1	Common Themes in Centriole and Centrosome Movements	
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#### 11 Abstract

12 Centrioles are found in nearly all eukaryotic cells and are required for growth and 13 maintenance of the radial array of microtubules, the mitotic spindle, and cilia and flagella. 14 Different types of microtubule structures are often required at different places in a given cell; 15 centrioles must move around to nucleate these varied structures. Here we draw together 16 recent data on diverse centriole movements to decipher common themes in how centrioles move. Par proteins establish and maintain the required cellular asymmetry. The actin 17 18 cytoskeleton facilitates movement of multiple basal bodies. Microtubule forces acting on the 19 cell cortex, and nuclear-cytoskeletal links, are important for positioning individual 20 centrosomes, and during cell division. Knowledge of these common mechanisms can inform 21 the study of centriole movements across biology.

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### 23 Introduction

In metazoan cells the major microtubule organising centre (MTOC) of the cell is the 24 25 centrosome (Figure 1 and glossary). It is composed of a pair of microtubule-based centrioles 26 surrounded by a pericentriolar matrix (PCM). Centrioles were once thought to be static organelles located in the centre of the cell, hence their name. In fact, they move around the 27 28 cell to fulfil their functions and correct centrille and centrosome positioning is vital for many 29 biological processes. There are now many examples of centriole/centrosome movements in 30 various physiological contexts and many different cell types. We will use the term 31 centrosome in situations where the centricle pair and PCM all move together, and centricle 32 where only a single centriole moves. Basal body is the term often used for a centriole that 33 assembles a cilium or flagellum. While many advances have been made over the last few 34 years in understanding what controls centrosome, centrille and basal body position, often, 35 these fields are investigated in isolation. If the data from these disparate fields are analysed together, it becomes apparent that there are common themes in both the mechanics of movement and the regulatory mechanisms involved. Here, we first provide a brief overview of the contexts in which centriole, centrosome and basal body movements are seen, and then elaborate on how some of the common themes that are emerging across eukaryotic biology are applied in each case.

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Centrosome location is critical for many biological processes (Figure 2-4) and also impacts 42 43 the position of other organelles. The centrosome and the nucleus are closely associated, and 44 the Golgi apparatus is also found near the centrosome [1] enabling polarization of membrane 45 trafficking and secretory machineries [2-3]. While centrosomes are not absolutely required to 46 organise the mitotic spindle [4-6], their position is important in symmetric and asymmetric 47 cell divisions (Figure 3) as movement of the two centrosomes to opposite sides of the nucleus 48 defines both the axis of division, and spindle position. In *Caenorhabditis elegans*, centrosome 49 positioning is key to the polarity establishment required during asymmetric cell divisions and 50 defines the anterior-posterior axis of the embryo [7]. The African trypanosome does not use 51 its centrioles (located at the base of the flagellum and termed basal bodies) to organise the 52 mitotic spindle; however, basal body positioning and segregation control cell morphogenesis 53 by influencing cytoskeletal construction and directly positioning the kinetoplast 54 (mitochondrial genome) [8] (Figure 3).

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56 Centriole/centrosome position also contributes to the spatial organisation of many cells in 57 G1/interphase. biflagellate In the green alga Chlamydomonas reinhardtii 58 centriole/centrosome position maintains overall cell geometry [9], and in metazoans it 59 organises the radial microtubule array during interphase (Figure 2). In many types of migrating cells, centrosome position between the nucleus and the leading edge is key to 60

migration [10-13]. Formation of the specialised immunological and virological synapses 61 62 involves centrosome re-orientation. During immunological synapse formation, the 63 centrosome migrates to the contact site between the T-cell and the antigen presenting cell 64 (Figure 4), where, in cytotoxic T-cells, it is involved in directed secretion of lytic granules [14]. During virological synapse formation, which mediates the cell-cell transfer of viral 65 66 particles between an infected cell and a target cell, the centrosome of the infected cell likewise re-orients towards the site of contact between the cells [15]. Finally, while centrioles 67 68 are not absolutely essential for cell division [16], they are critical for ciliogenesis. Humans 69 build motile and sensory cilia, and ciliogenesis of both kinds requires centriole/basal body 70 movement to the cell surface [17] (Figures 2 and 4). Most branches of the tree of life are 71 ciliate, and in many cases failure to build a cilium is incompatible with life. Even in 72 Drosophila, which can develop without centrioles, death eventually occurs due to the lack of 73 cilia on sensory neurons [16].

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Thus, there is a considerable diversity of centriole/centrosome movements in biology. All of the studied mechanisms of centriole movement require the actin or microtubule cytoskeletons, or both. Are there common mechanisms that apply across these varied processes or are there specialised mechanisms to facilitate movement in each physiological context?

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## 81 The microtubule cytoskeleton and centriole movement

The position of the centrosome in both migrating and non-migrating interphase cells requires a polarised radial microtubule array [18-19]. During cell migration, centrosome position is actin-independent [10, 12], while the cortical pool of the microtubule minus-end directed motor cytoplasmic dynein is implicated in centrosome position in several different cell types 86 [10, 12-13, 20]. During astrocyte migration, the small GTPase Cdc42 controls centrosome 87 and Golgi re-orientation towards the direction of migration through the microtubule 88 cytoskeleton and cytoplasmic dynein [10, 21]; however, in fibroblasts, the centrosome 89 remains at the cell centre while the nucleus moves rearward [20]. Nonetheless, centrosome 90 maintenance at this position is dependent on microtubules and dynein as inhibition of either 91 causes a rearward centrosome displacement [20].

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93 Microtubules and cytoplasmic dynein are also implicated in centrosome positioning in non-94 migrating cells [19, 22-24], suggesting that, when the centrosome needs to be central within 95 the cell, its position is actively maintained using microtubules and dynein to stabilise the 96 centrosome-associated microtubule array. In interphase cells, the centriole pair do not 97 necessarily remain together. The older, mature, centriole can remain stationary while the 98 vounger, immature centriole moves around the cell [25]. The Rho-associated kinase 99 p160Rock, is proposed to regulate the central position of the mature centriole [26]. p160Rock 100 is a major regulator of myosin II but it has many different substrates [27] and during cell 101 migration it can mediate both microtubule-dependent centrosome re-orientation [28] and 102 actin rearrangements; consequently, the way in which it affects centrosome position remains 103 unclear.

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In summary, microtubules and cytoplasmic dynein regulate centrosome position when a single centrosome needs to be positioned in the centre of the cell. How about when centrioles need to move away from the centre, or if there are multiple centrioles to move? In these cases, several studies have highlighted a primary role for the actin cytoskeleton in centriole positioning.

## 111 Actin in centrosome/centriole movements

While the first cell division of many organisms is symmetric, that of C. elegans is 112 asymmetric. During the first division of the C. elegans zygote, the cell divides 113 114 asymmetrically along its anterior-posterior axis to give rise to two cells that are committed to different cell fates. The centriole pair are key factors in the polarity establishment required 115 116 during these divisions. They are derived from the sperm cell rather than the oocyte and become embedded in the actin cortex underlying the plasma membrane by a mechanism that 117 118 may not involve interactions with microtubules. RNAi ablation of tubulin does not prevent 119 polarity establishment and is not required for centriole-cortex interaction [7, 29], suggesting 120 that another cytoskeletal polymer mediates centrille position. However, another study found 121 that tubulin disruption delays polarity induction, which requires a small centrosomal 122 microtubule aster [30]; therefore the precise role of microtubules in this process remains to be 123 clarified.

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125 The trachea, oviduct and ependymal epithelium of mammals are composed of a multi-ciliated 126 epithelium where each cell may have hundreds of cilia, each grown from a basal body. These 127 form de novo and migrate simultaneously to the apical cell surface. Drugs that target 128 microtubules do not directly stop basal body movement in oviduct [31], although the 129 contribution of microtubules to basal body movements in other cell types is unclear. In 130 contrast, much evidence from several cell types implicates the actin cytoskeleton in basal 131 body migration and docking at the cell surface. Actin and myosin associate with either basal 132 bodies, or the material surrounding them [32-34] and basal body migration is blocked by 133 treatment with inhibitors of actin [35] or myosin [36]. Basal body docking with the cell 134 membrane is also actin dependent. The Wnt planar cell polarity pathway and its effectors are 135 implicated in membrane trafficking during ciliogenesis and the formation of an actin array essential for basal body docking [37-39] [40]. Finally, myosin II localises to basal body
accessory structures in multi-ciliated epithelia [32], and is needed for basal body migration
[41].

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There are emerging similarities in the structure and function of basal bodies of cilia and 140 141 centrioles at the immunological synapse [42], such as the requirement for intraflagellar 142 transport components. These were previously thought of as cilium assembly and maintenance 143 proteins [43], but recent data have also demonstrated a role in polarised recycling at the 144 immunological synapse [44]. This suggests that there may be similar principles guiding 145 centriole movement and membrane trafficking in both cases. During immunological synapse 146 formation, receptor-mediated engagement between the immune cell and its target triggers a 147 transient aggregation of actin across the nascent synapse [45]. The polarisation of the 148 centriole pair to the synapse is accompanied by concomitant actin clearance from the inner 149 part of the forming synapse, to produce an outer ring of actin [14, 46] and it has been 150 suggested that the forces generated by actin clearance are used to move the centrioles forward 151 [2].Centrioles are always docked at the centre of the synapse and it is unclear whether this is 152 due to radial actin reorganisation that localises them to this region by default, or whether 153 there are other actin binding proteins that regulate the site of docking. Much less is known 154 about the role of the cytoskeleton in virological synapse formation, although it is an actin-155 dependent process [47] and integrity of both the actin and microtubule cytoskeletons is 156 crucial [48-49].

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158 In general it is over-simplistic to consider the actin and microtubule cytoskeletons in 159 isolation. Extensive cross-talk exists and many biological processes are carried out by actin

and microtubules acting together. Much research has shown that interactions of microtubulesthat are anchored into the actin cortex are often responsible for centrile positioning.

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## 163 Microtubule interactions with the actin cortex and centrille and centrosome positioning

164 Interaction between microtubules and the actin cytoskeleton at the cell cortex are essential 165 for maintaining the physical position of the centrosome within the cell and for orchestrating placement of the duplicated centrosomes during symmetric and asymmetric cell divisions. 166 167 Pulling forces of microtubules anchored to the cell cortex provide a mechanism for 168 centrosome positioning [11, 50]. Studies of male germline stem cells in Drosophila have 169 revealed differential centrosome behaviour during the asymmetric cell divisions that 170 characterise development. After centrosome duplication, the older of the two centrosomes 171 retains a well-defined microtubule array and remains in place, while the younger centrosome 172 nucleates few microtubules and migrates away to set up the symmetrical plane of the mitotic 173 spindle [51].

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175 The pulling forces are provided by dynein on microtubules that extend to and are anchored 176 into the cell cortex regulated by both dynein and the actin motor myosin II. Myosin II is one 177 of several proteins that organise the cortex, and cortical organisation is critical for providing the mechanical support needed for centrosome positioning. The importance of cortical 178 179 rigidity has been highlighted by recent studies that showed that myosin II- and moesin-180 dependent cortical rigidity are required for spindle positioning [52-54]. Moreover, cell shape 181 plays an important part in spindle orientation, which is highly dependent on cell-substrate 182 adhesions [55] that are communicated to the cytoskeleton via integrins and actin-microtubule 183 linkers including EB1 and myosin X [56-57].

185 Taken together, these studies provide evidence for a conserved pathway that explains 186 centrosome movement in cells containing a centralised centrosome with a radial array of 187 microtubules that are in contact with the plasma membrane. However, not all eukaryotic cells 188 have a radial array of microtubules emanating from a centralised MTOC. Protists such as the 189 African trypanosome represent the polar opposite, with microtubules arranged as a 190 subpellicular sheet underlying the plasma membrane [58]. Basal bodies extend a microtubule 191 axoneme for the flagellum in this organism, but not a radial array of microtubules within the 192 cell body. Despite these differences, microtubules do provide the mechanism for basal body 193 movements and segregation during cell division [59] and these microtubules are closely 194 associated with the plasma membrane in the same way that microtubules associate with the 195 actin cortex underlying the plasma membrane in other organisms.

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197 Thus, the idea that microtubule interactions with a cortical cytoskeleton are used to move 198 single or paired centrioles/centrosomes is conserved across eukaryotic biology.

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## 200 The role of the nuclear envelope in centrille movements.

201 It is increasingly recognised that the nucleus, as well as the cortex, plays a role in positioning 202 centrioles. In many cells, the centrioles are tightly associated with the nuclear envelope. This 203 connection is observed in lower eukarvotes as a physical linkage of striated fibres called the basal body-nucleus connector (rhizoplast, [60]). During the cell division cycle of 204 205 *Chlamydomonas*, the two flagella are cleaved from the cell. The two pairs of centrioles move 206 from the apical surface to the poles of the spindle during mitosis, and after division return to 207 the apical surface and grow two new flagella. The basal body-nucleus connector exists as 208 striated fibres that connect both pairs of centrioles, which are in turn connected to the nucleus 209 by centrin-containing rhizoplasts [61]. Is the nucleus, the centrille or both involved in the 210 movement to and from the apical surface? In cells lacking the rhizoplast connection, the 211 nucleus is mispositioned, but the mature centriole is correctly localised, indicating that 212 nuclear mis-positioning has little impact on mature centrille positioning [9] and suggesting that the centriole regulates nuclear positioning rather than vice versa. In a migrating 213 214 mammalian cell the opposite appears true. Studies on centrosome reorientation in migrating 215 fibroblasts suggest that the centrosome might remain relatively central while the nucleus 216 moves rearwards [20]. Centrosome position is maintained by dynein-mediated cortical 217 tethering of microtubules [62]; however, it is unclear whether centrosome rotation drives 218 nuclear movement or vice versa.

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In higher eukaryotes, the link between the nuclear envelope and the centrosome is essential for development [63] and is robust enough to withstand cell lysis and nuclear isolation [64]. This linkage is required for nuclear migration and the control of cell cycle timing [64] and in *Drosophila* there is evidence that the centrosomes can reach the cell cortex during development with the aid of the nucleus [65].

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226 In the organisms studied to date, the centrosome-nucleus linkage is mediated by proteins 227 containing paired KASH (Klarsicht-anchorage protein1-Syne homology) and SUN (Sad1-228 UNC84) domains (Figure 5). First discovered in C. elegans, proteins with these domains are 229 found across eukaryotes and localise to the nuclear envelope and centrosomes [63, 66-68]. 230 Multiple SUN and KASH proteins exist that provide links between the nuclear envelope and 231 cytoskeletal polymers [69-70] (Figure 5). One of the diverse roles of the SUN-KASH 232 complex is the regulation of centrosome position. The C. elegans KASH protein ZYG-12 233 anchors the centrosome to the nuclear envelope during embryogenesis [63]. ZYG-12 is not 234 found in mammalian cells; however several proteins fulfill the role of nucleus-centrosome 235 linkers including the nuclear membrane protein emerin [71], and the multi-isoform KASH 236 protein Nesprin 2 [66]. Several Nesprin isoforms contain calponin-homology domains that 237 allow them to bind actin, and these are implicated in positioning the centrosome during 238 ciliogenesis of sensory cilia [72], suggesting that actin-dependent nuclear re-positioning or 239 rotation may re-orient the centrosome apically. A novel epithelial-specific Nesprin isoform, 240 Nesprin 4, interacts with the microtubule motor kinesin-1, and this link is proposed to 241 contribute to nucleus and centrosome positioning in interphase cells [73]. It will be interesting to see if Nesprin 4 and kinesins are also involved in centrosome/centriole 242 243 positioning during ciliogenesis as Nesprin-microtubule links are also important in cell 244 migration. In migrating neurons, a SUN1/2-Nesprin1/2 complex acts with the lissencephaly-245 associated proteins Lis1 and Doublecortin to couple the centrosome and nucleus through 246 cytoplasmic dynein [66, 74]

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## 248 **Regulation of centriole movements**

Given that centrioles and centrosomes can track around the nuclear envelope, change their position relative to the nucleus, embed themselves in the cortex, or move around the cell, how do they know where to go? In the absence of other cues, cell-cell contacts are the main mediators of centrosome positioning [75-76]. The extracellular signals that trigger centrosome and centriole movements are varied; however there is now much evidence from a variety of systems that these signals converge on the Par (partitioning) proteins [77] and the Rho family of small GTPases.

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The Par proteins are a key set of polarity proteins that were identified in screens for mutants affecting the first asymmetric cell division in *C. elegans* [78-79]. In both *C. elegans* and *Drosophila*, the Par complex acts through the Rho family of small GTPases and the actin 260 cytoskeleton to establish the cortical polarity that is needed for spindle positioning [50, 80]. 261 Once this initial polarity is established, the same mechanisms act together with microtubule-262 cortical interactions to produce the forces that result in the asymmetrically placed spindle. 263 External signalling cues from neighbouring epithelial cells are needed to regulate the 264 localisation of polarity markers - and hence the axis of the mitotic spindle - during 265 asymmetric cell division in Drosophila neuroblasts [81]. Two Rho GTPases act together to 266 regulate polarity establishment in C. elegans. Rho1 mediates the centrosome-dependent 267 cortical actomyosin rearrangements that lead to contractile asymmetry within the cortex. 268 Cdc42 mediates the link between the cortex and Par6 proteins, and coordinates Par protein 269 segregation as the cortical asymmetry develops [80]. During cell migration, integrin 270 signalling through Cdc42 to Par6 and aPKC is required for the microtubule-dependent 271 centrosome localisation observed during astrocyte migration [10, 21] and blocking Cdc42 272 prevents macrophage polarization towards a chemotactic signal [82]. A Par3-Par6-aPKC 273 complex stabilises microtubule-dependent cell polarity during keratinocyte migration, 274 although its role in centrosome movement is unclear [83]. During development, aPKC is 275 needed during neuronal repolarization [84] and Pard3 controls centrosome positioning during 276 neurulation [85]. During ciliogenesis of multi-ciliated epithelia, the Par3-Par6-aPKC polarity 277 complex localises to cilia and regulates ciliogenesis via association with kinesin-II [86], one 278 of the motors required to build cilia by intraflagellar transport [43]. Rho is not needed for 279 centriole re-orientation during virological synapse formation, however, inhibition of Rac and 280 Cdc42 prevents centrioles from re-orientating [49]. Cdc42 and Par proteins are also 281 implicated in immunological synapse formation. Cdc42 inhibition blocks centrosome re-282 orientation [87], while Par3 is recruited to the synapse [88] and overexpression of a 283 dominant-negative form of Par1b blocks centrosome re-orientation [89], suggesting that Par3 284 localisation is functionally relevant to immunological synapse formation. The signalling events that regulate the Par proteins in this case are unclear, however strength of signalling via the T cell receptor is important [90] and when more than one contact is present, the centrosome can oscillate between the possible targets [91] until the decision is made to kill the target that produces the strongest signal [92]. It is therefore reasonable to suggest that this provides the required external cue.

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### 291 Concluding remarks

292 While the importance of the cytoskeleton, polarity proteins, and the nuclear envelope in 293 centriole movements has long been recognised in several different fields, the idea of common 294 themes has been slower to emerge. Research carried out over the last few years has 295 highlighted that, even though centrioles and centrosomes are positioned to achieve very 296 different outcomes, much of the basic machinery that is used is remarkably similar. It seems 297 likely that disparate signalling events might converge on the recruitment of the Par proteins to 298 establish and maintain the asymmetry that is a key feature of these centrosome re-orientation 299 events. In general, where multiple centrioles need to be moved, there is a requirement for the 300 actin cytoskeleton, while microtubule forces acting on the cell cortex are particularly 301 important for positioning individual centrosomes, and during cell division. Finally, the 302 involvement of KASH proteins in multiple centrosome positioning contexts suggests that 303 they too may represent a conserved mechanism for regulating centrosome location, and their 304 potential roles in mediating other centrosome movements warrants investigation. A challenge 305 for the future is to identify the polarity cues that regulate centrosome position in organisms 306 outside the metazoa that lack the Par proteins. It will be interesting to see if there are 307 conserved mechanisms to set up asymmetry in these systems. These might include examples 308 of cytotaxis such as those that are involved in polarity replication during trypanosome 309 morphogenesis [93] or the inheritance of cortical organisation in ciliates [94].

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311 Several proteins that are implicated in control of centrosome/centriole positioning have been 312 linked to human inherited disease. Lissencephaly, or "smooth brain," is a brain malformation disorder caused by abnormal neuronal migration early in development. Two of the underlying 313 314 proteins, Lis1 and Doublecortin, mediate the centrosome-nucleus linkage [74] and it seems 315 likely that polarity problems caused by disruption of this link might contribute to the disease. 316 Other neuronal migration disorders can also result in structurally abnormal or missing areas 317 of the brain including midline defects such as agenesis of the corpus callosum and hypoplasia 318 of the cerebellar vermis. Many the ciliopathies, or diseases of cilium dysfunction, present 319 with midline defects as part of the phenotype, and two ciliopathies, Meckel-Gruber syndrome 320 and hydrolethalus syndrome, have been linked to centrosome/basal body-positioning defects 321 [95-96]. How these fit in to the pathways and processes described here remains to be seen, 322 however, the Meckel-Gruber syndrome proteins are implicated in planar cell polarity 323 signaling [97-98] and may regulate centrosome re-orientation during ciliogenesis through 324 actin cytoskeleton remodeling and maintaining the centrosome-nuclear envelope connection 325 [72]. Finally, there are other diseases that have been linked to centrosome dysfunction [99] 326 and it will be fascinating to discover if centriole/centrosome position is also compromised in 327 these cases.

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As more details of the molecular control of polarity establishment are uncovered, it will become possible to understand which of the activities in *C. elegans* represent general principles in polarity establishment, and which are specialized to the particular case of embryonic polarity establishment. Despite the likelihood of cell-type specific specializations, analysis reveals a commonality in the mechanisms used to move centrioles and centrosomes throughout eukaryotic biology. Notwithstanding the very different contexts in which centriole

- 335 movements are observed, these commonalities have the potential to contribute to our
- 336 understanding of centriole movements in less well-studied systems.
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- 545 546

# 547 Figure Legends

548 Fig 1: Centriole duplication cycle. During interphase/G1 centrioles function in organizing 549 microtubules, and in many eukaryotic cells the mature centriole assembles a primary 550 cilium/flagellum. The centriole pair must duplicate only once during the cell cycle and this 551 begins at the G1/S-phase transition with a pro-centriole assembled orthogonal to each mature 552 centriole. Recent studies have dissected the molecules required for initial pro-centriole assembly in C. elegans early embryos and humans (pro-centriole assembly: C. elegans: SPD-553 2, ZYG-1 Sas-6, Sas-5, Sas-4, α-, β-, and γ-tubulin; Humans: Cep192, Plk4/Sak, Sas-6, Sas-554 555 4, CPAP,  $\alpha$ -,  $\beta$ -,  $\gamma$ -,  $\delta$ -,  $\epsilon$ -tubulin, CP110, and Cep135. Regulatory molecules: Cdk1/Cyclin B, Aurora-A, Plks. Control of centriole duplication: Separase, Plk1, Plk4, SAS-6.) [99]. Pro-556 557 centriole elongation continues through G2 until there are two pairs of centrioles that migrate 558 to the pole of the spindle. Following mitosis the tight association and orthogonal orientation 559 of the mature centriole and pro-centriole is no longer apparent. This is a stage termed 'disengagement' and is a crucial stage in the control of centrille duplication [100]. (a) 560 561 Microtubule arrays/primary cilium functions. (b) New centriole formation. (c) New centriole 562 elongation. (d) Centrosome segregation. (e) Centrosomes move to spindle poles. (f)563 Centrosome inheritance to daughter cells.

564

565 Fig 2: Centrosome migration in G1/interphase cells. In many animal cells the centrosome migrates from a central position within the cell to the cell cortex, where a primary cilium 566 567 assembles from the mature centriole. The primary cilium acts as an antenna for the cell that 568 senses the environment and is needed to transduce certain signaling pathways. Migration of 569 the centrosome can involve interaction with radial microtubules and actomyosin at the cell 570 cortex, however, the role of the cortex in primary cilium formation is not known. (a) 571 G1/interphase cell. The centrosome is located centrally within the cell. Central location is 572 maintained by microtubules/dynein and regulated by p160Rock. (b) The centrosome moves 573 to the cell surface in some cell types via interaction of microtubules with the actomyosin cell 574 cortex. (c) A microtubule-based primary cilium is assembled from the mature centriole.

575

576 Fig 3: Centrosome/basal body migration during cell division. (a). Symmetric & 577 asymmetric cell division. Migration of the duplicated centrosomes to the opposite poles of the 578 spindle requires actin-microtubule interactions with the cell cortex (left). The Par proteins are 579 important in modulating these interactions in order to promote asymmetric positioning 580 (right). (b). Interphase African trypanosome cell with a single flagellum assembled from the 581 mature basal body (left). G1/S-phase basal body duplication occurs and a new flagellum 582 assembles alongside the old flagellum. Migration is microtubule-dependent via subpellicular 583 microtubules at the cell cortex (middle). Intriguingly, actin and myosin II are not involved in 584 either basal body migration or cytokinesis in T. brucei (right). (c). Bi-flagellated interphase 585 C. reinhardtii cell (left). Flagella are cleaved and the centrioles migrate to the poles of the mitotic spindle via a nucleus-centriole connector (rhizoplast; middle). Centrioles return to 586

the cell cortex and two new flagella are assembled for each daughter cell prior to cytokinesis(right).

589

590 Fig. 4: Centriole/basal body migration in terminally differentiated cells. (a). The process 591 of ciliogenesis produces thousands of motile or immotile cilia on many specialized terminally 592 differentiated cells. Large numbers of basal bodies are formed within a single cell (left). 593 Basal bodies migrate and dock with the cell membrane. Movement requires actomyosin, and 594 is regulated by GTPase RhoA (middle). Motile or immotile cilia are assembled from the 595 docked basal bodies (right). Basal bodies form via a combination of the centriolar and 596 acentriolar pathways (see text box). The role of the existing centrosome is unknown (b). 597 Cytotoxic T-cells form an immunological synapse to facilitate killing a target cell. 598 Centrosome migration is required during the early stages of synapse formation and occurs by 599 interactions between microtubules and the cell cortex. Recognition of a target cell by a 600 cytotoxic T-cell and assembly of the synapse (left; arrow points to nascent synapse). 601 Movement of the centrosome to the synapse requires both microtubules and actomyosin 602 (middle). The centrosome docks at the plasma membrane of the immunological synapse and 603 lytic granules (black) travel along microtubules to the synapse to kill the target cell (right).

604

Fig 5: SUN and KASH domain proteins couple the nucleus to the actin and microtubule cytoskeletons. The SUN domain-KASH domain interaction occurs within the space between the inner and outer nuclear membranes. Many different KASH-domain proteins exist and can provide a physical linkage between the nuclear lamina and the cytoskeleton. KASH proteins with an N-terminal actin-binding domain link the actin cytoskeleton to the nucleus. Other KASH proteins link microtubules to the nucleus via interactions with kinesin or dynein. The 611 SUN-KASH interaction is evolutionarily conserved and ha many roles within cells,612 including nuclear migration and centrosome orientation.

613

614 Figure I

615 Transmission electron micrograph of a centriole pair from a mouse kidney cell. Scale bar:616 100nm.

617

618 **Glossary:** 

619 Actin motors: myosins are actin motors that carry cargo along actin and are ATP-dependent.

620 With the exception of myosin VI all other myosins studied to date are plus-end directed.

Astral microtubules: extend out from each centrosome at opposite poles of the mitoticspindle pole to the cell cortex and are required for mitotic spindle orientation.

623 **Basal body:** a microtubule organizing centre that subtends a cilium or flagellum.

624 **Cell cortex:** a specialized area of the cell underlying the plasma membrane that is required

625 for mechanical support of cell shape and form. Microtubules (called cortical microtubules),

actin (called cortical actin) or both are found at the cell cortex in a wide range of eukaryoticcells.

628 **Centriole:** a microtubule-based barrel-shaped structure generally composed of 9 triplet 629 microtubules that is found in many cells (Figure I).

630 Centrosome: the major microtubule organizing centre in mammalian cells. It organizes
 631 radial arrays of microtubules, mitotic spindle microtubules and astral microtubules, and
 632 contains a pair of centrioles.

633 Immunological/Virological synapse: named for their similarity to classical neurological
634 synapses, the immunological synapse is the interface between an antigen-presenting cell and

a lymphocyte, while the virological synapse is the interface between infected cells and targetcells that can mediate cell-cell spread of viruses.

Microtubule motors: kinesin motors move along microtubules towards the plus-end of
 microtubules and dynein motors move towards the minus-end of microtubules. Both are
 ATP-dependent motors.

640 Pericentriolar material: the matrix that surrounds the centrioles within the centrosome. It 641 contains proteins responsible for microtubule nucleation and anchoring and plays a role in 642 centrosome duplication.

643

## 644 Box 1. Basal body production during ciliogenesis

Many metazoan organisms build two types of cilium: non-motile sensory, or primary, cilia and motile cilia. Each assembles from a basal body, which is analogous to the mitotic centrioles.

648

• Primary cilia are solitary organelles that assemble from a basal body derived from the
 pre-existing mature centriole, which moves to the cell surface and docks before extending
 the ciliary axoneme.

652

In contrast, there can be hundreds of motile cilia on a single cell and each needs a basal body. Basal body formation is linked to differentiation rather than proliferation and multiple basal bodies are formed in the cytoplasm and then simultaneously migrate to the cell surface. Basal bodies are formed *de novo* by a combination of the centriolar pathway and the acentriolar pathway, both of which can occur in a single cell. In the centriolar pathway, new basal bodies are produced around an existing centriole
 template, just as observed during cell cycle-dependent centriole duplication. However,
 more than one new basal body can form around a single centriole.

In the acentriolar pathway, basal body formation is not templated. Here, multiple basal
 bodies form around an intermediary structure called a deuterosome rather than around an
 existing centriole.



Figure 1









