figure 3A). In particular, two critical regulatory proteins, PLK-1 271 (Polo-like Kinase) and SPD-2 (Cep192 homolog), interact with the scaffold protein SPD-272 5 and enhance its self-assembly into supramolecular structures (Cabral et al., 2019; 273 Decker et al., 2011; Woodruff et al., 2017; Woodruff et al., 2015). PLK-1 and SPD-2, as 274 well as other PCM-localized client proteins, might also reinforce the mature SPD-5

scaffold. On the other hand, loss or inactivation of these proteins could weaken the PCMscaffold.

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278 To analyze PCM composition changes during anaphase, we visualized 9 different GFP-279 labeled PCM proteins relative to a standard PCM marker, mCherry:: y-tubulin. We then 280 measured the integrated density of PCM-localized mCherry and GFP signals during 281 mitosis (Figure 3B). The curves in Figure 3D and Figure S3 represent averages for >10 282 experiments (mean +/- 95% CI). mCherry::γ-tubulin signal peaked ~75-100s after 283 anaphase onset, then declined, indicating its departure from PCM. GFP::v-tubulin 284 behaved similarly, as expected (Figure 3C,D and Figure S3). PLK-1 signal decreased 285 immediately after anaphase onset and was no longer detectable ~100s later. SPD-2 also 286 departed from the PCM prior to  $\gamma$ -tubulin. However, the main scaffold protein SPD-5 287 departed afterward. All other proteins departed coincidentally with  $\gamma$ -tubulin or soon 288 afterward. TPXL-1 and AIR-1 departed in a biphasic manner: an initial loss of signal 289 occurred prior to  $\gamma$ -tubulin departure, then a second phase occurred after  $\gamma$ -tubulin 290 departure. To compare departure kinetics across all experiments, we determined the 291 halfway point of disassembly for each individual GFP and mCherry curve per experiment, 292 then calculated the time differential between halfway points ( $\Delta t_{\text{EXIT}}$ ; Figure 3E). The 293 results for anterior and posterior PCM proteins are summarized in Figure 3F and 3G, 294 respectively. A negative  $\Delta t_{EXIT}$  value indicates GFP::PCM protein departure before  $\gamma$ -295 tubulin, and a positive value indicates departure after  $\gamma$ -tubulin. Our results reveal that 296 PCM composition changes in a stepwise manner during anaphase: PLK-1 departs first, 297 followed by SPD-2,  $\gamma$ -tubulin, TAC-1, and finally SPD-5 and proteins that form tight complexes with SPD-5 (RSA-1, RSA-2). TPXL-1 and AIR-1 were more variable in their 298 299 departure, possibly because they localize both to PCM and microtubules that remain after 300 disassembly of the PCM scaffold (Hannak et al., 2001; Ozlu et al., 2005).

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## Polo Kinase and SPD-2 reinforce the PCM scaffold by increasing its strength and ductility

PLK-1 and SPD-2 are the first proteins to depart the PCM during anaphase, when the
 PCM begins to weaken. Thus, we hypothesized that PLK-1 and SPD-2 normally reinforce

the PCM to achieve full strength and stability in metaphase. If this idea is correct, then acute inhibition of PLK-1 phosphorylation or SPD-2 in metaphase might prematurely weaken the PCM, accelerate its disassembly, or reveal hidden material states not previously visible.

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311 For acute inhibition of PLK-1, we treated semi-permeable embryos (via perm-1(RNAi)) 312 with 10 μM Polo Kinase inhibitor BI-2536 in prometaphase (Carvalho et al., 2011). After 313 2 minutes in drug solution, we applied low, medium, and high-flow FLUCS to centrosomes 314 (Figure 4A-C). Under these conditions, low and medium-flow FLUCS deformed 315 metaphase PCM in BI-2536-treated embryos, in contrast to wild-type embryos (Figure 316 4B); in both cases, PCM fracture did not occur. Under high-flow FLUCS, BI2536 treatment 317 increased PCM deformation rate as much as ~11-fold with fracture occurring only in a 318 minority of the cases (30%) (Figure 4A,C). The fact that PLK-1 inhibition enabled PCM to 319 be deformed easily without necessarily fracturing suggests that PLK-1 mostly determines 320 PCM strength. This experiment also reveals that wild-type PCM is ductile during 321 metaphase; this could not be observed previously because the deformation resistance of 322 wild-type PCM was too high. BI-2536 treatment also caused premature disassembly of 323 the SPD-5 scaffold in metaphase-arrested embryos, consistent with previous findings 324 (Figure 4D)(Cabral et al., 2019). Our results show that continuous PLK-1 activity is 325 needed for PCM to achieve full strength and maintain integrity until chromosome are 326 separated.

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328 Next, we analyzed embryos expressing a temperature-sensitive version of SPD-2 (spd-329 2(or188ts))(Kemp et al., 2004). We mounted spd-2(or188ts) afp::spd-5 embryos in cold 330 media and maintained the sample at 17°C while imaging until prometaphase, then upshifted the embryos to 25°C for 1 min to inactivate SPD-2<sup>or188ts</sup>. We then lowered the 331 temperature to 17°C to perform FLUCS in metaphase as per usual (Figure 4E)(Note: 332 333 because of the local heating caused by FLUCS, the treated centrosome remained at 334 ~23°C throughout the experiments). The absence of fully functional SPD-2 made PCM 335 more susceptible to FLUCS-induced fracture and disintegration at all applied flow 336 velocities (Figure 4E-G). Even in PCM that did not fracture into observable pieces, the

337 GFP::SPD-5 signal decayed after application of FLUCS (Figure 4F). We did not observe this phenotype in wild-type PCM or spd-2 mutant PCM not treated with FLUCS (Figure 338 339 4F). Our interpretation of this data is that SPD-2 is required for PCM ductility and strength. 340 Without SPD-2, PCM becomes brittle and susceptible to fracture and diffusion-driven 341 departure of constituents after modest mechanical agitation. In line with this view, 342 inactivation of SPD-2 caused premature disassembly of the SPD-5 scaffold in early 343 anaphase, even without FLUCS perturbations (Figure 4H). Deformation rates were 344 difficult to measure because rapid fracture and vanishing GFP::SPD-5 signal precluded a flow analysis. We conclude that both Polo Kinase and SPD-2 help PCM achieve 345 346 maximal strength and ductility to prevent disassembly.

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348 We next used a minimal in vitro system to test if PLK-1 and SPD-2 directly affect the 349 mechanical properties of the SPD-5 scaffold. When incubated in a crowded environment 350 (e.g. >4% PEG), purified recombinant SPD-5 assembles into micron-scale condensates 351 that recruit PLK-1, SPD-2, and other PCM proteins (Woodruff et al., 2017). We could not 352 assess SPD-5 condensates using FLUCS because the condensates were propelled 353 guickly away from the flow path (data not shown); thus, our simplified in vitro conditions 354 do not exactly match those found in native cytoplasm. As another way to assess the 355 strength of SPD-5 interactions, we induced disassembly of young RFP-labeled SPD-5 356 condensates (500 nM SPD-5::RFP; 5 min after formation) through application of pipetting 357 shear forces and dilution, then measured the amount of condensates that survived using 358 fluorescence microscopy (Figure 5A)(note: dilution is required to prevent SPD-5 re-359 assembly; thus, this assay tests resistance to disassembly only)(Enos et al., 2018). This 360 treatment completely disassembled condensates composed solely of SPD-5 (Figure 361 5B,C). Addition of constitutively active PLK-1 (PLK-1<sup>CA</sup>; T194D T-loop phospho-mimic) or 362 SPD-2 prevented SPD-5 condensate disassembly, with the combination of the two yielding the greatest protection (Figure 5B,C). Kinase-dead PLK-1 (PLK-1<sup>KD</sup>; K67M 363 364 mutant) did not promote SPD-5 condensate survival. These results suggest that PLK-1 365 phosphorylation of SPD-5, along with direct binding of SPD-2, reinforce the interactions 366 between SPD-5 molecules and thus enhance the ability of PCM to resist disassembly.

367 Our *in vitro* and *in vivo* data together suggest that PLK-1 and SPD-2 tune PCM load-

- 368 bearing capacity by conferring strength and ductility to the SPD-5 scaffold.
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## Phosphatase PP2A<sup>SUR-6</sup> promotes PCM disassembly by compromising scaffold ductility

We next investigated how embryos promote the PCM weakening transition during anaphase. PP2A phosphatase localizes to the PCM and plays multiple roles in centriole biogenesis, spindle assembly, and mitotic exit (Wlodarchak and Xing, 2016). The *C. elegans* homolog of PP2A (LET-92) complexed with the B55 $\alpha$  regulatory subunit SUR-6 (PP2A<sup>SUR-6</sup>) is required for complete PCM disassembly (Enos et al., 2018; Magescas et al., 2019). We thus tested if PP2A<sup>SUR-6</sup> drives PCM disassembly by compromising the mechanical properties of PCM.

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380 We treated semi-permeable one-cell embryos with 10  $\mu$ M PP2A inhibitor (LB-100) in 381 metaphase, then performed high-flow FLUCS in anaphase. Unlike in wild-type embryos, 382 where PCM fractured guickly after high-flow FLUCS, the PCM in LB-100-treated embryos 383 stretched orthogonal to the induced flow but resisted fracture (Figure 6A). PCM 384 deformation velocity was ~2-fold higher (0.26 vs. 0.12 µm/min), initially suggesting that 385 PCM is easier to deform when PP2A is inhibited (Figure 6B). However, PP2A inhibition 386 also elevated the ductility of PCM 1.5-fold (final length divided by the original length) and 387 lowered the fracture probability >2-fold in all cell cycle stages (Figure 6C-D). In 2/10 388 anaphase embryos treated with LB-100, PCM stretched as much as 4-fold in length after 389 FLUCS, reaching up to 10 µm while staying connected. PCM was also more resistant to 390 fracture in embryos depleted of the PP2A regulatory subunit SUR-6 (Figure S4A-C). 391 These results suggest that, when PP2A is inhibited, the ductile nature of PCM is 392 preserved throughout anaphase, allowing PCM to absorb more energy overall without 393 fracturing. This is likely achieved through "self-healing", or the breakage and reformation 394 of weak inter-scaffold interactions. The increase in deformation velocity may then result 395 from ductile PCM becoming easier to stretch as it becomes more extended. On the other 396 hand, wild-type PCM is brittle during anaphase and can only be extended short distances 397 before fracturing. We conclude that PP2A normally functions to eliminate "self-healing"

398 PCM scaffold interactions, thus making PCM brittle during anaphase and susceptible to 399 microtubule-mediated fracture in telophase. Consistent with this conclusion, LB-100 400 inhibition of PP2A or depletion of its regulatory subunit SUR-6 inhibited SPD-5 scaffold 401 disassembly in telophase (Figure 6E) (Enos et al., 2018). We speculate that PCM may 402 be less porous in this mutant ductile state compared to the wild-type brittle state, which 403 could delay PCM disassembly further by preventing access of additional disassembly 404 machinery and/or delaying the departure of PLK-1 and SPD-2. Consistent with the latter 405 concept, both let-92 RNAi and sur-6 RNAi impaired SPD-2 and PLK-1 departure from 406 PCM during anaphase (Figure S4D-G) (Magescas et al., 2019).

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408 To determine when and where PP2A might dephosphorylate PCM proteins, we visualized 409 embryos expressing GFP::LET-92, the PP2A catalytic subunit in C. elegans (Schlaitz et 410 al., 2007). GFP::LET-92 localized to the PCM and persisted there until SPD-5 scaffold 411 disassembly, approximately 100s after PLK-1 had departed from the PCM (Figure 6F), 412 consistent with previous observations (Magescas et al., 2019). Our results suggest that, 413 during anaphase, Polo Kinase activity at the PCM ceases and PP2A removes Polo-414 delivered phosphates and contributes to SPD-2 departure. This shift in the balance of 415 phosphorylation and dephosphorylation changes the mechanical properties of the PCM, 416 making it more brittle and susceptible to fracture and dissolution.

417

### 418 **DISCUSSION**

419 Mitotic spindle assembly and positioning require that centrosomes bear tensile 420 microtubule-dependent forces without structural failure. As mitosis ends, however, these 421 same forces are sufficient to deform and fracture centrosomes, facilitating their 422 disassembly. Disassembly is essential to release centrioles and avoid accumulation of 423 old centrosomes over successive rounds of cell division. Here, we combined flow-driven 424 mechanical perturbations in vivo with biochemical reconstitution in vitro to determine the 425 molecular mechanisms regulating deformation resistance and fracture resistance of 426 PCM, the outer and most massive layer of a centrosome.

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## 428 PCM mechanical properties, function, and renewal can be achieved through 429 transient reinforcement of the PCM scaffold

430 Using C. elegans embryos as a model system, we found that PCM deformation 431 resistance, fracture resistance, and composition are tuned in a cell-cycle-dependent 432 manner (Figure 7A,B). During metaphase, PCM resists both microtubule-mediated forces 433 and induced flow perturbations without deforming or fracturing. In this state, PCM is 434 structured as a reinforced composite, comprising a non-dynamic scaffold of SPD-5 435 molecules filled with a dynamic phase of regulatory molecules like SPD-2 and PLK-1, which frequently bind and unbind the scaffold (Figure 7B). During anaphase, PCM loses 436 437 PLK-1 and SPD-2 and then becomes susceptible to deformation and fracture. During 438 telophase, PCM is at its weakest and is easily fractured and dispersed by microtubule-439 mediated forces, a hallmark step in the PCM disassembly process.

440

441 Our implementation of flow perturbations in vivo using FLUCS reveals how PLK-1, SPD-442 2, and PP2A contribute to the dynamic mechanical properties of the PCM. We interpret 443 deformation resistance as an indicator of strength and fracture resistance and strain as 444 indicators of ductility. Wild-type metaphase PCM is highly resistant to flow perturbations, 445 but underlying features appear in different mutant states (Figure 7C). When PLK-1 is 446 inhibited, the PCM scaffold is easily deformed by FLUCS and stretches without fracturing. 447 Thus, PLK-1 normally maintains PCM strength. On the other hand, when SPD-2 is 448 inhibited, the PCM scaffold is easily fractured and dissolved by FLUCS but does not 449 stretch. Thus, SPD-2 normally maintains PCM ductility and strength. Elimination of either 450 PLK-1 or SPD-2 causes premature PCM disassembly, suggesting that the combination 451 of strength and ductility is necessary for PCM function and maintenance during spindle 452 assembly in metaphase. In anaphase, wild-type PCM is more easily deformed and 453 fractured by FLUCS. Yet, when PP2A is inhibited, PCM is difficult to fracture by FLUCS 454 and instead stretches up to 4 times its original length, revealing that the high ductility of 455 the PCM, which was established prior to metaphase, is preserved. Thus, PP2A normally 456 functions to drive PCM disassembly by reducing PCM ductility.

457

458 We propose that the balance of PLK-1, SPD-2, and PP2A activities determine the 459 mechanical properties and assembly/disassembly state of the PCM (Figure 7D). In 460 metaphase, PLK-1 phosphorylation of SPD-5 and direct binding of SPD-2 reinforce the 461 SPD-5 scaffold, conferring strength and ductility. During anaphase, PLK-1 and SPD-2 462 depart from the PCM, while PP2A phosphatase remains and removes PLK-1-derived phosphates. As a result, PCM becomes progressively brittle and weak, allowing 463 464 microtubule-dependent forces to deform and fracture it in telophase. Since PLK-1 and 465 SPD-2 stabilize the SPD-5 scaffold, but are themselves dynamic, we call this mode of 466 PCM regulation "transient reinforcement".

467

468 Transient reinforcement of the PCM scaffold, in theory, could enable cell-cycle regulated 469 PCM assembly, function, and renewal. In preparation for mitosis, PCM must rapidly 470 assemble and provide a solid foundation for nucleating and anchoring microtubules. If 471 PCM assembly fails, then mitotic spindle assembly and chromosome segregation is 472 severely impaired (Doxsey et al., 1994; Hamill et al., 2002; Sunkel and Glover, 1988). 473 PLK-1 and SPD-2 thus play dual roles in PCM functionality: 1) catalyzing assembly of the 474 PCM scaffold and 2) strengthening it to withstand microtubule-dependent pulling forces 475 (shown here). While PCM is stable during spindle assembly, PCM disassembles in 476 telophase only to be rebuilt in the next cell cycle. PCM disassembly is essential for entry 477 into various post-mitotic states, including the formation of acentriolar oocytes and heart 478 tissue (Pimenta-Margues et al., 2016; Zebrowski et al., 2015). How might transient 479 reinforcement enable PCM disassembly and renewal? Based on in vivo and in vitro FRAP 480 data, PLK-1 and SPD-2 are mobile within the PCM, suggesting that they frequently bind 481 and unbind the SPD-5 scaffold (Laos et al., 2015; Woodruff et al., 2017). Either 482 decreasing their association or increasing their dissociation rates with the SPD-5 scaffold 483 would reduce SPD-2 and PLK-1 levels at the PCM. One potential mechanism is through 484 localized ubiquitination and degradation, which controls Polo Kinase levels at 485 centrosomes in human tissue culture cells during anaphase (Lindon and Pines, 2004). It 486 is currently unknown how PCM levels of SPD-2 or its homolog Cep192 are tuned. The 487 completeness and speed of SPD-2 and PLK-1 removal during anaphase is also 488 suggestive of feedback. Thus, the system could be set up such that minor changes to

SPD-2 and PLK-1 affinity elicit large, switch-like changes in PCM structure and mechanical properties. Further experiments are needed to define how the other numerous PCM proteins and centriole tethers contribute to PCM mechanical properties. One possible control point is linkage of the SPD-5 scaffold to the centriole via PCMD-1; inactivation of PCMD-1 or laser ablation of centrioles leads to aberrant PCM deformation, presumably because PCM can no longer fully resist microtubule-pulling forces (Cabral et al., 2019; Erpf et al., 2019).

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### 497 **PCM** mechanical properties in other eukaryotes

498 The mechanical properties of PCM in other species have yet to be determined. We 499 speculate that our transient reinforcement model for tuning PCM load bearing capacity 500 may be conserved for three reasons. First, diverse eukaryotic species-nematodes, 501 frogs, flies, and human cells-use both Polo Kinase and SPD-2/Cep192 to enhance 502 assembly of the PCM scaffold (Conduit et al., 2014b; Decker et al., 2011; Haren et al., 503 2009; Joukov et al., 2014; Kemp et al., 2004; Pelletier et al., 2004; Woodruff et al., 2015). 504 Second, PP2A is highly conserved in eukaryotes and is required for mitotic exit in these 505 species (Wlodarchak and Xing, 2016). Third, in *Drosophila* embryos, PCM undergoes 506 cell-cycle-regulated deformation and fracture, termed "flaring", which appears similar to 507 disassembling PCM in C. elegans (Megraw et al., 2002). PCM flares are visible during 508 interphase, cease during metaphase and anaphase, and then escalate during telophase. 509 Flares also require dynamic microtubules. Thus, *Drosophila* PCM flaring may be due to 510 decreasing PCM strength and ductility during telophase, such that PCM can no longer 511 resist microtubule-mediated forces.

512

### 513 Parallels between PCM and common soft materials in engineering

The mechanical properties and structure of mature PCM are analogous to common composite materials such as flexible plastics and hydrogels. Most modern plastics comprise cross-linked polymer chains embedded with plasticizers, chemicals that make the plastic more flexible and ductile. Over time, these plasticizers exit by diffusion, making the remaining plastic brittle and weak, which is a form of material aging. The rubber sole on a shoe will crack with age; flexibility of the old sole can be restored by impregnating

the rubber with a plasticizer such as silicone. For PCM, the SPD-5 scaffold is most similar to the polymer chains, whereas PLK-1 and SPD-2 could act as plasticizers. Similar to aging rubber losing its plasticizers, our results show that the PCM scaffold becomes brittle and weak during anaphase, coincident with both PLK-1 and SPD-2 leaving the PCM.

524

525 PCM is also similar to physical composites like polyampholyte hydrogels, which exhibit a 526 unique combination of high tensile strength and flexibility. Polyampholyte gels comprise 527 polymers cross-linked with a combination of high- and low-affinity non-covalent bonds. 528 Upon stress, the low-affinity bonds break and dissipate energy, while the high-affinity 529 bonds maintain the overall supramolecular structure. The low affinity bonds then guickly 530 reform, resulting in self-healing that prevents structural fatigue from repeated stresses 531 (Sun et al., 2013). These bonds also make the material more ductile, such that it will 532 undergo plastic deformation instead of fracturing. For PCM, it is possible that PLK-1, 533 SPD-2, and other PCM-resident proteins dissipate stress by unbinding from the PCM 534 scaffold, then re-binding to achieve self-healing. Eliminating these weak interactions 535 would make the PCM weaker and more brittle. This would naturally occur in anaphase as 536 PLK-1 and SPD-2 depart from PCM. This concept could also explain why FLUCS induces 537 PCM fracture and deformation in metaphase when we acutely inhibit PLK-1 and SPD-2. 538 Alternatively, in PP2A-inhibited embryos, PCM is ductile due to the preservation of low 539 affinity, self-healing bonds. Although stronger and more ductile than normal, this mutant 540 PCM still weakens and disassembles in telophase, suggesting that another yet unknown 541 process disrupts the strong interactions between SPD-5 molecules.

542

### 543 Conclusion

This work establishes that PCM, the most substantial layer of the centrosome, transitions from a strong, ductile state in metaphase to a weak, brittle state in telophase. This transition is driven by PP2A phosphatase and inactivation of Polo Kinase and SPD-2/Cep-192, which are essential for centrosome assembly and reinforce the PCM scaffold during metaphase. This mode of mechanical regulation, which we term "transient reinforcement", is a functional form of material aging that allows PCM to resist microtubule-mediated tensile stresses during spindle assembly and then to be fractured and disassembled by

similar forces during mitotic exit. Implicitly, our work demonstrates how flow perturbations
 can reveal functional mechanical states of membrane-less organelles in vivo.

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#### 554

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560

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569

## 570 AUTHOR CONTRIBUTIONS

571 M.M., A.W.F., and M.K. built the FLUCS microscope. M.M. and V.M.T. performed and 572 analyzed the FLUCS experiments on *C. elegans* embryos. M.U.R. performed the *spd-2* 573 temperature-sensitive experiments. B.F-G. and S.J.E. performed two-color imaging of 574 PCM proteins during anaphase and wrote the image analysis scripts. A.B. performed 575 imaging of *sur-6(RNAi)* mutants. J.B.W. purified proteins and performed the *in vitro* 576 assays. V.M.T. performed all other in vivo experiments and analysis. J.B.W. and M.K. 577 analyzed data and wrote the manuscript.

578

## 579 **FIGURE LEGENDS**

580 Figure 1. FLUCS reveals changes in PCM deformation resistance and fracture

581 resistance during mitosis in *C. elegans* embryos.

A. Diagram of mitotic progression in a one-cell *C. elegans* embryo (top panels). Force generation (arrows) by cortically anchored microtubules aid in chromosome segregation, spindle positioning, and PCM disassembly during telophase. Livecell confocal microscopy images of *C. elegans* embryos expressing a PCM marker (mMaple::SPD-5) and DNA marker (mCherry::HistoneH2B).

- 587 B. Diagram of the FLUCS microscope setup (left) and generation of intracellular flows 588 after unidirectional scanning of a 1455 nm laser at 1500 Hz. Scan path is 589 represented by the magenta line. The magnitudes of local flow velocities are 590 reflected by arrow size.
- 591 C. Using FLUCS, flows (red arrows) are generated in the cytoplasm and pass through 592 the PCM (green ball). Microtubule-derived pulling forces (grey arrows) also exert 593 tensile stresses on PCM.
- 594 D. Tuning the power of the 1455 nm laser (25, 32, and 40 mW) generates three tiers 595 of flow velocity (LOW, MEDIUM, and HIGH). Individual data points are plotted with 596 mean +/- 95% CI; n = 5 measurements per condition.
- 597 E. Time lapse images of PCM morphology in anaphase after application of no flow 598 (OFF) or low, medium, and high flow. Orange heading boxes indicate when flow 599 occurs. Arrows indicate flow path and direction. Blue heading boxes indicate when 600 flow is turned off.
- 601 F. PCM was subjected to high-flow FLUCS during metaphase, anaphase, and 602 telophase.
- G. For each FLUCS experiment, we measured the change in PCM length over time
   (Deformation rate) and the frequency of complete separations in PCM (Fracture
   probability).
- 606 H. PCM deformation rates were measured in metaphase, anaphase, and telophase 607 using low, medium, and high flow. Individual data points are plotted with mean +/-608 95% CI; n = 7,6,7 (metaphase; LOW, MED, HIGH flow), n = 7,8,7 (anaphase), n = 609 7,8,9 (telophase) measurements per condition. 10 kHz bidirectional scanning of 610 the 1455 nm laser using 40 mW power, generates heat without producing flows 611 (TEMP control; n = 4,5,5 (metaphase, anaphase, telophase)). For high flow, 612 differences are statistically significant using a one-way ANOVA followed by a

613	Tukey's multiple comparison test (metaphase vs. anaphase, p= 0.04; metaphase
614	vs. telophase, p = 0.0001).
615	I. PCM fracture probabilities from experiments in (H). Sample numbers are the same
616	as in (H).
617	
618	Figure 2. PCM undergoes structural weakening during anaphase, independent of
619	cortical pulling forces.
620	A. Time-lapse images of gpr-1/2(RNAi) embryos treated with 40 mW FLUCS (high
621	flow).
622	B. PCM deformation rates in metaphase (M), anaphase (A), and telophase (T) using
623	high flow in wild-type and <i>gpr-1/2(RNAi)</i> embryos. Wild-type data are from
624	experiments in Figure 1. Individual data points are plotted with mean +/- 95% CI;
625	n = 7,7,9 (wild-type; metaphase, anaphase, telophase) and n = 10,12,14 ( <i>gpr</i> -
626	1/2(RNAi)).
627	C. PCM fracture probabilities from experiments in (B).
628	D. Permeabilized embryos were treated in metaphase or telophase with 10 $\mu$ g/ml
629	nocodazole, then subjected to high-flow FLUCS. Representative images are
630	shown on the left, line scans (dotted line in the inset) of fluorescence intensity
631	before and after FLUCS are on the right.
632	E. Zoomed in time-lapse images of PCM deformation under high-flow FLUCS in a
633	<i>gpr-1/2(RNAi)</i> embryo.
634	F. Plots comparing PCM length on the long axis orthogonal to flow direction over time.
635	Both high flow and medium flow induce PCM deformation.
636	
637	Figure 3. Discrete changes in PCM composition correlate with the PCM
638	weakening transition in anaphase.
639	A. Diagram of <i>C. elegans</i> centrosome architecture and composition.
640	B. Worm lines were generated that express mCherry-labeled $\gamma$ -tubulin (as a standard)
641	and 9 different GFP-labeled PCM proteins (left panels). Fluorescence intensity at
642	the PCM was measured over time (right panels).

643 C. Example images from dual-color, time-lapse recording of PCM disassembly in 9 644 different embryo lines described in (B).

D. Quantification of the experiments in B-C. For each strain, the plots represent the 645 646 normalized integrated fluorescence density of PCM-localized mCherry-tagged y-647 tubulin compared to the GFP-tagged protein from anaphase onward. Anaphase was indicated by spindle rocking. Shown are the analyses for anterior-localized 648 649 centrosomes. Data were normalized to the maxima for each individual curve, then 650 these curves were averaged (mean +/- 95% C.I. n = 13 ( $\gamma$ -tubulin), 9 (PLK-1), 10 651 (SPD-2), 12 (SPD-5), 14, (RSA-1), 12 (RSA-2), 7 (TAC-1), 10 (TPXL-1), 13 (AIR-652 1)).

- E. The order of PCM protein departure was determined by calculating the time lag between halfway points of PCM protein departure per strain ( $\Delta t_{EXIT}$ ). Halfway points were determined by fitting each curve during the window of linear departure.
- F. Departure time lag ( $\Delta t_{EXIT}$ ) of GFP-labeled PCM proteins relative to mCherry::γtubulin. A negative value indicates that the GFP-labeled protein departed before γtubulin. A positive value indicates that the GFP-labeled protein departed after γtubulin. Results for anterior centrosomes are shown (mean +/- 95% C.I.; sample number is the same as in (D)). Statistical analyses are shown in Table S3.
- 661 G. Departure time lag ( $\Delta t_{EXIT}$ ) of posterior-localized PCM proteins relative to 662 mCherry::  $\gamma$ -tubulin. Sample number is the same as in (D). Statistical analyses are 663 shown in Table S4.
- 664

# Figure 4. Acute inhibition of PLK-1 and SPD-2 induces premature weakening and disassembly of the PCM scaffold.

- A. PCM was subjected to high-flow FLUCS during metaphase in wild-type embryos
   or permeabilized embryos treated with 10 μM BI-2536 (inhibitor of Polo Kinases).
   Permeabilized embryos behaved as wild-type embryos during the first cell division
   (see methods; Carvalho 2011).
- B. PCM deformation rates in metaphase using low, medium, and high flow in wild-type and BI-2536-treated embryos. Wild-type data are from experiments in Figure
  1. Individual data points are plotted showing mean +/- 95% CI; n = 6-7 (wild-type)

- and n = 5-7 (BI-2536-treated). P-values were calculated using a Mann-Whitney test.
- 676 C. PCM fracture probabilities from experiments in (B).
- D. Permeabilized embryos were arrested in metaphase using 10  $\mu$ M MG-132, then treated with 0.1% ethanol (no drug) or 10  $\mu$ M BI-2536. Data are plotted as normalized lines representing mean +/- 95% CI; n = 8 (no drug) and n = 10 (BI-2536-treated).
- E. Embryos expressing GFP::SPD-5 and a temperature-sensitive version of SPD-2
   (*spd-2(or188ts)*) were allowed to assemble centrosomes at the permissive
   temperature (16°C), upshifted to the non-permissive temperature (25°C) for 1 min
   during prometaphase, then subjected to high-flow FLUCS during metaphase.
- F. For each experiment in *spd-2(or188ts)* embryos, one centrosome was subjected
   to FLUCS and the other left alone (control). Integrated fluorescent intensities of
   the SPD-5 signal were tracked over time, then normalized to the starting value.
   Each curve represents a single experiment.
- 689 G. PCM fracture probabilities using low, medium, and high flow. Wild-type data are 690 reproduced from Figure 1. n = 7,7,8 (wild-type) and 5,6,8 (*spd-2(or188ts)*).
- H. Embryos were upshifted from 16°C to 23°C during metaphase, then imaged during
  anaphase. Data show integrated fluorescence densities of PCM-localized signal,
  plotted as normalized lines representing mean +/- 95% CI; n = 24 centrosomes in
  both wild-type and *spd-2(or188ts)* embryos.
- 695

## Figure 5. Polo Kinase and SPD-2 protect *in vitro* reconstituted PCM from induceddisassembly.

- A. *In vitro* SPD-5 condensate disassembly experiment. 500 nM SPD-5::TagRFP was
   incubated in 9% PEG to induce spontaneous formation of micron-scale SPD-5
   condensates (1. before). After 5 min, the condensates were pipetted harshly and
   diluted 1:10 in PEG-free buffer, incubated for 10 min, then imaged (2. after).
- B. Quantification of total SPD-5 condensate mass per field of view remaining after
   dilution-induced disassembly. Buffer, 240 nM SPD-2, 500 nM constitutively active
   PLK-1 (PLK-1<sup>CA</sup>), and/or 500 nM kinase dead PLK-1 (PLK-1<sup>KD</sup>) were added at the

705	beginning. The plot shows total integrated fluorescence density for each field of
706	view (red bars indicate mean +/- 95% C.I.; n >22 images per experiment).
707	C. Representative images from (B) before and after dilution-induced disassembly.
708	
709	Figure 6. PCM becomes fracture-resistant and ductile in anaphase after inhibition
710	of PP2A phosphatase.
711	A. PCM was subjected to high-flow FLUCS during anaphase in wild-type embryos
712	or permeabilized embryos treated with 10 $\mu$ M LB-100 (inhibitor of PP2A
713	phosphatase).
714	B. PCM deformation rates in anaphase using high flow in wild-type and LB-100-
715	treated embryos. Wild-type data are from experiments in Figure 1. Individual data
716	points are plotted with bars representing mean +/- 95% CI; n = 7 (wild-type) and
717	10 (LB-100-treated) centrosomes. P-values were calculated using a Mann-
718	Whitney test.
719	C. Ratio of final PCM length to original length in experiments from (B). Original PCM
720	length was measured before flow began and final PCM length was measured once
721	flow was turned off. P-values were calculated using a Mann-Whitney test.
722	D. PCM fracture probabilities for high-flow FLUCS in metaphase (M), anaphase (A),
723	and telophase (T). Wild-type data are from experiments in Figure 1; n= 8,11,11
724	(wild-type) and 7,9,6 (LB-100-treated) centrosomes.
725	E. Permeabilized embryos were treated with no drug or 10 $\mu$ M LB-100 in metaphase,
726	then imaged until 300s after anaphase onset. Data are plotted as normalized lines
727	representing mean +/- 95% CI; n = 24 (no drug) and n = 21 (LB-100-treated)
728	centrosomes.
729	F. Dual-color imaging of embryos expressing GFP-tagged LET-92, the PP2A
730	catalytic subunit in C. elegans (GFP::PP2Ac), and SPD-5::mCherry. Data are
731	plotted as normalized lines representing mean +/- 95% CI; n = 10 centrosomes.
732	
733	Figure 7. The balance of PLK-1, SPD-2, and PP2A activities tune PCM strength and
734	ductility.

- A. PCM resistance to microtubule-mediated forces peaks in metaphase during
   spindle assembly, then declines in anaphase and telophase, corresponding to
   PCM disassembly.
- B. PCM-localized levels of 8 different proteins during mitotic progression. In
  anaphase, PLK-1 levels decline first, followed by SPD-2. The catalytic subunit of
  PP2A phosphatase (PP2Ac), as well as the main scaffold protein SPD-5, remain
  at the PCM until late telophase.
- C. In metaphase, FLUCS cannot fracture or deform wild-type PCM. However, FLUCS
  can fracture PCM in *spd-2* mutant embryos (i.e., PCM is less ductile) and stretch
  and deform PCM in PLK-1 inhibited embryos (i.e., PCM is less strong but still
  ductile). In anaphase, FLUCS easily deforms and fractures wild-type PCM, while
  it deforms and stretches PCM in PP2A-inhibited embryos (i.e., PCM is more
  ductile).
- D. The combination of PP2A phosphatase activity and the departure of PLK-1 and
   SPD-2 transitions PCM from a strong, ductile state in metaphase to a weak, brittle
   state in telophase. This transition enables PCM disassembly and dispersal through
   microtubule-mediated pulling forces.
- 752

## 753 SUPPLEMENTAL FIGURE LEGENDS

## 754 **Figure S1. FLUCS control experiments.**

- A. Power law scaling of cytoplasmic flow with increasing FLUCS laser power.
   Individual data points represent mean +/- 95% CI; n = 5 embryos per laser
   condition. Flow velocities were fit with a second-order polynomial.
- B. Application of high-flow FLUCS in an anaphase 1-cell embryo. Images are pseudo colored to highlight the subtle increase in cytoplasmic mMaple::SPD-5
   fluorescence after flow begins. Note: the centrosome goes out of focus in the first
   frame when FLUCS begins.
- C. Application of high-flow FLUCS in a prometaphase 1-cell embryo. Single plane
   images are shown. Flow causes the centrosome to leave the plane of focus at t=0s
   and t=20s. Flow then displaces the centrosome toward the cortex.

D. High-flow FLUCS was applied in metaphase, continuing into anaphase (indicated
 by chromosome segregation at t=12s).

- E. Example images from the experiment in Figure 1H. Bidirectional scanning of a 40
   mW laser (1455 nm) at 10 kHz creates local temperature gradients without
   generating flow.
- F. Time-lapse fluorescence and brightfield images of an embryo after cessation of
   FLUCS. Application of high-flow FLUCS does not affect embryonic development.
- 772

773 Figure S2. Contributions of microtubule pulling forces to PCM deformation.

- 774 A. PCM deformation rates in metaphase (M), anaphase (A), and telophase (T) 775 using high flow in wild-type and *gpr-1/2(RNAi)* embryos or 40 mW bidirectional 776 laser scanning (temperature control; no flow). Data are from experiments in 777 Figures 1 and 2. Individual data points are plotted with mean +/-95% CI; n = 7.7.9 (wild-type; metaphase, anaphase, telophase), n = 10,12,13 (gpr-778 779 1/2(RNAi) and n = 4,5,5 (temperature control). P values were calculated using 780 Brown-Forsythe and Welch ANOVA tests followed by Dunnett's T3 multiple 781 comparisons tests.
- B. Time-lapse images of centrosomes in a *csnk-1(RNAi)* embryo, where microtubule-mediated pulling forces at the cortex are ~1.5-fold elevated compared to wild type (Panbianco et al., 2008). PCM deformation does not occur prematurely in metaphase.
- 786

# Figure S3. Localization profiles of PCM proteins in the posterior embryo during PCM disassembly.

789Quantification of PCM localization in the posterior side of 1-cell embryos in Figure7903. For each strain, the plots represent the normalized integrated fluorescence791density of PCM-localized mCherry-tagged  $\gamma$ -tubulin compared to the GFP-tagged792protein from anaphase onward. Anaphase was indicated by spindle rocking. Data793are normalized to maxima for each individual curve, then averaged; mean +/- 95%794C.I. n = 13 ( $\gamma$ -tubulin), 9 (PLK-1), 10 (SPD-2), 12 (SPD-5), 14, (RSA-1), 12 (RSA-7952), 7 (TAC-1), 10 (TPXL-1), 13 (AIR-1)).

796	
797	Figure S4. FLUCS and localization experiments in <i>sur-6(RNAi)</i> embryos.
798	A. High-flow FLUCS was applied to a centrosome in an embryo depleted of SUR-6,
799	a PP2A regulatory subunit involved in PCM disassembly.
800	B. PCM deformation rates in anaphase during high-flow FLUCS in wild-type and sur-
801	6(RNAi) embryos. Wild-type data are from experiments in Figure 1. Individual data
802	points are plotted with bars representing mean +/- 95% CI; n = 7 (wild-type) and
803	10 ( <i>sur-6(RNAi</i> )) centrosomes.
804	C. PCM fracture probabilities in metaphase (M), anaphase (A), and telophase (T)
805	during high-flow FLUCS experiments. N= 8-11 (wild-type) and 8-17 ( <i>sur-6(RNAi</i> ))
806	centrosomes.
807	D. Images of PCM-localized GFP::PLK-1 in wild-type and <i>sur-6(RNAi)</i> embryos.
808	E. Normalized integrated fluorescence intensity of PCM-localized GFP::SPD-2
809	during anaphase. Data are plotted as mean +/- 95% CI; n= 22 (wild-type) and 32
810	( <i>sur-6(RNAi)</i> ) centrosomes.
811	F. Images of PCM-localized GFP::SPD-2 in wild-type and <i>sur-6(RNAi)</i> embryos.
812	G. Normalized integrated fluorescence intensity of PCM-localized GFP::SPD-2
813	during anaphase. Data are plotted as mean +/- 95% CI; n= 20 (wild-type) and 24
814	( <i>sur-6(RNAi)</i> ) centrosomes.
815	
816	SUPPLEMENTAL MOVIES
817	Movie S1. FLUCS flow control using 25 mW, 32 mW, and 40 mW laser scans at 1.5
818	kHz. Flows were generated in C. elegans 1-cell embryos using three different 1455 nm
819	laser powers (25 mW, 32 mW, and 40 mW).
820	Movie S2. High-flow FLUCS targeting the centrosome in a prometaphase 1-cell
821	embryo. Prometaphase C. elegans embryos expressing mCherry::histoneH2B
822	(magenta) and GFP::SPD-5 (green) were subjected to high-flow FLUCS (40 mW).
823	Images are of a single confocal plane.
824	Movie S3. High-flow FLUCS targeting the centrosome in a metaphase 1-cell
825	embryo. Metaphase C. elegans embryos expressing mCherry::histoneH2B (magenta)

- and GFP::SPD-5 (green) were subjected to high-flow FLUCS (40 mW). Images are of a
- single confocal plane.
- 828 Movie S4. High-flow FLUCS targeting the centrosome in an anaphase 1-cell
- 829 **embryo.** Anaphase *C. elegans* embryos expressing mCherry::histoneH2B (magenta)
- and GFP::SPD-5 (green) were subjected to high-flow FLUCS (40 mW). Images are of a
- single confocal plane.
- 832 Movie S5. High-flow FLUCS targeting the centrosome in a telophase 1-cell
- 833 embryo. Telophase *C. elegans* embryos expressing mCherry::histoneH2B (magenta)
- and GFP::SPD-5 (green) were subjected to high-flow FLUCS (40 mW). Images are of a
- single confocal plane.
- 836 Movie S6. High-flow FLUCS targeting the centrosome during the metaphase to
- 837 anaphase transition in a 1-cell embryo. C. elegans embryos expressing
- mCherry::histoneH2B (magenta) and GFP::SPD-5 (green) were subjected to high-flow
- FLUCS (40 mW) during the metaphase-anaphase transition. Images are of a single
- 840 confocal plane.
- 841

## 842 TABLE S1. C. elegans strains used in this study

Strain name	genotype	Creation method	Origin
DAM858	vie11[pAD676; gfp::tac-1]II	CRISPR	Alexander Dammermann
EU584	spd-2(or188ts) I	mutagenesis	Bruce Bowerman
JWW1	utsw2[mMaple::spd-5] I	CRISPR	This study
JWW13	spd-2(or188ts) I; unc-119(ed9) III; ItSi202[pVV103/ pOD1021; Pspd-2::GFP::SPD-5 RNAiresistant;cb- unc-119(+)]II	Cross of EU584 and OD847	This study
JWW35	ItSi202[pVV103/ pOD1021; Pspd-2::GFP::SPD-5 RNAiresistant;cb-unc-119(+)]II ; unc-119(ed3)III; ddIs44[WRM0614cB02 GLCherry::tbg-1;Cbr-unc- 119(+)]	Cross of OD847 and TH169	This study
JWW64	utsw2[mMaple::spd-5] l; ltls37 [(pAA64) pie- 1p::mCherry::his-58 + unc-119(+)] IV.	Cross of JWW1 and OD95	This study
JWW65	It17[plk-1::gfp+loxP]III ; ItIs37 [(pAA64) pie- 1p::mCherry::his-58 + unc-119(+)] IV; unc-119(ed3) III	Cross of OD2425 and OD95	This study
JWW66	ItSi203[pVV60; Pspd-2::GFP::SPD-2 reencoded; cb-unc-119(+)]II; ItIs37 [(pAA64) pie- 1p::mCherry::his-58 + unc-119(+)] IV; unc-119(ed3) III	Cross of OD824 and OD95	This study

JWW67	unc-119(ed9) III; utsw1[pJWB56; Pspd- 2::GFP::SPD-5(530E, 627E, 653E, 658E) re-	MosSCI, into EG6699	This study
JWW69	encoded; cb-unc-119(+)]II unc-119(ed9) III; ItSi202[pVV103/ pOD1021; Pspd- 2::GFP::SPD-5 RNAiresistant;cb-unc-119(+)]II; ItIs37 [(pAA64) pie-1p::mCherry::his-58 + unc-	Cross of OD847 and OD95	This study
JWW70	119(+)] IV. unc-119(ed9) III; utsw1[pJWB56; Pspd- 2::GFP::SPD-5(530E, 627E, 653E, 658E) re- encoded; cb-unc-119(+)]II; ItIs37 [(pAA64) pie-	Cross of JWW1 and OD95	This study
JWW71	1p::mCherry::his-58 + unc-119(+)] IV. It17[plk-1::gfp+loxP]III; unc-119(ed3)III; ddls44[WRM0614cB02 GLCherry::tbg-1;Cbr-unc- 119(+)]	Cross of OD2425 and TH169	This study
JWW72	vie11[pAD676; gfp::tac-1]II ; unc-119(ed3)III; ddls44[WRM0614cB02 GLCherry::tbg-1;Cbr-unc- 119(+)]	Cross of DAM858 and TH169	This study
JWW89	spd-2(or188ts) I; ItSi202[pVV103/ pOD1021; Pspd- 2::GFP::SPD-5 RNAiresistant;cb-unc-119(+)]II; ItIs37 [(pAA64) pie-1p::mCherry::his-58 + unc- 119(+)] IV.	Cross of JWW13 and OD95	This study
OD2425	It17[plk-1::gfp+loxP]III	CRISPR	Karen Oegema
OD823	ItSi203[pVV60; Pspd-2::GFP::SPD-2 reencoded; cb-unc-119(+)]II; unc-119(ed3) III	MosSCI, into EG6699	Karen Oegema
OD847	unc-119(ed9) III; ItSi202[pVV103/ pOD1021; Pspd- 2::GFP::SPD-5 RNAiresistant;cb-unc-119(+)]II	MosSCI, into EG6699	(Woodruff et al., 2015)
OD95	unc-119(ed3) III; ltls37 [(pAA64) pie- 1p::mCherry::his-58 + unc-119(+)] IV; ltls38 [pie- 1p::GFP::PH(PLC1delta1) + unc-119(+)]	Microparticle bombardment	CGC
TH169	unc-119(ed3)III; ddls44[WRM0614cB02 GLCherry::tbg-1;Cbr-unc-119(+)]	Microparticle bombardment	Anthony Hyman
TH447	unc-119(ed9) III; ddls243[pie-1p::LAP::LET-92; unc-119(+)]; ddls247[pie-1p::SPD-5(synthetic introns, CAI 0.65)::mCherry; unc-119(+)]	Microparticle bombardment	Anthony Hyman
TH530	rsa-1::LAP; unc-119(ed3)III; ddls44[WRM0614cB02 GLCherry::tbg-1;Cbr-unc- 119(+)]	Microparticle bombardment	Anthony Hyman
TH531	rsa-2::LAP; unc-119(ed3)III; ddls44[WRM0614cB02 GLCherry::tbg-1;Cbr-unc- 119(+)]	Microparticle bombardment	Anthony Hyman
TH539	spd-2::GFP; unc-119(ed3)III; ddls44[WRM0614cB02 GLCherry::tbg-1;Cbr-unc- 119(+)]	Microparticle bombardment	Anthony Hyman
TH571	unc-119(ed3)III; ddls12[pie-1p::tpxl-1::GFP;unc- 119(+)]; ddls44[WRM0614cB02 GLCherry::tbg- 1;Cbr-unc-119(+)]	Microparticle bombardment	Anthony Hyman
TH630	ddls44[WRM0614cB02 GLCherry::tbg-1;Cbr-unc- 119(+)]; ddls62[pie-1p::AIR-1(synthetic introns, CAI 1.0)::GFP; unc-119(+)]; unc-119(ed3)III	Microparticle bombardment	Anthony Hyman
EG6699	ttTi5605 II; unc-119(ed3) III; oxEx1578.		CGC

## **TABLE S2.** Protein expression plasmids used in this study

Plasmid	Gene	N-term tag	C-term tag	Origin
name				
JWV11	<i>plk-1(T194D)</i> constitutively active		PreScission-6xHis	(Woodruff et al., 2015)
JWV12	<i>plk-1(K67M)</i> kinase dead		PreScission-6xHis	(Woodruff et al., 2015)
JWV2	spd-5(wt)	MBP- PreScission	PreScission-6xHis	(Woodruff et al., 2015)
JWV3	spd-5(wt)	MBP- PreScission	tagRFP- PreScission-6xHis	(Woodruff et al., 2015)
JWV6	spd-2(wt)	MBP-TEV	TEV-6xHis	(Woodruff et al., 2015)

846

## 847 **TABLE S3. One-way ANOVA and post-hoc tests of anterior PCM disassembly**

## 848 profiles from Figure 3F

Holm-Sidak's				
multiple		0:	0	Adjusted P
comparisons test	Mean Diff.	Significant?	Summary	Value
AIR-1 vs. TPXL-1	-25.55	No	ns	0.2725
AIR-1 vs. RSA-1	-8.310	No	ns	0.9615
AIR-1 vs. RSA-2	-14.38	No	ns	0.8174
AIR-1 vs. SPD-2	38.26	Yes	**	0.0079
AIR-1 vs. SPD-5	-24.73	No	ns	0.2183
AIR-1 vs. TBG-1	8.728	No	ns	0.9598
AIR-1 vs. TAC-1	0.4412	No	ns	0.9963
AIR-1 vs. PLK-1	92.29	Yes	****	<0.0001
TPXL-1 vs. RSA-1	17.24	No	ns	0.8060
TPXL-1 vs. RSA-2	11.16	No	ns	0.9598
TPXL-1 vs. SPD-2	63.81	Yes	****	<0.0001
TPXL-1 vs. SPD-5	0.8171	No	ns	0.9963
TPXL-1 vs. TBG-1	34.28	Yes	*	0.0356
TPXL-1 vs. TAC-1	25.99	No	ns	0.4307
TPXL-1 vs. PLK-1	117.8	Yes	****	<0.0001
RSA-1 vs. RSA-2	-6.074	No	ns	0.9615
RSA-1 vs. SPD-2	46.57	Yes	**	0.0012
RSA-1 vs. SPD-5	-16.42	No	ns	0.8060
RSA-1 vs. TBG-1	17.04	No	ns	0.7693
RSA-1 vs. TAC-1	8.751	No	ns	0.9615
RSA-1 vs. PLK-1	100.6	Yes	****	< 0.0001
RSA-2 vs. SPD-2	52.64	Yes	****	< 0.0001
RSA-2 vs. SPD-5	-10.35	No	ns	0.9598
RSA-2 vs. TBG-1	23.11	No	ns	0.2887
RSA-2 vs. TAC-1	14.83	No	ns	0.8971
RSA-2 vs. PLK-1	106.7	Yes	****	< 0.0001
SPD-2 vs. SPD-5	-62.99	Yes	****	< 0.0001
SPD-2 vs. TBG-1	-29.53	No	ns	0.0940

SPD-2 vs. TAC-1	-37.82	Yes	*	0.0454
SPD-2 vs. PLK-1	54.03	Yes	***	0.0002
SPD-5 vs. TBG-1	33.46	Yes	*	0.0207
SPD-5 vs. TAC-1	25.17	No	ns	0.4085
SPD-5 vs. PLK-1	117.0	Yes	****	<0.0001
TBG-1 vs. TAC-1	-8.286	No	ns	0.9615
TBG-1 vs. PLK-1	83.56	Yes	****	<0.0001
TAC-1 vs. PLK-1	91.85	Yes	****	<0.0001

849

### 850 **TABLE S4. One-way ANOVA and post-hoc tests of posterior PCM disassembly**

### 851 profiles from Figure 3G

Holm-Sidak's multiple				Adjusted P
comparisons test	Mean Diff.	Significant?	Summary	Value
AIR-1 vs. TPXL-1	-9.235	No	ns	0.9002
AIR-1 vs. RSA-1	-36.70	Yes	***	0.0003
AIR-1 vs. RSA-2	-46.04	Yes	****	<0.0001
AIR-1 vs. SPD-2	5.014	No	ns	0.9652
AIR-1 vs. SPD-5	-42.30	Yes	****	<0.0001
AIR-1 vs. TBG-1	-13.68	No	ns	0.5679
AIR-1 vs. TAC-1	-34.04	Yes	**	0.0041
AIR-1 vs. PLK-1	40.38	Yes	****	<0.0001
TPXL-1 vs. RSA-1	-27.46	Yes	*	0.0334
TPXL-1 vs. RSA-2	-36.81	Yes	***	0.0006
TPXL-1 vs. SPD-2	14.25	No	ns	0.6655
TPXL-1 vs. SPD-5	-33.07	Yes	**	0.0028
TPXL-1 vs. TBG-1	-4.446	No	ns	0.9652
TPXL-1 vs. TAC-1	-24.80	No	ns	0.1399
TPXL-1 vs. PLK-1	49.61	Yes	****	<0.0001
RSA-1 vs. RSA-2	-9.346	No	ns	0.9002
RSA-1 vs. SPD-2	41.71	Yes	****	<0.0001
RSA-1 vs. SPD-5	-5.608	No	ns	0.9652
RSA-1 vs. TBG-1	23.02	No	ns	0.0790
RSA-1 vs. TAC-1	2.661	No	ns	0.9652
RSA-1 vs. PLK-1	77.07	Yes	****	<0.0001
RSA-2 vs. SPD-2	51.06	Yes	****	<0.0001
RSA-2 vs. SPD-5	3.739	No	ns	0.9652
RSA-2 vs. TBG-1	32.36	Yes	**	0.0014
RSA-2 vs. TAC-1	12.01	No	ns	0.8399
RSA-2 vs. PLK-1	86.42	Yes	****	<0.0001
SPD-2 vs. SPD-5	-47.32	Yes	****	<0.0001
SPD-2 vs. TBG-1	-18.70	No	ns	0.2609
SPD-2 vs. TAC-1	-39.05	Yes	**	0.0014
SPD-2 vs. PLK-1	35.36	Yes	**	0.0020
SPD-5 vs. TBG-1	28.62	Yes	**	0.0061
SPD-5 vs. TAC-1	8.269	No	ns	0.9307
SPD-5 vs. PLK-1	82.68	Yes	****	<0.0001

TBG-1 vs. TAC-1	-20.35	No	ns	0.2689
TBG-1 vs. PLK-1	54.06	Yes	****	<0.0001
TAC-1 vs. PLK-1	74.41	Yes	****	<0.0001

852

## 853

### 854 **METHODS**

- 855 Contact for reagent and resource sharing
- 856 Further requests and information for resources and reagents should be directed to and
- 857 will be fulfilled by the Lead Contact, Jeffrey Woodruff
- 858 (Jeffrey.woodruff@utsouthwestern.edu).
- 859

## 860 Experimental model and subject details

861 C. elegans worm strains were grown on nematode growth media (NGM) plates at 16-23°C, following standard protocols (www.wormbook.org). Worm strains used in this study 862 are listed in Table S1 and created using CRISPR (Paix et al., 2015; Paix et al., 2017), 863 864 MosSCI (Frokjaer-Jensen et al., 2008), or microparticle bombardment. Cas9 enzyme was 865 purified by the Protein Expression Facility at MPI-CBG. For expression of recombinant 866 proteins, we used suspended SF9-ESF S. frugiperda insect cells grown at 27°C in ESF 867 921 Insect Cell Culture Medium, Protein-Free (Expression Systems), supplemented with 868 Fetal Bovine Serum (2% final concentration).

869

## 870 RNAi treatment

871 RNAi was done by feeding using sur-6, gpr-2, csnk-1, and perm-1 feeding clones from 872 the Ahringer and Vidal collections (Source BioScience)(Rual et al., 2004). The spd-5 873 feeding clone targets a region that is reencoded in our MosSCI transgenes (Woodruff et 874 al., 2015). Bacteria were seeded onto nematode growth media (NGM) supplemented 875 with 1 mM isopropyl  $\beta$ -D-1-thiogalactopyranoside (IPTG) and 100  $\mu$ g mL<sup>-1</sup> ampicillin. 876 For *perm-1* feeding plates, 0.1 mM IPTG was used. L4 hermaphrodites were grown at 877 23°C for 24-28 hours for all conditions except for perm-1, which was at 20°C for 18-19 878 hours.

879

## 880 Drug treatment of semi-permeable embryos

881 For all drug treatments, C. elegans embryos were permeabilized using perm-1 RNAi 882 (Carvalho et al., 2011) and dissected into a 62% solution of ESF-921 Media (Expression 883 Systems). To arrest the embryos at metaphase, MG-132 (EMD Millipore) was used at 10 884 µM in 62% ESF, diluted from a 10 mM stock concentration in EtOH. To inhibit PLK-1, 885 BI-2536 (Advanced ChemBlocks Inc.) was used at 10 µM, diluted from a 10 mM stock 886 in ethanol. To inhibit PP2A, LB-100 (SelleckChem) was used at 10 µM, diluted from a 887 10 mM stock in dH<sub>2</sub>O. Nocodazole (Sigma) was diluted from a 5 mg/ml stock 888 concentration in DMSO. Samples were flushed with water or M9 after each experiment 889 to test for permeability (water will cause swelling and M9 will cause shrinking).

890

### 891 Construction of the FLUCS Microscope

To measure the physical material state of centrosomes inside living *C. elegans* embryos, we performed intracellular flow perturbations by employing the previously published technology FLUCS (Mittasch et al., 2018). The FLUCS setup consisted of three major modules: (i) an infrared laser scanning unit for thermal manipulations, (ii) a microscope allowing to simultaneously induce thermal patterns and to perform high-sensitivity fluorescence imaging, and (iii) a heat management stage.

898 (i) The infrared laser scanning unit consists of a fiber-based infrared Raman laser (CRFL-899 20-1455-OM1, 20 Watts, near TEM00 mode profile, Keopsys, France) with a wavelength 900 of 1455 nm, operated in continuous-wave mode and linearly polarized using a polarizing 901 beam splitter cube (CCM1-PBS254, Thorlabs, USA). To precisely correct for the 902 divergence of the laser beam, a telescope was used, composed of two telescope lenses 903 with focal lengths of f 1=100 mm and f 2=150 mm (AC254-C series, Thorlabs, USA), 904 respectively. A lambda-half plate (waveplate, 1/2 1550 Edmund optics, USA) was used 905 to rotate the linearly polarized laser light to match the optical axis of the acoustic-optical 906 deflector (AOD). A variable optical beam expander (4x expander, 36100, Edmund optics, 907 USA) allows control of the beam diameter (~1.5 mm beam diameter at back-focal-plane 908 was used) without changing the size of the scan pattern. Rapid (up to 1 MHz update rate) 909 and precise (down to 100 nm) infrared laser scanning was achieved by utilizing two-910 dimensional AOD (AA.DTSXY-A6-145, Pegasus Optik, Germany), electronic oscillators (AA.DRFAI0Y-B-0-x, Pegasus Optik, Germany), and electronic amplifiers 2.5 W 911

912 (AA.AMPA-B-34-20.4, Pegasus Optik, Germany). The AOD was controlled by generating 913 analog signals using a custom software in LabVIEW (National instruments, USA) in 914 combination with a PCI controller card (PCIe 6369, National Instruments, USA). To 915 precisely translate the AOD-induced beam scanning into the back-focal-plane of the 916 microscope objective lens a telescope composed of two telescope lenses with focal 917 lengths of f 3=f 4=300 mm (AC254-C series, Thorlabs, USA) was used. A dichroic mirror 918 (F73-705, AHF, Germany) was used to couple the infrared laser beam into the light path 919 of the microscope (IX83, Olympus, Japan), by selectively reflecting the infrared light but 920 transmitting visible wavelengths which were used for fluorescence imaging.

921

922 (ii) The microscope was equipped with Brightfield (BF) and fluorescent imaging optics. 923 For simultaneous high-resolution fluorescence imaging and precise infrared laser 924 scanning, an infrared-coated microscope objective lens (60x UPLSAPO NA=1.2, W-IR 925 coating. Olympus, Japan) was used, which was operated with heavy water (D2O) as 926 immersion liquid to reduce undesired infrared laser light absorption in the immersion 927 layer. For Brightfield illumination a high-power LED (M565L3, Thorlabs, USA) in 928 combination with dedicated LED driver (LEDD1B, Thorlabs, USA) was used. For confocal 929 fluorescence imaging, a VisiScope confocal imaging system (Visitron, Germany) coupled 930 to a Yokogawa CSU-X1- A12 scan head and an iXON Ultra EMCCD camera (Andor, 931 Ireland) were used.

932

933 (iii) The heat management unit consisted of a thin sample mounting chamber based on a 934 standard cover slip  $(18 \times 18 \times 0.17 \text{ mm})$  (Menzel, Germany) facing the objective lens, and 935 a high thermal conductive sapphire cover slide (thermal conductivity of 27.1  $W/m \cdot K$ , 936 SMS-7521, UQG Optics, UK) closing the sandwich-like chamber from the top. To 937 efficiently remove the induced heat from the samples, e.g. C. elegans embryos, the 938 sapphire slide was actively cooled from room temperature to 17 °C. This active cooling 939 was performed by using Peltier elements (TES1-127021, TEC, Conrad) glued to the 940 sapphire slides. The cooling power of the Peltier elements was controlled by a PID 941 hardware controller (TEC-1089-SV, Meerstetter Engineering, Swiss). A custom-built 942 water-cooling stage was used to dissipate the heat produced by the Peltier elements. The

height of the buffer-filled chamber was defined using polystyrene beads (Polybead,

Polysciences, Germany) with a diameter of  $15 \mu m$ . The height of the resulting chamber

945 was measured by locating the upper and lower chamber surface using a piezo stage.

946

### 947 Application of FLUCS within embryos

948 Late L4 hermaphrodites were grown for 18-19 hours on standard NGM or *perm-1* feeding 949 plates. Worms were then dissected on an 18 mm x 18 mm coverslip (0.17 mm thickness) 950 in 6 µL of M9 buffer or 62% ESF 921 (for permeabilized embryos) with 15 µm polystyrene 951 beads. The sample was placed onto a sapphire microscope slide equipped with Peltier 952 cooling elements, then the coverslip sealed with dental silicone (Picodent twinsil, Picodent, 953 Germany). The cooling stage and sample were then mounted on the FLUCS microscope 954 stage. Embryos were identified and staged using a 10x air objective, then imaged with a 955 60x 1.2 NA Plan Apochromat water immersion objective (Olympus) using 488 nm and 956 561 nm laser illumination, 1X1 binning, and 2s intervals.

957 Hydrodynamic flows were generated by scanning the 1455 nm laser through either 1) 958 center of centrosome or 2) through the cytoplasm for velocity calibration. Custom-written 959 LabVIEW software superimposes the scan path of the infrared laser with the high-960 resolution image of the camera. The sub-pixel alignment of the induced flow field and the 961 camera image was verified routinely before the embryonic experiments by using 962 fluorescent tracer particles immersed in a highly viscous sucrose solution. FLUCS 963 experiments used unidirectional but repeated laser scans with  $1.5 \ kHz$  scan frequency, a 964 scan length of 30 µm, and three different laser powers (25 mW, 32 mW, and 40 mW).

965 Centrosomes were targeted for FLUCS at metaphase, anaphase, or telophase. 966 Centrosomes were affected by FLUCS between 30-60 s. For experiments requiring drug 967 treatment, worms were dissected in 6 µL of the specific drug solution and quickly placed 968 on the microscope within 1-2 minutes. To maintain consistency of drug treatment 969 duration, only embryos found exactly at prometaphase (for metaphase experiments) and 970 metaphase (for anaphase experiments) were then targeted for FLUCS. Temperature-971 sensitive worms were dissected in cold 62% ESF-921 media on a cooled dissecting scope 972 and quickly mounted onto the cooling stage, which was maintained at 17°C. At

973 prometaphase, temperature was upshifted to 25°C for 1 minute, then decreased to 17°C.
974 Centrosomes were then targeted for FLUCS at metaphase.

975

### 976 Confocal microscopy and live-cell imaging

977 Adult worms were dissected in M9 before being mounted on a 5% agar pad for imaging. 978 For live cell imaging with drug treatments, perm-1 adult worms were dissected in 8-10 µL 979 of 62% ESF 921 with 15 µm polystyrene beads (Sigma-Aldrich) on a 22 x 50 mm 980 coverslip. Samples were mounted on a 1 mm thick glass slide with 2 x 6 mm laser cut 981 holes 30 mm apart (Potomac), to produce a flow chamber. In one open chamber, 40 µL 982 of the drug solution in 62% ESF was added during prometaphase. Liquid was wicked 983 from the opposite chamber using a Kimwipe to then allow more of the drug solution to be 984 added to the sample. To arrest embryos in metaphase, perm-1 adult worms were 985 dissected in 10 µM MG-132 solution. Cell cycle stage was indicated based on mCherry::HIS-58 fluorescence and cell morphology (metaphase = aligned chromosomes; 986 987 anaphase = chromosomes separate; telophase = chromosomes de-condense and 988 cytokinetic furrow ingresses).

989

990 Time-lapse images were taken using an inverted Nikon Eclipse Ti microscope with a 991 Yokogawa spinning disk confocal head (CSU-X1), piezo Z stage, and an iXon Ultra 992 EMCCD camera (Andor), controlled by Metamorph software. On this system, the 60x 1.4 993 NA Apochromat oil objective was used to acquire 36 x 0.5 µm Z-stacks every 10 seconds 994 with 100 ms exposures and 2X2 binning. For PCM localization in csnk-1(RNAi) embryos, 995 and PP2A localization, time-lapse images were acquired with an inverted Nikon Eclipse 996 Ti2-E microscope with a Yokogawa confocal scanner unit (CSU-W1), piezo Z stage, and 997 an iXon Ultra 888 EMCCD camera (Andor), controlled by Nikon Elements software. For 998 most experiments, we used a 60x 1.2 NA Plan Apochromat water immersion objective to 999 acquire 35 x 0.5 µm Z-stacks every 10 seconds with 100 ms exposures and 2X2 binning. 1000 Simultaneous imaging with the 488 nm and 561 nm lasers was achieved using an 1001 OptoSplit II beam splitter (Cairn). For LET-92::GFP imaging, a 100x 1.35 NA Plan 1002 Apochromat silicone oil objective was used to acquire 11 x 0.5 µm Z-stacks in 20 second 1003 intervals with 100 ms exposures and 2X2 binning. Images in Figure 3 were taken using
an inverted Olympus IX81 microscope with a Yokogawa spinning-disk confocal head
(CSU-X1), a 60x 1.2 NA Plan Apochromat water objective, and an iXon EM + DU-897 BV
back illuminated EMCCD (Andor).

1007

### 1008 spd-2(or188ts) temperature shift assay

1009 JWW69 (control) and JWW89 (spd-2(or188ts)) strains were used for imaging. 1010 Sequencing of JWW89 confirmed a single point mutation in *spd-2* resulting in a glycine to glutamic acid amino acid substitution (G615E) as described in Kemp et. al. Both worm 1011 1012 strains propagated at 16°C, which is the permissive temperature for spd-2(or188ts). To 1013 prepare the embryos for imaging, a metal block was buried halfway in wet ice. A 24x60 mm glass coverslip (thickness of 1) and a flow chamber slide were placed over the cold 1014 1015 block. To prevent sticking of the glass to the cold block due to water condensation, two 1016 Kimwipes were placed between the glass and the cold block. To minimize exposure to 1017 elevated temperatures during embryo dissection, the glass stage on the dissecting 1018 microscope stand was placed in a 4°C fridge and left to cool for approximately 10 min. 1019 Once everything was cold, 10µL of cold M9 plus 15 µm polystyrene (Sigma) beads was 1020 pipetted to the middle of the 24x60 mm cover slip.

1021

1022 For each worm strain, plates were transported inside the ice bucket directly contacting 1023 ice to the dissecting microscope area. The microscope glass stage was taken out from 1024 the fridge and assembled into its place. Three to four adult worms containing a single row 1025 of eggs were transferred to the M9 plus beads on the cover slip still located on top of the 1026 cold block. The coverslip was transferred to the dissecting scope and the worms cut open 1027 using 22G needles. The coverslip was mounted on the flow chamber slide, then the edges 1028 of the cover slip were sealed using clear nail polish. The sample was moved to the 1029 imaging room on the cold block.

1030

1031 The Nikon Eclipse Ti2 microscope described above was used for imaging. Embryos were 1032 staged using a 10X air objective, then imaged with a 60X NA 1.2 water objective. To 1033 rapidly raise the temperature of the sample (up-shift), 40  $\mu$ l of 25°C M9 was pipetted into 1034 the flow chamber well. 30 x 0.5  $\mu$ m Z stacks were collected every 10 s using simultaneous

illumination with 488 nm and 561 nm lasers (14.7% and 17.7% intensity respectively),2x2 binning, 100 ms exposures.

1037

#### 1038 **Protein expression and purification**

All expression plasmids are listed in Table S2. SPD-5, SPD-2, and PLK-1 proteins were expressed using the FlexiBAC baculovirus system (Lemaitre et al., 2019) and purified as previously described (Woodruff and Hyman, 2015; Woodruff et al., 2015), with the following exception: SPD-2 was stored in its uncleaved form (MBP-TEV-SPD-2).

1043

#### 1044 *In vitro* SPD-5 condensate disassembly assay

1045 SPD-5 condensates were formed by diluting 10 µM SPD-5 (1:10 mixture of SPD-5 and SPD-5::TagRFP) in Condensate buffer (25 mM HEPES, pH 7.4, 150 mM KCI) containing 1046 polyethylene glycol 3350 (Sigma) and fresh 0.5 mM DTT. Before use, the SPD-5 stock 1047 solution was centrifuged for 5 min at 80,000 rpm to remove residual aggregates. 5 min 1048 1049 after formation, SPD-5 condensates were placed in glass-bottom 96-well dishes (Corning, 1050 4850, high content imaging dish) pre-cleaned with 2% Hellmanex and washed in water. 1051 For each sample, half was placed in the well undisturbed (control), and the other half was 1052 diluted 10-fold, pipetted 5 times, then placed in a well (induced disassembly). 96-well 1053 plates were imaged on an inverted Nikon Ti-E microscope using a 60x NA 1.4 Plan 1054 Apochromat oil objective, a Zyla cMOS camera (Andor), and MicroManager control 1055 software. For each image, SPD-5 condensates were identified through applying a 1056 threshold then using the particle analyzer function in FIJI. When analyzing condensate 1057 formation, we report the sum of the integrated intensities of each condensate per image 1058 (total condensate mass). Survival % plotted in Figure 5 assumes a 10-fold loss in total 1059 condensate mass due to dilution.

1060

## 1061 **Quantification and statistical analysis**

Images were analyzed with FIJI (<u>https://fiji.sc/</u>), R (<u>https://www.r-project.org/</u>), and GraphPad Prism (<u>https://www.graphpad.com</u>). For FLUCS experiments, centrosome deformation was calculated by measuring the long axis (orthogonal to the flow direction) of PCM-localized SPD-5 at the initial and final time points of PCM deformation, prior to 1066 fracture (defined below). The deformation rate equaled the difference in PCM lengths 1067 divided by the time interval. Centrosome fracture was measured using line scans across the long axis of PCM-localized SPD-5. Fracture was scored if signal dropped to 1068 1069 cytoplasmic levels over three consecutive pixels on the long axis across the entire flow 1070 path, and if this signal gap persisted for the rest of the images. For all other experiments, centrosome tracking and measurement was conducted using max intensity projections, 1071 1072 correction for photobleaching, followed by thresholding and particle analysis. Thresholds 1073 were determined using: mean background intensity of the cytoplasm + b\*(standard 1074 deviation of background), where b represents an integer value that is identical for all 1075 samples within an experiment. The integrated fluorescence density for the centrosome-1076 localized signals were normalized to either the first intensity value or max intensity value 1077 (Figure 3) and plotted over time.

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All data are expressed as the mean ± 95% confidence intervals as stated in the figure legends and results. The value of n and what n represents (e.g., number of images, condensates or experimental replicates) is stated in figure legends and results. Normality tests were first performed before applying statistical tests. Statistical tests were performed with GraphPad Prism.

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## 1085 **REFERENCES**

- Begasse, M.L., M. Leaver, F. Vazquez, S.W. Grill, and A.A. Hyman. 2015. Temperature
   Dependence of Cell Division Timing Accounts for a Shift in the Thermal Limits of C.
   elegans and C. briggsae. *Cell Rep.* 10:647-653.
- 1089Cabral, G., T. Laos, J. Dumont, and A. Dammermann. 2019. Differential Requirements for1090Centrioles in Mitotic Centrosome Growth and Maintenance. *Dev Cell*. 50:355-366 e356.
- Carvalho, A., S.K. Olson, E. Gutierrez, K. Zhang, L.B. Noble, E. Zanin, A. Desai, A. Groisman, and
  K. Oegema. 2011. Acute drug treatment in the early C. elegans embryo. *PLoS One*.
  6:e24656.
- Colombo, K., S.W. Grill, R.J. Kimple, F.S. Willard, D.P. Siderovski, and P. Gonczy. 2003.
   Translation of polarity cues into asymmetric spindle positioning in Caenorhabditis
   elegans embryos. *Science*. 300:1957-1961.
- Conduit, P.T., K. Brunk, J. Dobbelaere, C.I. Dix, E.P. Lucas, and J.W. Raff. 2010. Centrioles
   regulate centrosome size by controlling the rate of Cnn incorporation into the PCM.
   *Current biology : CB*. 20:2178-2186.

- Conduit, P.T., Z. Feng, J.H. Richens, J. Baumbach, A. Wainman, S.D. Bakshi, J. Dobbelaere, S.
   Johnson, S.M. Lea, and J.W. Raff. 2014a. The centrosome-specific phosphorylation of
   Cnn by Polo/Plk1 drives Cnn scaffold assembly and centrosome maturation. *Dev Cell*.
   28:659-669.
- Conduit, P.T., J.H. Richens, A. Wainman, J. Holder, C.C. Vicente, M.B. Pratt, C.I. Dix, Z.A. Novak,
  I.M. Dobbie, L. Schermelleh, and J.W. Raff. 2014b. A molecular mechanism of mitotic
  centrosome assembly in Drosophila. *eLife*. 3:e03399.
- Conduit, P.T., A. Wainman, and J.W. Raff. 2015. Centrosome function and assembly in animal
   cells. *Nature reviews. Molecular cell biology*. 16:611-624.
- Decker, M., S. Jaensch, A. Pozniakovsky, A. Zinke, K.F. O'Connell, W. Zachariae, E. Myers, and
   A.A. Hyman. 2011. Limiting amounts of centrosome material set centrosome size in C.
   elegans embryos. *Current biology : CB*. 21:1259-1267.
- Doxsey, S.J., P. Stein, L. Evans, P.D. Calarco, and M. Kirschner. 1994. Pericentrin, a highly
   conserved centrosome protein involved in microtubule organization. *Cell*. 76:639-650.
- Enos, S.J., M. Dressler, B.F. Gomes, A.A. Hyman, and J.B. Woodruff. 2018. Phosphatase PP2A
   and microtubule-mediated pulling forces disassemble centrosomes during mitotic exit.
   *Biol Open*. 7.
- Erpf, A.C., L. Stenzel, N. Memar, M. Antoniolli, M. Osepashvili, R. Schnabel, B. Conradt, and T.
   Mikeladze-Dvali. 2019. PCMD-1 Organizes Centrosome Matrix Assembly in C. elegans.
   *Current biology : CB*. 29:1324-1336 e1326.
- Fong, K.W., Y.K. Choi, J.B. Rattner, and R.Z. Qi. 2008. CDK5RAP2 is a pericentriolar protein that
   functions in centrosomal attachment of the gamma-tubulin ring complex. *Mol Biol Cell*.
   19:115-125.
- Frokjaer-Jensen, C., M.W. Davis, C.E. Hopkins, B.J. Newman, J.M. Thummel, S.P. Olesen, M.
  Grunnet, and E.M. Jorgensen. 2008. Single-copy insertion of transgenes in
  Caenorhabditis elegans. *Nat Genet*. 40:1375-1383.
- 1126 George, E.P., D. Raabe, and R.O. Ritchie. 2019. High-entropy alloys. *Nature Reviews Materials*.
- Gomez-Ferreria, M.A., U. Rath, D.W. Buster, S.K. Chanda, J.S. Caldwell, D.R. Rines, and D.J.
   Sharp. 2007. Human Cep192 is required for mitotic centrosome and spindle assembly.
   *Current biology : CB*. 17:1960-1966.
- Gonczy, P., S. Grill, E.H. Stelzer, M. Kirkham, and A.A. Hyman. 2001. Spindle positioning during
   the asymmetric first cell division of Caenorhabditis elegans embryos. *Novartis Found Symp.* 237:164-175; discussion 176-181.
- Grill, S.W., P. Gonczy, E.H. Stelzer, and A.A. Hyman. 2001. Polarity controls forces governing
   asymmetric spindle positioning in the Caenorhabditis elegans embryo. *Nature*. 409:630 633.
- 1136Grill, S.W., J. Howard, E. Schaffer, E.H. Stelzer, and A.A. Hyman. 2003. The distribution of active1137force generators controls mitotic spindle position. Science. 301:518-521.
- Hamill, D.R., A.F. Severson, J.C. Carter, and B. Bowerman. 2002. Centrosome maturation and
   mitotic spindle assembly in C. elegans require SPD-5, a protein with multiple coiled-coil
   domains. *Dev Cell*. 3:673-684.
- 1141Hannak, E., M. Kirkham, A.A. Hyman, and K. Oegema. 2001. Aurora-A kinase is required for1142centrosome maturation in Caenorhabditis elegans. J Cell Biol. 155:1109-1116.

1143 Harashima, H., N. Dissmeyer, and A. Schnittger. 2013. Cell cycle control across the eukaryotic 1144 kingdom. Trends Cell Biol. 23:345-356. 1145 Haren, L., T. Stearns, and J. Luders. 2009. Plk1-dependent recruitment of gamma-tubulin 1146 complexes to mitotic centrosomes involves multiple PCM components. PLoS One. 1147 4:e5976. 1148 Joukov, V., J.C. Walter, and A. De Nicolo. 2014. The Cep192-organized aurora A-Plk1 cascade is 1149 essential for centrosome cycle and bipolar spindle assembly. Mol Cell. 55:578-591. 1150 Kemp, C.A., K.R. Kopish, P. Zipperlen, J. Ahringer, and K.F. O'Connell. 2004. Centrosome 1151 maturation and duplication in C. elegans require the coiled-coil protein SPD-2. Dev Cell. 1152 6:511-523. 1153 Labbe, J.C., E.K. McCarthy, and B. Goldstein. 2004. The forces that position a mitotic spindle 1154 asymmetrically are tethered until after the time of spindle assembly. J Cell Biol. 167:245-1155 256. 1156 Laos, T., G. Cabral, and A. Dammermann. 2015. Isotropic incorporation of SPD-5 underlies 1157 centrosome assembly in C. elegans. *Current biology : CB*. 25:R648-649. 1158 Lee, K., and K. Rhee. 2011. PLK1 phosphorylation of pericentrin initiates centrosome maturation 1159 at the onset of mitosis. J Cell Biol. 195:1093-1101. 1160 Lemaitre, R.P., A. Bogdanova, B. Borgonovo, J.B. Woodruff, and D.N. Drechsel. 2019. FlexiBAC: a 1161 versatile, open-source baculovirus vector system for protein expression, secretion, and 1162 proteolytic processing. BMC Biotechnol. 19:20. Lindon, C., and J. Pines. 2004. Ordered proteolysis in anaphase inactivates Plk1 to contribute to 1163 1164 proper mitotic exit in human cells. J Cell Biol. 164:233-241. 1165 Magescas, J., J.C. Zonka, and J.L. Feldman. 2019. A two-step mechanism for the inactivation of 1166 microtubule organizing center function at the centrosome. eLife. 8. 1167 McNally, F.J. 2013. Mechanisms of spindle positioning. J Cell Biol. 200:131-140. 1168 Megraw, T.L., S. Kilaru, F.R. Turner, and T.C. Kaufman. 2002. The centrosome is a dynamic 1169 structure that ejects PCM flares. J Cell Sci. 115:4707-4718. Mittasch, M., P. Gross, M. Nestler, A.W. Fritsch, C. Iserman, M. Kar, M. Munder, A. Voigt, S. 1170 1171 Alberti, S.W. Grill, and M. Kreysing. 2018. Non-invasive perturbations of intracellular 1172 flow reveal physical principles of cell organization. Nat Cell Biol. 20:344-351. 1173 Nguyen-Ngoc, T., K. Afshar, and P. Gonczy. 2007. Coupling of cortical dynein and G alpha 1174 proteins mediates spindle positioning in Caenorhabditis elegans. Nat Cell Biol. 9:1294-1175 1302. 1176 Ozlu, N., M. Sravko, K. Kinoshita, B. Habermann, T. O'Toole E, T. Muller-Reichert, N. Schmalz, A. 1177 Desai, and A.A. Hyman. 2005. An essential function of the C. elegans ortholog of TPX2 is 1178 to localize activated aurora A kinase to mitotic spindles. Dev Cell. 9:237-248. 1179 Paix, A., A. Folkmann, D. Rasoloson, and G. Seydoux. 2015. High Efficiency, Homology-Directed 1180 Genome Editing in Caenorhabditis elegans Using CRISPR-Cas9 Ribonucleoprotein 1181 Complexes. Genetics. 201:47-54. 1182 Paix, A., A. Folkmann, and G. Seydoux. 2017. Precision genome editing using CRISPR-Cas9 and 1183 linear repair templates in C. elegans. *Methods*. 121-122:86-93. 1184 Panbianco, C., D. Weinkove, E. Zanin, D. Jones, N. Divecha, M. Gotta, and J. Ahringer. 2008. A 1185 casein kinase 1 and PAR proteins regulate asymmetry of a PIP(2) synthesis enzyme for 1186 asymmetric spindle positioning. Dev Cell. 15:198-208.

1187 Pecreaux, J., J.C. Roper, K. Kruse, F. Julicher, A.A. Hyman, S.W. Grill, and J. Howard. 2006. 1188 Spindle oscillations during asymmetric cell division require a threshold number of active 1189 cortical force generators. *Current biology : CB*. 16:2111-2122. 1190 Pelletier, L., N. Ozlu, E. Hannak, C. Cowan, B. Habermann, M. Ruer, T. Muller-Reichert, and A.A. 1191 Hyman. 2004. The Caenorhabditis elegans centrosomal protein SPD-2 is required for 1192 both pericentriolar material recruitment and centriole duplication. Current biology : CB. 1193 14:863-873. 1194 Pimenta-Margues, A., I. Bento, C.A. Lopes, P. Duarte, S.C. Jana, and M. Bettencourt-Dias. 2016. 1195 A mechanism for the elimination of the female gamete centrosome in Drosophila 1196 melanogaster. Science. 353:aaf4866. 1197 Rual, J.F., J. Ceron, J. Koreth, T. Hao, A.S. Nicot, T. Hirozane-Kishikawa, J. Vandenhaute, S.H. 1198 Orkin, D.E. Hill, S. van den Heuvel, and M. Vidal. 2004. Toward improving Caenorhabditis 1199 elegans phenome mapping with an ORFeome-based RNAi library. Genome Res. 14:2162-1200 2168. 1201 Schlaitz, A.L., M. Srayko, A. Dammermann, S. Quintin, N. Wielsch, I. MacLeod, Q. de Robillard, A. 1202 Zinke, J.R. Yates, 3rd, T. Muller-Reichert, A. Shevchenko, K. Oegema, and A.A. Hyman. 1203 2007. The C. elegans RSA complex localizes protein phosphatase 2A to centrosomes and 1204 regulates mitotic spindle assembly. Cell. 128:115-127. 1205 Severson, A.F., and B. Bowerman. 2003. Myosin and the PAR proteins polarize microfilament-1206 dependent forces that shape and position mitotic spindles in Caenorhabditis elegans. J 1207 Cell Biol. 161:21-26. 1208 Sun, T.L., T. Kurokawa, S. Kuroda, A.B. Ihsan, T. Akasaki, K. Sato, M.A. Hague, T. Nakajima, and 1209 J.P. Gong. 2013. Physical hydrogels composed of polyampholytes demonstrate high 1210 toughness and viscoelasticity. Nat Mater. 12:932-937. 1211 Sunkel, C.E., and D.M. Glover. 1988. polo, a mitotic mutant of Drosophila displaying abnormal 1212 spindle poles. J Cell Sci. 89 (Pt 1):25-38. 1213 Weinert, F.M., J.A. Kraus, T. Franosch, and D. Braun. 2008. Microscale fluid flow induced by 1214 thermoviscous expansion along a traveling wave. Phys Rev Lett. 100:164501. 1215 Wlodarchak, N., and Y. Xing. 2016. PP2A as a master regulator of the cell cycle. Crit Rev Biochem 1216 *Mol Biol*. 51:162-184. 1217 Woodruff, J.B., B. Ferreira Gomes, P.O. Widlund, J. Mahamid, A. Honigmann, and A.A. Hyman. 1218 2017. The Centrosome Is a Selective Condensate that Nucleates Microtubules by 1219 Concentrating Tubulin. Cell. 169:1066-1077 e1010. 1220 Woodruff, J.B., and A.A. Hyman. 2015. Method: In vitro analysis of pericentriolar material 1221 assembly. Methods Cell Biol. 129:369-382. 1222 Woodruff, J.B., O. Wueseke, and A.A. Hyman. 2014. Pericentriolar material structure and 1223 dynamics. Philos Trans R Soc Lond B Biol Sci. 369. 1224 Woodruff, J.B., O. Wueseke, V. Viscardi, J. Mahamid, S.D. Ochoa, J. Bunkenborg, P.O. Widlund, 1225 A. Pozniakovsky, E. Zanin, S. Bahmanyar, A. Zinke, S.H. Hong, M. Decker, W. Baumeister, 1226 J.S. Andersen, K. Oegema, and A.A. Hyman. 2015. Centrosomes. Regulated assembly of a 1227 supramolecular centrosome scaffold in vitro. Science. 348:808-812. 1228 Zebrowski, D.C., S. Vergarajauregui, C.C. Wu, T. Piatkowski, R. Becker, M. Leone, S. Hirth, F. 1229 Ricciardi, N. Falk, A. Giessl, S. Just, T. Braun, G. Weidinger, and F.B. Engel. 2015.

- 1230 Developmental alterations in centrosome integrity contribute to the post-mitotic state
- 1231 of mammalian cardiomyocytes. *eLife*. 4.
- Zhu, F., S. Lawo, A. Bird, D. Pinchev, A. Ralph, C. Richter, T. Muller-Reichert, R. Kittler, A.A.
   Hyman, and L. Pelletier. 2008. The mammalian SPD-2 ortholog Cep192 regulates
- 1234 centrosome biogenesis. *Current biology : CB*. 18:136-141.
- 1235











## Mittasch et al. Figure 5



10 µm







#### Continuous FLUCS during metaphase-anaphase transition D

Metaphase-Anaphase mMaple::SPD-5 mCherry::H2B OFF 169 **10 μm** 

#### 10 kHz, bidirectional laser scan---no flow, only heat Ε



#### Embryo survival after FLUCS in one-cell stage F

### **Brightfield images**





csnk-1RNAi) --- increased pulling forces throughout the cell cycle



# Mittasch et al. Figure S3



# **Disassembly profiles for posterior PCM proteins**

