Recent advances in pericentriolar material organization: ordered layers and scaffolding gels [version 1; referees: 3 approved]

Andrew M. Fry, Josephina Sampson, Caroline Shak, Sue Shackleton
Department of Molecular and Cell Biology, University of Leicester, Leicester, UK

Abstract
The centrosome is an unusual organelle that lacks a surrounding membrane, raising the question of what limits its size and shape. Moreover, while electron microscopy (EM) has provided a detailed view of centriole architecture, there has been limited understanding of how the second major component of centrosomes, the pericentriolar material (PCM), is organized. Here, we summarize exciting recent findings from super-resolution fluorescence imaging, structural biology, and biochemical reconstitution that together reveal the presence of ordered layers and complex gel-like scaffolds in the PCM. Moreover, we discuss how this is leading to a better understanding of the process of microtubule nucleation, how alterations in PCM size are regulated in cycling and differentiated cells, and why mutations in PCM components lead to specific human pathologies.
Introduction
The centrosome is a single copy organelle present in the majority of animal cells. Through concentrating proteins required for microtubule nucleation, most notably γ-tubulin and its γ-tubulin ring complex (γ-TuRC) partners, it serves as the primary microtubule organizing center (MTOC) of the cell. Centrosomes have two major structural elements: a centriole pair, consisting of two approximately 200 by 400 nm barrels that are each composed of nine highly stable microtubule triplets, and a surrounding protein-rich matrix, the PCM, that is attached to the centrioles but extends outward to a diameter of about 1 micron. Unlike most organelles, the centrosome lacks an encompassing membrane, and although EM has shown the elegant ultrastructure of the centrioles, it has told us very little about how the PCM is organized or what defines its size and shape. Indeed, with the large coiled-coil and electron-dense nature of most PCM components, our view of the PCM has remained frustratingly opaque. Yet we have long known that the PCM is the site from which microtubules are nucleated and that microtubule nucleation capacity can be precisely modulated according to specific cues. Hence, recent technological breakthroughs in subdiffraction super-resolution imaging, together with structural biology and biochemical reconstitution approaches, now provide us with a much more detailed view of PCM architecture that is both exciting and transformative to our understanding.

The PCM proximal layer is highly organized in the interphase centrosome
Prior to super-resolution imaging, the most accurate and useful conception of the PCM was as a salt-resistant “centro-matrix” of 12–15 nm wide filaments in which circular γ-TuRCs of 25–30 nm diameter were embedded. However, the development of different modalities of “optical nanoscopy” that allowed imaging below the standard diffraction limit of fluorescent light, including structured illumination microscopy (SIM), stimulated emission depletion (STED), stochastic optical-reconstruction microscopy (STORM), and photoactivated localization microscopy (PALM), led to a completely new understanding of how the PCM is organized. Indeed, these super-resolution immunofluorescence microscopy approaches, mainly undertaken in Drosophila (fly) and human cells, have dramatically changed our perception of the PCM in interphase cells from being an amorphous mass lacking definition to a remarkably ordered assembly.

The first discovery using these approaches was that major components of the PCM in an interphase centrosome, including pericentrin (pericentrin-like protein [PLP] in flies), Cep152 (Asterless in flies), Cep192 (SPD-2 in flies and worms), and Cdk5Rap2 (centrosomin in flies), form annular concentric rings around the centrioles. These rings differ in diameter, ranging in cross-section from approximately 200 nm, the diameter of the centriole itself, to about 500 nm. In other words, some are located close to the centriole surface, whereas others are positioned further out. Together, these proteins form a well-organized PCM proximal layer that, from the centriole surface, extends approximately 150–200 nm in width. Importantly, using domain-specific antibodies, it was shown that for two of these proximal layer proteins, namely pericentrin and Cep152, their C-termini are closely associated with the centrioles, whereas their N-termini are further away. Indeed, the physical distances measured between the N- and C-termini match the predicted lengths of these elongated proteins (about 150 nm), suggesting that they form rod-like filaments with one end (C-terminus) anchored to the centriole and the other end (N-terminus) extending out towards the cytoplasm. This model provides a rationale for how these proteins can define the diameter of the PCM through acting as molecular rulers that also limit the width of the PCM proximal layer in the interphase centrosome (Figure 1).

While this explains earlier structure–function studies on pericentrin showing that its C-terminal pericentrin-AKAP450 centrosome targeting (PACT) domain is required for centrosome binding, it remains to be determined at the molecular level how this, or the C-terminus of Cep152, interacts with centrioles. Current evidence for how the proximal layer attaches to centrioles suggests that it is likely to involve proteins that also play key roles in centriole duplication, such as CPAP (SAS-4 in flies and worms) and Plk4. Two populations of Cep192 have also been described: one closely attached to centriole walls that contributes to both centriole duplication and attachment of the PCM proximal layer, and a second further out in the PCM proximal layer that contributes to proximal layer organization. In vertebrate cells, PCM integrity is important for centriole duplication; however, this dependency may not be universal, as many species duplicate centrioles in interphase with very little PCM. Moreover, our current knowledge of how PCM proximal layer proteins functionally interact with core centriole duplication factors is sketchy at best. That said, genetic studies in worms recently identified a new factor, SAS-7, that potentially bridges between centrioles and SPD-2 acting upstream of both centriole duplication and PCM assembly.

The second discovery from the super-resolution microscopy studies was that, while some PCM proximal layer proteins form elongated filaments that extend from the surface of the centrioles, others are distributed amongst these filaments as, what have been termed, “branched matrix” components. These include Cdk5Rap2 and the second population of Cep192, and they depend on the filament proteins pericentrin and Cep152 for assembly into the PCM. However, the fact that pericentrin and Cep152 bind centrosomes independently suggests that there are at least two pathways for building the PCM; there is also evidence that branched matrix proteins show selectivity for particular filament proteins. Importantly, the branched matrix proteins directly anchor the γ-TuRCs, which are themselves highly organized macromolecular machines optimized for overcoming the nucleation barrier for tubulin polymerization. Furthermore, adaptor proteins are recruited, including NEDD1 and MZT1, that enable attachment of the γ-TuRC to the N-terminal centrosomin motif 1 (CM1) domain of Cdk5Rap2 while also stimulating microtubule-nucleating activity of the γ-TuRC.

Through structural biology-led studies on the fly homologue centrosomin, data from the Raft lab have identified how Cdk5Rap2 can self-assemble into a branched matrix. Tetramers of centrosomin are formed via interaction of two separate dimerization
Figure 1. Expansion and disassembly of the PCM upon mitotic progression and differentiation. This schematic figure provides a simplified overview of PCM organization. During interphase (center panel), the two centrioles (orange cylinders) are connected by an inter-centriolar linker with the bulk of the PCM associated with the older (mother) centriole. The size and shape of the PCM is defined by the two filament proteins, pericentrin and Cep152, which together create a branched matrix that provides binding sites for γ-TuRCs and their adaptor proteins. A second population of Cep192 is closely associated with proteins involved in centriole duplication at the centriole surface. In mitosis (right panel), PCM expansion results from the phosphorylation of multiple proteins, including pericentrin, Cdk5Rap2, and Cep192, by Plk1. This creates an outer expansive layer with gel-like properties (blue hatched circle) that is less well ordered but contains scaffolds that increase the microtubule nucleation capacity necessary for spindle assembly. Indeed, increased levels of these PCM proteins, together with additional centrosomally localized tubulin-binding proteins, such as chTOG and TPX2, may well allow microtubule nucleation to occur independently of γ-TuRCs (see inset). In contrast, in certain differentiated cells (left panel), disassembly of the proximal layer occurs with PCM proteins recruited to other non-centrosomal MTOCs. Note that the interphase centrosome is shown with two unduplicated centrioles typical of a G1 cell, whereas the mitotic centrosome contains a duplicated centriole pair that has also lost the inter-centriolar linker (mitotic cells have four centrioles, two in each spindle pole). CTD, C-terminal domain; NTD, N-terminal domain.

motifs, a leucine zipper (LZ) in the phospho-regulated multimerization (PReM) domain at the center of the protein and the centrosomin motif 2 (CM2) domain at the C-terminus12,23. However, tetramer formation requires phosphorylation within the PReM domain by the mitotic kinase Plk1 (or Polo in flies). Without this phosphorylation, centrosomin most likely exists as a closed intramolecular homodimer, and this may explain why very little PCM is assembled around interphase centrioles in flies.

Expansion of the mitotic PCM results from the assembly of disordered gel-like scaffolds

Super-resolution imaging suggests that, in contrast to interphase PCM, mitosis PCM is much less ordered. What has been clear for considerable time is that PCM expansion occurs through a process known as centrosome maturation that is absolutely dependent on Plk1 kinase activity, with Plk1 phosphorylating multiple PCM components, including pericentrin, Cdk5Rap2, Cep192, and, in worms, SPD-526–30. Worms lack pericentrin, Cep152, and Cdk5Rap2 but use the SPD-5 protein to perform analogous functions to Cdk5Rap2 and centrosomin in assembling the expansive PCM in mitosis13. Plk1 phosphorylation promotes the recruitment of significantly increased amounts of these proteins, thereby creating a new expansive outer layer to the PCM that substantially increases the overall centrosome diameter.

What had been less clear until recently is the molecular basis for how phosphorylation leads to PCM expansion. However, as indicated above, structural biology studies have now revealed how the phosphorylation of centrosomin promotes the formation of intermolecular tetramers25. Plk1 phosphorylation triggers the assembly of the 2:2 complex between the CM2 and LZ domains and leads, at least in vitro, to the generation of micron-scale assemblies reminiscent of centrosomes. Meanwhile, biochemical reconstitution experiments with the Caenorhabditis elegans PCM protein, SPD-5, have led to the elegant theory that the PCM can exist without a membrane by forming a phase-separated condensate in the cytoplasm31. Macromolecular crowding with agents such as polyethylene glycol drove purified SPD-5 to assemble into micron-scale assemblies that again were similar in size and shape to the PCM in vivo. Strikingly, these could form in suboptimal macromolecular crowding conditions if Plk1 was added, while SPD-2 also facilitated their assembly. Taken together, the current model is that Plk1 phosphorylation promotes the formation of gel-like, phase-separated condensates that, although lacking in uniform organization, most likely contain oligomeric scaffolds that enhance the capacity for microtubule nucleation.

What limits the steady-state size of this expanded mitotic PCM is likely to be a combination of a limiting cytoplasmic...
concentration of PCM components, together with the balance of localized Plk1 kinase activity and competing phosphatase activity. Aurora-A is another kinase that works in concert with Plk1 to promote centrosome maturation\textsuperscript{31}. In this case, different PCM proteins have been identified as Aurora-A substrates that directly facilitate γ-TuRC recruitment or, in the case of TACC and chTOG, stabilize microtubules. Plk1 can also phosphorylate NEDD1 either directly or indirectly via the activation of another mitotic kinase, NEK9, which increases the rate of γ-TuRC recruitment to the PCM\textsuperscript{41,33,34}. Importantly, many PCM components, including Cdk5Rap2, Cep152, and pericentrin, as well as additional effectors such as chTOG and TPX2, can directly bind α/β-tubulin heterodimers and promote microtubule nucleation independently of γ-tubulin by raising the local concentration required for spontaneous nucleation\textsuperscript{31,33,38}. Indeed, chTOG and TPX2 partition into the SPD-5 condensates generated \textit{in vitro} using purified proteins, where they concentrate tubulin and promote microtubule nucleation in the absence of γ-tubulin\textsuperscript{31}.

Interestingly, the additional proteins present in the mitotic PCM do not simply stick to the outside of the PCM. In flies, centrosomin is initially recruited to the inner regions of the PCM before moving outwards in a microtubule-dependent flaring mechanism that involves flux from the inner to the outer regions of the centrosome\textsuperscript{37}. This recruitment to the centriole surface is dependent on the Cep192 homologue, SPD-2, while centrosomin is in turn required to maintain SPD-2 at the centrosome\textsuperscript{39}. This could very well be crucial to limiting the centrosome size as Plk1 is tightly localized to the centriole surface\textsuperscript{31}. So, as PCM proteins are recruited in mitosis, they will initially be phosphorylated by Plk1 when close to the centriole; however, this phosphorylation is gradually lost as these proteins flux outwards. Hence, assuming the competing phosphatase is evenly distributed, there will be a diminishing gradient of Plk1-mediated substrate phosphorylation as one moves out through the PCM until eventually the threshold is passed for maintaining a phase-separated condensate\textsuperscript{31}.

As well as undergoing maturation at mitotic onset, centrosomes execute a process known as disjunction when the duplicated pairs of centrioles lose a proteinaceous tether, or linker, that holds the two centrosomes together during interphase\textsuperscript{39}. This raises the question of whether this centrosome linker is part of the PCM or rather an extension of the centrioles. To answer this question, we need to understand how the molecular elements of the linker, particularly the large coiled-coil proteins C-Nap1, rootletin, and LRRC45, physically interact with well-characterized PCM and centriole proteins\textsuperscript{40-42}. The linker extends between the proximal ends of the parental centrioles, and C-Nap1 can associate with the proximal-end centriole cartwheel component, Cep135. However, whether this interaction is direct or dependent on other PCM or centriole proteins is unknown. Meanwhile, rootletin has been described by immuno-EM to form an oligomeric filament that forms the major structural element of the linker, connecting the centriole proximal ends via C-Nap1\textsuperscript{43,44}. Other, smaller proteins have been identified as components of the linker, including centrin, Cep68, and β-catenin, and it will be important to explore how the linker is organized using super-resolution microscopy\textsuperscript{45-47}.

Unlike the outer expansive PCM layer that is assembled in mitosis, the linker structure is disassembled upon mitotic onset as a result of phosphorylation by the NEK2, and potentially NEK5, kinase\textsuperscript{46-50}. The coating of linker proteins with negative charge as a result of multi-site phosphorylation is the current favored model for linker disassembly\textsuperscript{51}. However, there are several lines of evidence that suggest that centrosome linker disassembly does not occur in isolation but is associated with reorganization of the PCM. Cdk5Rap2 and γ-tubulin levels are disturbed at the interphase centrosome by altered activity of NEK2 or NEK5; meanwhile, the linker protein Cep68 can interact with pericentrin and Cdk5Rap2, and loss of Cdk5Rap2 promotes premature centrosome disjunction\textsuperscript{49,51-53}.

**PCM disassembly in differentiated cells**

At the end of mitosis, the inactivation of Plk1 and subsequent dephosphorylation of PCM components, together with the loss of PCM fragments through flaring, lead to a return to the smaller size of PCM as defined by the proximal layer. The mechanism through which this expanded PCM is disassembled remains very poorly understood besides the requirement for the inactivation of Plk1, the dephosphorylation of Plk1 substrates, and the consequent reversal of scaffold assembly processes. Indeed, artificial tethering of Plk1 to centrioles through a PACT domain fusion prevents PCM disassembly and centrosome elimination in fly oocytes\textsuperscript{51}. However, there are also times in metazoan development when the proximal layer itself is disassembled under specific differentiation states and alternative MTOCs are formed in different cellular locations. A good example of this occurs during myogenesis. As myoblasts commit to differentiation, centrosome function is attenuated and microtubules instead become nucleated from the nuclear envelope, co-incident with the recruitment of PCM components to this membrane\textsuperscript{44-56}. Importantly, though, the mechanisms of PCM disassembly in these specialized circumstances are currently unknown.

Considerable attention is now being turned to the questions of, first, how the PCM proximal layer is disassembled and, second, which specific PCM proteins are recruited to the non-centrosomal MTOCs to enable microtubule nucleation from these sites. Muscle cells, for example, recruit a number of PCM proteins, including pericentrin and Cdk5Rap2, to the nuclear envelope\textsuperscript{55,56}. Hence, an important and intriguing question is whether a PCM-like proximal layer is assembled at non-centrosomal sites. Alternatively, these sites may assemble an oligomeric meshwork more reminiscent of the outer expansive PCM present in mitotic centrosomes. Like centrosome maturation and disjunction, the disassembly of the PCM proximal layer and then re-assembly of non-centrosomal MTOCs almost certainly depend upon post-translational modification of proximal layer proteins, although it could also involve changes in gene expression and protein degradation. In support of this, cyclin-dependent kinases regulate the formation of an MTOC at the apical membrane in \textit{C. elegans} intestinal cells, which in turn requires SPD-2 recruitment to the membrane\textsuperscript{57}.

The formation of a non-centrosomal MTOC also requires a site-specific receptor for anchoring microtubule nucleation
complexes in an alternative location. Recent studies have identified a muscle-specific isoform of the mammalian nuclear envelope nespin protein family, nespin-1α, that is required for the recruitment of pericentrin and Cdk5Rap2 as well as microtubule motor proteins to the nuclear envelope in muscle. Similarly, microtubule nucleation from the Golgi apparatus occurs in various differentiated cell types. In this instance, AKAP450, which shares homology with pericentrin through the conserved C-terminal PACT domain, is necessary for microtubule nucleation at the Golgi and its recruitment is dependent upon the Golgi-specific protein GM130. It remains to be determined, though, how these membrane proteins provide a platform for the assembly of these non-centrosomal MTOCs.

When considering the organization of non-centrosomal MTOCs, it is worth noting that land plants lack centrioles yet organize functional microtubule arrays. In interphase, plants possess cortical microtubule arrays that are distributed between the plasma membrane and large internal vacuole but without a focal point of organization. These microtubules appear to be primarily nucleated from γ-tubulin-associated complexes present at the plasma membrane or on other microtubules but then self-assemble into functional arrays through conserved microtubule-associated proteins, including chTOG. However, in early mitosis, a more organized and concentrated arrangement of microtubules is somehow formed, the metaphase band; this cortical ring of microtubules assembles at the cell equator at the site of future cell division, although the underlying assembly processes are not understood. As in animal cells, nuclear envelope breakdown allows these microtubules to interact with chromatin-nucleated microtubules to form the spindle. Interestingly, as in muscle cells, microtubule nucleation occurs from the nuclear, as well as plasma, membrane in interphase. However, while γ-tubulin complexes are again implicated, current evidence suggests that these are bound to nesprin family proteins rather than orthologues of classical PCM proteins. So, in this sense, some mechanisms of non-centrosomal MTOC organization may well be conserved; however, plant cells lack homologues of pericentrin and Cdk5Rap2, suggesting that other processes are involved.

**Pathological consequences of PCM disruption**

Increased centrosome numbers are a typical hallmark of cancer cells and can promote genomic instability through cell division errors and metaphase events through disturbance of cell polarity and migration. Meanwhile, inherited mutations in genes that encode core centriolar components can interfere with primary cilia function, causing multi-organ syndromes known as ciliopathies. In both cases, there are reasonably clear explanations for why centrosome defects contribute to disease pathology. Provocatively, though, some proteins encoded by ciliopathy genes also localize to the nucleus and have roles in the DNA damage response, suggesting an alternative pathological mechanism. What is less clear is why loss-of-function mutations in PCM components underlie two forms of growth defects: one that affects the whole body, primordial dwarfism, and one that is restricted to the brain, microcephaly.

Inherited mutations in either of the two proximal layer filament proteins pericentrin or Cep152, as well as the CPAP centriole duplication factor, lead to Seckel syndrome and microcephalic osteodysplastic primordial dwarfism (MOPD) type II. These primordial dwarfism syndromes exhibit profound growth retardation in every organ of the body and lead to miniature individuals. As overall size in mammals is dependent on cell number, primordial dwarfism reflects a reduction in cell number that must result from either decreased proliferation or increased cell death (or both). The simplest explanation would be that PCM defects prevent cell cycle progression by activating cell cycle checkpoints. Indeed, there is a wealth of evidence that centrosomes act as a meeting place to facilitate many different intracellular signaling events, including checkpoint activation, that have no direct role in microtubule nucleation or organization. Moreover, as for ciliopathies, some primordial dwarfism syndromes are associated with mutations in genes that regulate the DNA damage response, and the replication stress response in particular. For example, PCM proteins are necessary to recruit ATR, a key checkpoint protein that monitors replication fork integrity, to the centrosome to facilitate DNA damage signaling and loss-of-function ATR mutations also lead to Seckel syndrome. Hence, in the absence of an intact centrosome, cell cycle progression halts.

An alternative explanation is that defects prevent the PCM expansion necessary for proper spindle assembly and mitotic progression, thus leading to mitotic catastrophe and increasing cell death. However, a major conundrum in explaining primordial dwarfism is that these processes would have to affect all organs equally. It is possible that a uniform effect on stem cells during early development could cause a similar reduction in the progenitor pools that control particular tissue sizes. Close-range homeostatic mechanisms might also ensure an appropriate balance in the sizes of neighboring tissues, while more long-range effects could be exerted through endocrine organs that control tissue growth throughout the body. However, these are just ideas, and it is safe to say that we remain far from understanding the pathological basis for why PCM defects cause primordial dwarfism.

In contrast to primordial dwarfism, microcephaly is a specific reduction in brain size without affecting other organs. Intriguingly, the majority of genes implicated in autosomal recessive primary microcephaly encode centrosomal proteins, including both centriolar and PCM components. Notably, a different set of mutations in Cep152 cause microcephaly to those that cause Seckel syndrome, while mutations in Cdk5Rap2 were amongst the first to be identified in microcephaly patients. The current hypothesis for why centrosomal proteins are particularly prominent in this disease focuses very much on the disturbance of cell divisions within the neural progenitor pool. These undergo not only symmetric divisions in early development to expand the pool but also asymmetric divisions to generate differentiated neurons whilst concomitantly replenishing the progenitor pool. As with primordial dwarfism, PCM defects that interfere with cell cycle progression by activating checkpoints or induce cell death as a result of mitotic defects would explain the reduced neural progenitor pool. Indeed, mutations in other centrosomal genes that cause primary microcephaly, such as STIL, result in centriole amplification, and the experimental induction of centriole amplification by overexpression of Plk4 leads to a microcephalic condition in mice primarily through mitotic errors and excessive apoptosis. Equally, though,
errors in spindle positioning that prevent asymmetric divisions would cause failure to either generate differentiated cells or replenish the progenitor pool. The organ-specific nature of this developmental disease does strongly suggest that neural progenitors are especially sensitive to perturbations in these processes, and one can speculate that this is due to the heavy reliance on asymmetric divisions to generate large brains.

**Future perspectives**

The combination of super-resolution microscopy, structural biology, and biochemical reconstitution is beginning to provide the long-sought-after mechanistic details of how the PCM is organized and regulated. It has also revealed important similarities and differences between species. The PCM proximal layer present in interphase centrosomes is largely assembled around pericentrin/PLP and Cep152/Asterless in mammals and flies. However, neither of these proteins is found in worms. Mammals, flies, and worms all have a Cep192/SPD-2 protein that, at least in mammals, aids the connection between the centrioles and proximal layer. However, the amount of PCM present in interphase in flies and worms appears to be minimal, and it is not clear whether SPD-2 has a role at this time. In the expanded mitotic PCM, mammals and flies use the related Cdk5Rap2 and centrosomin proteins to form extended scaffolds, while this function is performed in worms by SPD-5 that, at least by sequence, is unrelated. Cep192/SPD-2 have a more obviously conserved role across all species in the expanded mitotic PCM. Indeed, the principle of an ordered proximal layer at the PCM when present in interphase and a more disordered gel-like scaffold in the expanded PCM in mitosis does appear to be universally shared.

These findings are stimulating specific structure–function studies into, for example, how the proximal layer regulates centrosome size, how the filament proteins are anchored to centriole walls, and how expansion and disassembly of the PCM are regulated by post-translational modifications. Relevant to this is whether the electron-dense centriolar satellites that surround centrosomes in some vertebrate cells represent supra- assemblies of PCM complexes that are being trafficked to centrosomes. Indeed, understanding the functional relationship between centriolar satellites and the PCM should explain whether there are proteins, such as PCM-1, that have their primary role in centriolar satellite integrity and why proteins implicated in ciliopathies seem to be over-represented in centriolar satellites.

Unfortunately, apart from the importance of Plk1, we still know little about the molecular events that control PCM expansion, disassembly, and relocation. A biochemical understanding of the roles of individual sites phosphorylated by Plk1, and Aurora-A for that matter, will be required as well as identification of the competing phosphatases. The recent demonstration of how Plk1 phosphorylation of *Drosophila* Cdk5Rap2 may promote oligomerization through exposing a dimerization interface is an excellent example. However, other types of modification will almost certainly contribute to this, including, for example, the acetylation or ubiquitylation of PCM proteins. It is also not clear how pericentrin, which binds centrioles through its PACT domain to create the proximal layer in interphase, is recruited to the outer layer during PCM expansion in mitosis. Equally, the need to understand mechanisms that drive PCM disassembly is highlighted by the finding that formation of the nuclear MTOC in myotubes is disrupted in certain forms of muscular dystrophy and may contribute to disease pathophysiology.

Perhaps unexpectedly, we have developed a reasonably coherent model for PCM assembly and recruitment of γ-TuRCs based on just a handful of proteins. Yet more than 100 proteins have been described as PCM components; so what are the rest doing? On one hand, there is good evidence, as indicated earlier, that the centrosome can act as a meeting place for signaling proteins that are not directly involved in microtubule organization. However, ruling out a role for a particular PCM protein in microtubule organization is not easy. In this regard, the *in vitro* reconstitution of PCM assembly will be a particularly valuable approach and should identify the minimal set of components required for different PCM functions, including efficient microtubule nucleation and anchoring.

Finally, we need to understand how genetic mutations that affect PCM components give rise to growth-related pathologies. Our current knowledge can broadly explain the reduction in cell number, be it from checkpoint-mediated cell cycle arrest, increased cell death resulting from mitotic errors, or spindle orientation defects that perturb stem cell pools. However, the uniformity of organs affected in primordial dwarfism and, conversely, the tissue specificity in primary microcephaly are striking and difficult to explain, particularly considering that different mutations in the same protein (Cep152) can give rise to one or other pathology. Answering these questions will come at least in part from complementing what we have learnt from super-resolution, structural biology, and biochemical reconstitution studies with gene-editing approaches not just in cells but also in whole animals.

**Abbreviations**

CM1 and 2, centrosomin motif 1 and 2; EM, electron microscopy; LZ, leucine zipper; MTOC, microtubule organizing center; PACT, pericentrin-AKAP450 centrosome targeting; PCM, pericentriolar material; PLP, pericentrin-like protein; PReM, phospho-regulated multimerization domain; γ-TuRC, γ-tubulin ring complex.

**Competing interests**

The authors declare that they have no competing interests.

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The referees who approved this article are:

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1 Tony Hyman *, Max Planck Institute of Molecular Cell Biology and Genetics, Dresden, Germany
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2 Bruce Bowerman University of Oregon, Oregon, USA
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3 Jordan Raff Sir William Dunn School of Pathology, Oxford University, Oxford, UK
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