CPAP promotes timely cilium disassembly to maintain neural progenitor pool

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Abstract

A mutation in the centrosomal-P4.1-associated protein (CPAP) causes Seckel syndrome with microcephaly, which is suggested to arise from a decline in neural progenitor cells (NPCs) during development. However, mechanisms of NPCs maintenance remain unclear. Here, we report an unexpected role for the cilium in NPCs maintenance and identify CPAP as a negative regulator of ciliary length independent of its role in centrosome biogenesis. At the onset of cilium disassembly, CPAP provides a scaffold for the cilium disassembly complex (CDC), which includes Nde1, Aurora A, and OFD1, recruited to the ciliary base for timely cilium disassembly. In contrast, mutated CPAP fails to localize at the ciliary base associated with inefficient CDC recruitment, long cilia, retarded cilium disassembly, and delayed cell cycle re-entry leading to premature differentiation of patient iPS-derived NPCs. Aberrant CDC function also promotes premature differentiation of NPCs in Seckel iPS-derived organoids. Thus, our results suggest a role for cilia in microcephaly and its involvement during neurogenesis and brain size control.

Keywords  brain organoids; CPAP; cilium; microcephaly; neural progenitor cell maintenance

Subject Categories  Cell Adhesion, Polarity & Cytoskeleton; Development & Differentiation; Molecular Biology of Disease

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Introduction

During interphase, the centriole within a centrosome nucleates the formation of the primary cilium, which is known to play key roles in signaling pathways (Berbari et al, 2009; Nigg & Raff, 2009; Jackson, 2011). The cilium is the primary microtubule-based organelle that is dynamically regulated, with assembly occurring during cell cycle exit, and disassembly coinciding with cell cycle re-entry (Jackson, 2011; Kim et al, 2011; Paridaen et al, 2013; Guemez-Gamboa et al, 2014). Thus, a delay or failure in cilium disassembly could act as a brake, retaining cells in G0/G1 and transiently preventing cell cycle progression (Jackson, 2011; Kim et al, 2011; Li et al, 2011).

Mutations in centrosomal proteins cause developmental disorders such as primary microcephaly and Seckel syndrome (Thornton & Woods, 2009; Kalay et al, 2011; McIntyre et al, 2012; Lancaster et al, 2013; Alcantara & O’Driscoll, 2014; Bazzi & Anderson, 2014; Insolera et al, 2014; Martin et al, 2014). Microcephaly is a neuro-developmental disorder leading to extreme reduction in brain size. In Seckel syndrome, the reduced brain size is coupled with pre- and postnatal reduction in body size. A mutation in the conserved centrosomal protein CPAP causes Seckel syndrome, characterized by dwarfism, low birthweight, microcephaly and intellectual disability (Al-Dosari et al, 2010).

The cell biological basis for microcephaly-linked disorders remains largely unclear. Most studies suggest a decline in the neural stem cell pool to eventually lead to a structurally normal but small-sized brain (Thornton & Woods, 2009; Alcantara & O’Driscoll, 2014). In the developing neural epithelium, stem cell numbers are rapidly expanded through its own symmetrical division (Rakic, 1995). This step is crucial to generate a large enough pool of neural stem cells that undergo asymmetric division producing cells that form various cortical layers in later stages of development (Alcantara & O’Driscoll, 2014). By impairing stem cell division even mildly, the proportion of symmetric divisions is concomitantly reduced, depleting the progenitor pool and limiting the total number of neurons that can be generated (Thornton & Woods, 2009; Alcantara & O’Driscoll, 2014). On the other hand, altering the cell
cycle length, in particular the G1 phase, contributes to switching of neural progenitors from proliferative to differentiating states (Calegari et al, 2005; Lange et al, 2009; Li et al, 2011).

Taken together, defective cellular division seems to underlie the depletion of a specific neural stem cell pool. However, how the centrosome or cilium regulates stem cell division to maintain a stem cell pool is largely unknown. Interestingly, till this day neither Seckel syndrome nor microcephaly disorders have been directly associated with clinical features of ciliopathies. Thus, the role of centrosome in microcephaly disorders remains unclear. However, since centrosomes template ciliogenesis, it is conceivable that altered ciliogenesis could underlie the pathophysiology of microcephaly. Recent studies have uncovered that the proper timing of cilium disassembly mediated by Nde-1 and TcTex-1 is critically regulated during cell cycle (Kim et al, 2011; Li et al, 2011; Maskey et al, 2015). Interestingly, mutations in Nde-1 and microtubule and kinetochore components causing microcephaly and ciliary defects highlight a possible role for the cilium in regulating NPCs to control the number of neurons generated during brain development (Alkuraya et al, 2011; Hu et al, 2014; Waters et al, 2015). However, the biological significance of these spatiotemporally interlinked processes namely cilium disassembly, cell cycle re-entry and neural progenitor cells (NPCs) differentiation, and the molecular mechanisms linking these cellular events together remain unclear.

Here, we investigate the effects of cilium disassembly on differentiation of NPCs derived from Seckel patient-induced pluripotent stem (iPS) cells. To model Seckel syndrome in a disease-relevant tissue, we developed human Seckel iPS-derived organoids. Our initial studies indicate Seckel cells to contain numerically normal centrosomes and mitotic spindle poles. However, upon further analysis, unusually long cilia, retarded cilium disassembly and delayed cell cycle re-entry were observed. This promoted premature differentiation of NPCs, identifying a previously unknown role for CPAP-mediated cilium disassembly in maintaining the neural stem cell pool. Our findings also uncover a yet-unknown feature of NPCs differentiation that is coupled with a timely cilium disassembly.

Results

Seckel fibroblasts have long cilia and reduced proliferation

CPAP plays a role in centriole formation and provides a scaffold for protein complexes during centrosome assembly (Hung et al, 2004; Gopalakrishnan et al, 2011, 2012; Lin et al, 2013; Zheng et al, 2014). However, the consequences of how a CPAP mutation (Seckel variant CPAP) underlies Seckel syndrome are unknown. To address this, we examined the centrosomes of Seckel patient-derived fibroblasts carrying a splice-site mutation in CPAP introducing a homozygous G-to-C transition in the last nucleotide of intron 11 causing the deletion of exons 11, 12, and/or 13 (Appendix Fig S1A-C; Al-Dosari et al, 2010).

In contrast to other microcephaly or Seckel cells that display numerically aberrant centrosomes, cells in the current study displayed correct numbers of centrosomes and mitotic spindle poles (Fig 1A; Rauch et al, 2008; Kalay et al, 2011; Hussain et al, 2012; Martin et al, 2014). However, interphase centrosomes displayed faint or no CPAP immunoreactivity. Importantly, compared to wild-type (WT) fibroblasts, Seckel cells recruited similar levels of other centrosomal proteins (γ-tubulin, pericentrin, and Cep152) (Fig 1A). These data suggest an altered function for the Seckel variant CPAP in an interphase-specific process.

As centrioles template cilium formation in interphase, we examined the cilium in asynchronously proliferating cells cultured in the presence of serum and found a significant proportion of Seckel cells (134 out of 321 cells) to contain unusually long cilia measuring up to 30 μm in length with a median length of 4.46 μm compared to 3.74 μm for WT cells (Fig 1B and Appendix Fig S2A). Ultrastructural analyses of Seckel cilia revealed disorganized abnormal ciliary appendages, and discontinuity of microtubular blades of basal bodies. These observations indicate aberrant ciliogenesis in CPAP-mutated Seckel cells (Fig 1C and D).

In vertebrate cells, an elongated cilium is associated with suppression of cell division via a delay in G1-S transition (Jackson, 2011; Kim et al, 2011; Li et al, 2011). As Seckel cells have long cilia, we speculated that they would also have a delayed G1-S transition. Indeed, our 72-h pulse labeling experiments using ethynyl-deoxyuridine (EdU) revealed a reduced number of Seckel cells with EdU incorporation indicating that fewer cells are in S-phase (Fig 1E). Furthermore, our fluorescence ubiquitination cell cycle indicator (FUCCI)-based analyses revealed increased proportions of Seckel cells retained in G1 (Appendix Fig S2B).

To identify the transcript(s) responsible for the Seckel phenotype, individual transcripts were overexpressed in ciliated human retinal pigment epithelial cells (RPE1), causing long cilia accompanied with cell cycle delay, thus suggesting that all the transcripts might contribute to the observed phenotypes (Appendix Fig S3A).
Figure 1.
Defective cilia disassembly in microcephaly

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Figure 2.

A

Serum starvation

0h 96h 1h 24h

B

WT

Serum starve

1 hr after serum stimulation

24 hrs after serum stimulation

WT

Seckel

C

Cilium appearance

Cilium resorption

CPAP/Arl13b

Acet-α-tubulin / Cep152

CPAP Cep152

Serum starved 1h after serum stimulation 24h after serum stimulation

Intensity (A. U)

400 300 200 100 0

CPAP Cep152

Serum starve 1h after serum stimulation 24h after serum stimulation

Intensity (A. U)

400 300 200 100 0

CPAP Cep152

Cilium length (μm)

Wild type

Seckel

Seckel

48 hrs after serum stimulation

24 hrs after serum stimulation

CPAP/Ar13b γ-tub/Arl13b

WT

Seckel

% of mitotic cells after 24 hrs

0 5 10

WT

Seckel

% of cyclin A positive cells

0 25 50 75 100

Serum starved 1h After serum stimulation

0 24h

WT

Seckel

% of mitotic cells

0 5 10

WT

Seckel

Duration (hrs)

0 1 2 4 6 8 10 12 14 16 18 20 22 24

Cilium length (μm)

Wild type

Seckel

0 2 4 6

WT

Seckel

% of mitotic cells

0 5 10

WT

Seckel

% of mitotic cells

0 5 10
and B). Importantly, to some extent, expression of WT CPAP using bacterial artificial chromosome (BAC) recombineering in Seckel fibroblasts rescued interphase CPAP localization and cilium lengths (Appendix Fig S3C).

These data suggest that the long cilium caused by a delay or failure in cilium disassembly acts as a brake, transiently preventing cell cycle progression, and might thus underlie the pathophysiology of Seckel syndrome.

Seckel cells display retarded cilium disassembly and cell cycle re-entry

To study cilium disassembly and cell cycle re-entry, we synchronized WT and Seckel fibroblasts at G0 by serum starvation for 96 h. Both Seckel and WT cells responded to this treatment acquiring equivalent ciliary length allowing us to estimate the rate of cilium disassembly upon serum stimulation (Fig 2A). Ciliated G0 WT fibroblasts displayed faint or no CPAP labeling at the base of the cilium (Fig 2B). However, CPAP levels were elevated 1 h after serum stimulation (Fig 2B and Appendix Fig S4B). After 24 h, CPAP was focused at centrosomes, concurrent with cilium disassembly and cell cycle re-entry. Remarkably, the elevated levels of CPAP correlated with biphasic cilium disassembly, the first occurring 0-1 h after serum stimulation and the second after 4-24 h (Pugacheva et al., 2007; Fig 2B and Di-ii). In contrast, Seckel cells lacked dynamic CPAP localizations and exhibited retarded cilium disassembly and cell cycle re-entry upon serum stimulation as indicated by the significant number of cells negative for cyclin A immuno-reactivity, a G1-S transition marker (Fig 2C and E, and Appendix Fig S4C). While most WT cells entered mitosis 24 h after serum stimulation, mitotic Seckel cells were only observed after 48 h. Interestingly, a fraction of these Seckel fibroblasts still harbored Arl13b-positive ciliary remnants which are ciliary membrane structures observed near to a centriole (Paridaen et al., 2013; Fig 2F). These data suggest that CPAP plays a role in resorbing cilia before mitosis (Fig 2F).

To establish the link between cilium disassembly and cell cycle progression, we overexpressed CPAP’s CC5 domain, that is, the region of aa 1073-1159 deleted in Seckel variant CPAP, in RPE1 cells. We found a significant proportion of CC5-GFP-expressing cells to have either short or no cilia with a concurrent increase in EdU incorporation, suggesting that accelerating cilium disassembly concomitantly accelerates cell cycle progression (Appendix Fig S5A).

CPAP provides a scaffold for the cilium disassembly complex

To gain insight into how CPAP coordinates cilium disassembly, we tested whether CPAP could interact with proteins, which are known to play roles in cilium disassembly such as Nde-1, Aurora A, HDAC6 and OFD1 (Pugacheva et al., 2007; Kim et al., 2011; Tang et al., 2013; hereafter referred to as Cilium Disassembly Complex, CDC). Indeed, cytoplasmic CPAP co-immunopurified with components of the CDC (CPAP-CDC; Fig 3A). Interestingly, CDC component levels were elevated in WT fibroblasts specifically 1 h after serum stimulation (Fig 3B). The elevated levels of these proteins correlated with CPAP levels and expression of cyclin A (Figs 2B and 3B). All of these processes occur concomitantly with the onset of cilium disassembly (Figs 2B and 3B).

We then immunopurified endogenous cytoplasmic CPAP complexes from asynchronously proliferating WT and Seckel fibroblasts and found the Seckel variant CPAP to show reduced interaction with CDC components, although expression of CDC components was also increased by serum stimulation in Seckel cells (Fig 3B and C, and Appendix Fig S5D). The elevated levels of CDC components upon serum stimulation in Seckel cells indicate that they can respond to serum stimulation. Thus, the observed delay in cilium disassembly and cell cycle re-entry is likely due to either Seckel variant CPAP’s lack of scaffolding function or the absence of CPAP at the cilial base causing inefficient targeting of CDC components (Appendix Fig S5B and C). The delayed cell cycle re-entry is further reaffirmed by the low levels of nuclear cyclin A in Seckel cells even upon serum stimulation. By performing glutathione S-transferase (GST) pull-down and immunopurification assays with CC5, we found the CC5 domain to be sufficient to mediate these interactions (Fig 3D).

To confirm CC5-dependent recruitment of CDC components during cilium disassembly, we analyzed CDC components in WT and Seckel fibroblasts synchronized by serum starvation and stimulation. In WT fibroblasts, CDC component levels were elevated after serum stimulation and recruited to the cilial base, concurrent with cilium disassembly and cell cycle re-entry. In contrast, although expression of CDC components was elevated in Seckel cells after serum stimulation, they failed to efficiently localize to the cilial base, resulting in retarded cilium disassembly and cell cycle re-entry (Fig 3B and Appendix Fig S6). These results suggest CPAP to play a specific role in spatiotemporal cilium disassembly by providing a scaffold for CDC components, independent of its role in centrosome biogenesis. Evidence for this is that Seckel cells display correct numbers of centrosomes and mitotic spindle poles (Fig 1A).
CPAP provides a scaffold for the cilium disassembly complex.

A CPAP co-immunopurifies with CDC components from HeLa extracts. Centrosomal proteins Cap350 and ODF2 do not co-purify with CPAP.

B Levels of CDC components (CPAP, Aurora A, OFD1, and Nde-1) are differentially regulated in response to serum starvation and stimulation in WT and Seckel fibroblasts. Specifically, they are elevated 1 h after serum stimulation, thus enabling cilium disassembly. Elevated CDC levels in WT cells correlate with an elevated level of CPAP as shown in Fig 2 as well as with the expression of cyclin A, a G1-S marker. GAPDH was used as a loading control indicating an equivalent amount of WT and Seckel fibroblast total proteins were loaded.

C WT CPAP (WT cells IP) but not Seckel variant CPAP (Seckel cells IP) co-immunopurifies with CDC components from asynchronously grown WT and Seckel fibroblast cell extracts. The IP samples were adjusted so that equivalent amounts of immunoprecipitated CPAP were used in samples. Seckel (5x) and WT (1x) fibroblast cell extracts were used as input controls to show the efficacy of the various antibodies. Note that CPAP abundance in Seckel cells is ~5 times lower than that of WT cells. Centrosomal proteins Cap350 and ODF2 do not co-purify with CPAP indicating the specificity of CPAP interaction with CDC components.

D GST pull-down (i) and immunopurification assays (ii) of CPAP’s CC5 domain demonstrate its sufficiency in mediating CPAP-CDC protein interactions. The CC5 fragment is expressed as a GFP fusion protein in HeLa cells and immunopurified using anti-GFP antibodies. HeLa extracts were used as input controls. Cap350 does not co-purify with CPAP. Both Cap350 and ODF2 are not pulled down by CPAP’s CC5 domain.
Retarded cilium disassembly causes premature NPCs differentiation

To further study the effects of cilium-mediated cell cycle re-entry delay on NPCs, we generated iPS cells, differentiated them into neuroectoderm followed by expansion into neural epithelium (Appendix Figs S7 and S8). Although control and Seckel neural epithelium developed neural rosettes (Perrier et al., 2004), Seckel rosettes were not only decreased in size with disrupted structure (Appendix Fig S8B). Nonetheless, they could further be differentiated into NPCs. Interestingly, these Seckel rosettes already displayed the pan neuronal marker TUJ1 suggesting premature differentiation into neuronal cells (Appendix Fig S8B). Similar to our findings in fibroblasts, Seckel NPCs lacked CPAP at the base of the cilium and failed to efficiently recruit CDC components to the cilium (Fig 4A). As a result, an increased proportion of ciliated Seckel NPCs contained long cilia and exhibited reduced proliferation and delayed G1-S transition (Fig 4A–D).

Altering G1-S transition influences the switch of NPCs from proliferation to differentiation (Calegari et al., 2005; Lange et al., 2009). To corroborate this, we treated WT human NPCs with olomoucine, which lengthens the G1-S transition of NPCs (Calegari & Huttner, 2003), and found this sufficient to trigger differentiation of NPCs into neurons (Fig 4E). We thus speculated that Seckel NPCs, exhibiting retarded cilium disassembly and extended G1-S transition, undergo premature neural differentiation. Indeed, under non-differentiating conditions, Seckel NPCs but not control NPCs differentiated prematurely into neurons (Fig 4F). We thus speculated that Seckel NPCs, exhibiting retarded cilium disassembly and extended G1-S transition, undergo premature neural differentiation. Indeed, under non-differentiating conditions, Seckel NPCs but not control NPCs differentiated prematurely into neurons (Fig 4F).

We then tested whether retarded disassembly could be a common mechanism underlying a subset of microcephaly disorders. To do this, we first downregulated CDC components in WT NPCs using siRNA-mediated depletion (for Nde-1 and OFD1) or small molecule (VX680)-mediated inhibition (for Aurora A). The treatments specifically caused retarded cilium disassembly, long cilia, premature differentiation of NPCs, accompanied by reduced cell cycle progression, suggesting that mutations in these genes might underlie NPC depletion resulting in microcephaly (Fig 5A–D). A mutation in Nde-1 causes microcephaly for which the underlying mechanism remains unclear (Alkuraya et al., 2011). Together, these data suggest that cilia disassembly, cell cycle progression, and neuronal differentiation are causally related to each other.

We then tested an additional patient mutation in CPAP, a homozygous mutation in its TCP domain (CPAP E1235V; Bond et al., 2005; Gul et al., 2006). To do this, we expressed CPAP E1235V using RNAi-resistant BACs together with CPAP-specific siRNA treatment in WT NPCs as previously described (Zheng et al., 2014). This resulted in premature differentiation of NPCs due to retarded cilium disassembly as indicated by the presence of long cilia (Fig SE–G). Taken together, this suggests that defective cilium disassembly is not only limited to the Seckel mutation we describe here, but could potentially represent a common mechanism regulated by a group of microcephaly genes specifically implicated in ciliogenesis.

Human Seckel iPS-derived organoids model microcephaly

To determine whether aberrant CDC function could lead to brain defects, we adopted the recently described three-dimensional culture system to generate organoids in vitro (Lancaster et al., 2013; Mariani et al., 2015). Starting with the same number of WT and Seckel iPS cells, matrigel-embedded droplets of neural epithelium were grown in spinner bioreactors (Appendix Fig S9A). We could obtain 2.5–3.5-mm organoids exhibiting brain-like architectures within 14–16 days (Appendix Fig S9B and C). Immunostaining revealed that organoids contained complex heterogeneous tissues displaying regions reminiscent of the ventricle zone (VZ) exhibiting proliferating apical progenitors (apical radial glial aRG) expressing nestin, Pax6, and phospho-vimentin.

Importantly, we found that the morphology and sizes of ventricular zones of these iPS-derived organoids, the region of our interest,
Figure 4. The EMBO Journal Vol 35 | No 8 | 2016
progenitors (apical radial glial cells, aRGs) using p-vimentin (Calegari et al., 2016) targeting neurogenesis has been shown to occur due to cell cycle delay of mitotic apical progenitors in the ventricular zone via vertical orientation accompanied by premature disassembly of the cleavage plane from horizontal to vertical (Rakic, 1995; Tapias et al., 2005; Basto et al., 2013). A reduced NPC pool is an accepted cause underlying microcephaly-linked disorders (Lancaster et al., 2013). However, the mechanisms involved in NPCs maintenance still remain unclear. The finding that several known microcephaly proteins are found in centrosomes suggested that a centrosomal mechanism can control neuronal number in a developing brain (Bond et al., 2005; Basto et al., 2016). We hypothesized that a centrosomal mechanism can control neuronal number in a developing brain (Bond et al., 2005; Basto et al., 2016).

were fairly consistent between organoids as was also demonstrated recently using iPS-derived organoids (Mariani et al., 2015). While we observed proliferating RGs on the apical side toward the lumen, we observed neuronal cells exhibiting neuronal proteins such as doublecortin (DCX) and TUJ1 at the basal side away from the lumen (Fig 6A and B). Inhibition of Ofd1 specifically due to aberrant CDC function caused by the loss of CPAP's CC5 domain. From these results, we suggested that a centrosomal mechanism can control neuronal number in a developing brain (Bond et al., 2005; Basto et al., 2016). A reduced NPC pool is an accepted cause underlying microcephaly-linked disorders (Lancaster et al., 2013; Alcantara & O'Driscoll, 2013; Guemez-Gamboa et al., 2014). However, the mechanisms involved in NPCs maintenance still remain unclear. The finding that several known microcephaly proteins are found in centrosomes suggested that a centrosomal mechanism can control neuronal number in a developing brain (Bond et al., 2005; Basto et al., 2016).

To test whether expressing WT CPAP or CC5 alone could rescue the phenotype, we performed co-electroporation of CPAP-GFP, CC5-GFP, or GFP-alone into 6-day-old Seckel organoids and analyzed them 5 days later. As control, GFP-alone was electroporated. In analyzing regions of CPAP- or CC5-electroporated tissues, we found CPAP to partially rescue certain aspects of Seckel phenotype. This partial rescue could be due to CPAP overexpression itself (Kohlmair et al., 2009; Tang et al., 2009). However, the rescue effect was more prominent in CC5-transfected Seckel organoids as similarly observed with our NPCs experiments (Fig 6C).

Although Seckel organoids displayed distinct brain-like regions, they were smaller in size, exhibiting reduced neural epithelial tissues and progenitor regions with disorganized VZ and larger lumen (Fig 6B and C). Interestingly, these extended TUJ1-positive structures showed distinct GAD67-positive protrusions, dendritic spines of inhibitory GABAergic nature suggesting that our in vitro differentiating conditions are physiological (Guidotti et al., 2000).

Seckel organoids displayed increased numbers of mitotic aRGs with cleavage planes perpendicular to the ventricular surface suggesting a premature switching of the cleavage plane. These data agree with studies demonstrating that CPAP is critical for controlling the cleavage plane of mitotic neuronal progenitors (Insolera et al., 2014; Bazzi & Anderson, 2014; Garcez et al., 2015; Appendix Fig S10). Finally, when analyzing apical progenitor cilia extending into the lateral ventricle, Seckel organoids showed increased numbers and longer cilia compared to WT organoids, suggesting retarded cilium disassembly consistent with our results from fibroblasts and NPCs experiments (Fig 6C).

Figure 5. Downregulation of CDC components or a CPAP natural microcephaly mutation causes neural differentiation.

A-B: siRNA-mediated downregulation of CDC components in WT NPCs causes longer cilia (Ofd1 \( \ast \ast \ast P < 0.01 \), Nde1 \( \ast \ast \ast P < 0.01 \)) and differentiation of NPCs into TUJ1-positive cells (Ofd1 \( \ast \ast \ast P < 0.001 \), Nde1 \( \ast \ast \ast P < 0.01 \)). In controls, a non-targeting siRNA (scrambled) is used. Western blots on the right validate siRNAs-mediated Nde1 and Ofd1 depletion. Error bars are SEM. Ordinary two-way ANOVA followed by Sidak’s multiple comparisons test. \( n = 3 \) independent experiments. 275 cells for scrambled, 423 cells for Ofd1 siRNA, and 255 cells for Nde1 siRNA. Arrow marks the centrosome. Scale bar, 1 μm.

C: VX680 (0.5 μM), a concentration that will specifically allow neuronal differentiation was used to inhibit Aurora A activity. Bar diagrams show quantifications. Compared to DMSO-treated control NPCs, a higher proportion of VX680-treated NPCs are ciliated (\( \ast \ast \ast P < 0.003 \)), exhibiting longer cilia (\( \ast \ast \ast P < 0.001 \)), and show TuJ1-positive neurons (\( \ast \ast \ast P < 0.005 \)). Error bars are SEM. For ciliated cells and cilium length: \( n = 3 \) independent experiments. 164 cells for DMSO-treated and 77 cells for VX680-treated. For NPC differentiation: \( n = 4 \) independent experiments. 286 cells for DMSO-treated and 118 cells for VX680-treated. Scale bar, 1 μm.

D: Cell cycle progression of siRNA- or VX680-treated WT NPCs compared to control assessed by EdU incorporation for 24 h. Inhibition of Ofd1 (\( \ast \ast \ast P < 0.05 \), Nde1 \( \ast \ast \ast P < 0.01 \), and Aurora A \( \ast \ast \ast \ast P < 0.0003 \)) caused significantly reduced incorporation of EdU. Error bars are SEM.

E-G: Western blots show that CPAP-targeting siRNA treatment depletes endogenous CPAP. Both low and high exposures are given (f). Increased cilium length and NPC differentiation upon expression of RNAi-resistant CPAP E1235V compared to WT CPAP (f, G). Error bars are SEM. \( \ast \ast \ast P < 0.001 \) (cilium length), \( \ast \ast \ast P < 0.05 \) (differentiation), \( t \)-test. For cilium length: \( n = 4 \) independent experiments. 25 cells for CPAP, 47 cells for CPAP E1235V. For differentiation: \( n = 4 \) independent experiments. 145 for CPAP and 81 for CPAP E1235V. Scale bar, 1 μm.
Figure 5.

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illustrate a surprising role for the primary cilium in stem cell...incentive, what remains elusive is why Seckel CPAP lacking the CC5 domain results in long cilia. It is known that the CPAP-tubulin interaction is required for centriole duplication, linking the importance of timely cilium disassembly mediated by CPAP-CDC to cell cycle re-entry for neurogenesis and brain size control.

Most studies to date have focused on analyzing centrosomes, as microcephaly-related proteins are mainly centrosomal components, with clinical features of ciliopathies not reported to be associated with Seckel syndrome or microcephaly disorders. The results from our current study reveal that structurally and numerically aberrant centrosomes are not the only cause underlying microcephaly; instead, disrupted ciliary function could also lead to microcephaly-linked disorders. Our results also rule out the possibility of apoptosis-mediated NPCs loss in the currently studied microcephaly mutant. This suggests that disorders caused by defective cilium disassembly could potentially be grouped as ciliopathic diseases.

Although our experiments substantiate the previously unknown function of CPAP as a negative regulator of cilary length independent of its role in centrosome biogenesis, what remains elusive is why Seckel CPAP lacking the CC5 domain results in long cilia. It is known that the CPAP-tubulin interaction is required for centriole and cilium elongation (Tang et al., 2009; Wu & Tang, 2012). Evidence comes from a functional study where a non-tubulin-binding CPAP variant results in a short cilium, suggesting that CPAPs’ N-terminal tubulin binding capability could determine cilium length (Wu & Tang, 2012). From this observation, we speculate that the Seckel variant CPAP (where a part of its CC5 domain is deleted) could have an altered tubulin binding activity possibly accounting for an elongated cilium.

Alternatively, as shown by our ultra-structural studies that Seckel cilia display disrupted cilary appendages near the transition zone, we postulate that aberrant CDC recruitment could impair ciliary functions such as protein entry and exit from the cilium, thus regulating its length. Therefore, it would be important to study the ultra-localization of CDC components at the basal body and how they interact with each other to form structurally normal cilia. It is noteworthy that proteins mutated in ciliopathies are localized to the transition zone (Garcia-Gonzalo et al., 2011; Huang et al., 2011; Li et al., 2011).
Figure 6.
Our results also suggest a model where retarded cilium disassembly-mediated delay in cell cycle re-entry is sufficient to trigger premature differentiation of NPCs. In this context, it appears that the cilium plays a crucial role in determining asymmetric cell division. How timely cilium disassembly directly or indirectly regulates the apical progenitor cleavage plane remains unclear, thus warranting further study. Since the primary cilium is an extracellular antenna of vertebrate cells playing key roles in signaling pathways, it is likely that it traffics specific signaling cues in a spatiotemporal manner for faithful asymmetric cell division. We hope that our current findings provide new insights into these open questions and substantiate the importance of the cilium in asymmetric cell division.

Our CPAP mutant phenotypes seem to overlap with those observed in other genes involved in Seckel syndrome and microcephaly (Bond et al., 2005; Gul et al., 2006; Kalay et al., 2011; Alcantara & O’Driscoll, 2014; Martin et al., 2014). Thus, it would be interesting to test whether the observed “cilium-check point” in the current study serves as a common mechanism in organ size control. Indeed, our findings that depletion of CDC components leads to phenotypes similar to that observed in Seckel cells could represent a conserved mechanism among microcephaly causing genes that are implicated in ciliogenesis. This previously unknown role for cilium disassembly in fate determination of somatic stem cells may not only be restricted to NPCs but could also be true for other paradigms of organ size control and homeostasis during development.

Materials and Methods

Molecular biology and cell culture

WT skin fibroblast or RPE1 cells (Stemgent BJ Human Fibroblasts, Cat. No. 08-0027, Stemgent, San Diego, USA), and skin fibroblasts derived from a Seckel patient were grown in fibroblast medium containing Dulbecco’s modified Eagle’s medium (DMEM), 10% fetal bovine serum (FBS), 0.1 mM MEM non-essential amino acids (NAA), 100 U/ml penicillin, 100 μg/ml streptomycin (all from Life Technologies GmbH, Darmstadt, Germany), and 0.05 mM β-mercaptoethanol (Sigma-Aldrich, Munich, Germany). HeLa cells were grown under standard conditions in DMEM supplemented with 10% FBS. For serum starvation experiments, cells were grown in fibroblast medium without serum for the described time period and stimulated with 10% FBS-containing medium. For transfecting fibroblasts, the Neon Transfection System (Life Technologies) was used with the following parameters: 1 × 10^7 cells/ml in 10 μl buffer containing 1 μg plasmid DNA or 250 ng BAC DNA per 1 × 10^6 cells, pulse voltage 1,650 V, pulse width 10 ms, and 3 pulses.

For lentiviral transductions, GFP-tagged CPAP with different mutations were cloned into pLenti6.3 vector (3rd-generation lentivirus). Viral transductions were tested after 48 h and subsequently used for further analyses.

Western blot

Protein samples were resolved in 8 or 12% acrylamide gels and transferred to nitrocellulose membranes. After incubating with primary antibodies overnight at 4°C, the blots were treated with secondary antibodies at RT for 1 h. Super Signal West Pico or Femto Chemiluminescent substrates (Pierce) were used for detection. Antibody dilutions for Western blots: monoclonal mouse anti-CPAP (1:50), rabbit-CDK5RAP2 (1:3,000, T. K. Tang), mouse anti-γ-tubulin (1:3,000; Sigma-Aldrich), mouse anti-α-tubulin (1:3,000; Sigma), mouse anti-HDAC6 (1:1,000; Sigma-Aldrich), rabbit anti-Nde1 (1:1,000; Proteintech), mouse anti-Aurora A (1:1,000; Cell Signaling), rabbit anti-OFD1 (1:3,000; J. Reiter), mouse anti-cyclin A (1:1,000; Sigma-Aldrich), mouse anti-GFP (1:3,000; Roche), and peroxidase-conjugated secondary antibodies were used at 1:3,000 (Life Technologies).

Immunofluorescence and light microscopy

For immunofluorescence, mouse anti-γ-tubulin (1:500; Sigma-Aldrich), rat anti-α-tubulin (1:50, Chemicon), mouse anti-CPAP (1:25), rabbit anti-Cep152 (1:500, E. Nigg), rabbit anti-PCNT (ab4448, Abcam, Cambridge, USA; 1:1,000), rabbit anti-Arl13b (1:100, Proteintech, Manchester, UK), mouse anti-acetylated tubulin (1:500, Sigma-Aldrich), rabbit anti-Nde1 (1:50, Proteintech, Manchester, UK), mouse anti-Nastin 4D11 (1:100, Novobis Logicals, Cambridge, USA), rabbit anti-Tuj1 (1:100, Sigma-Aldrich), mouse anti-Oct4 (1:200, Santa Cruz), rabbit anti-Nanog (1:200, Santa Cruz, Heidelberg, Germany), mouse anti-Pax6 (1:50, DSHB, Iowa University, Iowa, USA), mouse anti-cyclin A2 (ab16726, Abcam), mouse anti-phosphorylated vimentin (1:100, 4A4, Biozol, Eching, Germany), rabbit anti-OFD1 (J. Reiter, 1: 50), goat anti-AFP (1:50 Santa Cruz), mouse anti-MF-20 (1:100, DSHB, Iowa) antibodies were used. Secondary antibodies Alexa Fluor dyes conjugated goat/ donkey anti-mouse or anti-rabbit were used at 1:1,000 (Life Technologies). DAPI (1 μg/ml, Sigma) stained DNA.

For light microscopy, cells were fixed in 3.7% paraformaldehyde (PFA) or ice-cold methanol, permeabilized with 0.5% Triton X-100 in PBS for 10 min, and blocked with 0.5% fish gelatin in PBS for 2 h at RT. Antibody labeling was performed for 1 h at RT or overnight at 4°C followed by three washes in PBS. Confocal images were collected using an Olympus Fluoview FV 1000 scanning confocal microscope and processed using Adobe Photoshop. Organoids were fixed in 3.7% PFA for 2 h, stored in 30% sucrose solution at 4°C overnight and then embedded in Tissue Tek (Weckert, Kitzingen, Germany), and stored at −80°C. Organoid cryosections were then cut in 20-μm-thin sections for immunofluorescent stainings.

To count nestin-positive neurons in organoid slices, we performed counting in Z-serial 1-μm slices. Special care was taken to prevent overlaps. When overlaps were seen, those cells were not considered. For clarity, we will describe it in the method section.

Statistics calculations

Statistical analyses, for example, Student’s t-test or one-way ANOVA followed by Tukey’s multiple comparisons were performed using GraphPad Prism version 6.0d.
Electron microscopy

Asynchronously grown fibroblasts were pelleted, fixed with 2% glutaraldehyde (Electron Microscopy Sciences), and processed for electron microscopy as previously described (Gopalakrishnan et al., 2010; Zheng et al., 2014). The embedded cell pellets were ultra-thin-sectioned (80 nm), counterstained, and visualized for ultra-structural details of centrosomes using a Zeiss 10A electron microscope. Images were processed using Adobe Photoshop.

Pulse chasing experiments, chemical compounds, and RNAi treatments

For cell proliferation estimation, EdU was pulsed in cells for 24–72 h before fixation, and stained according to the manufacturer’s instructions (Click-IT EdU assay kit, Life Technologies). For inhibitor experiments, wherever necessary, NPCs were treated with the Cdk1 inhibitor, olomoucine (Sigma-Aldrich; 25 μM), or VX680 (Selleckchem; 0.5 μM) for 4 days, fixed, and analyzed by microscopy.

For RNAi treatments, NPCs were electroporated using the Neon Transfection System with 100 nmol of siRNA targeting human NDE1 and OFD1 mRNA. NDE1: siGENOME Human NDE1 (54,820) siRNA-SMARTpool (M-020625-00-0005) OFD1: siGENOME Human OFD1 (8481) siRNA-SMARTpool (M-009300-02-0005). Negative controls were performed with non-targeting siRNA (scrambled; 100 nmol, AllStars Negative control siRNA; Qiagen). Electroporation using 100 nmol siRNA was repeated on day 3. siRNA-treated NPCs were fixed on day 7 for further analysis.

Protein expression, purification, and GST pull-down assays

The region deleted in the Seckel syndrome patient (residues 1,073–1,159 of CPAP) was cloned into the pGEX vector (Amersham Biosciences) containing an N-terminal GST tag and the eGFP-C1 vector (Clontech). The protein was expressed in E. coli BL21(DE3) (New England Biolabs) and induced by 0.5 mM IPTG overnight at 18°C. The recombinant protein was affinity-purified from supernatant using GST-sepharose beads (GE Healthcare). The conjugated protein was then treated with HeLa cell extracts and further analyzed by Western blots.

Immunopurification of CPAP complexes

Protein G beads were coated with anti-CPAP antibodies overnight at 4°C, mixed with HeLa, WT, or Seckel fibroblast cell extracts, respectively, and incubated at 4°C for 4 h. Cell extracts were prepared in a buffer containing 80 mM BRB, 100 mM KCl, 1 mM MgCl2, 1 mM EGTA, protease inhibitor cocktails, and 1 mM PMSF. The extracts were centrifuged at 100,000 g, and the supernatant was collected. These high-speed extracts were then used for immunopurification. The protein-bound beads were washed with BRB buffer containing 0.1% Triton X-100 and 100 mM NaCl and washed twice with high-salt buffer containing 500 mM salt. After a final wash with buffer containing 100 mM NaCl, the samples were eluted using Laemmli buffer for Western blot analyses.

To immunopurify endogenous proteins that interact with CPAP 1073–1159, stable HeLa cell line expressing the fragment GFP-1073-1159 (CC5-GFP) was generated. Cell extracts prepared from these stable cell lines were used for immunopurification using anti-GFP-coated magnetic beads as per the manufacturer’s instructions. Complexes were then eluted for Western blot analyses.

Patient iPS cell reprogramming

Skin fibroblasts were transfected with pCXLE-hOct3/4-shp53, pCXLE-hSox2-hKlf4, pCXLE-hLin28-hL-Myc (Addgene, Cambridge, USA) using the Neon Transfection System according to the manufacturer’s protocol. After 5 days, the medium was replaced with DMEM/F12 medium supplemented with Glutamax, 20% knockout serum replacer, 1% NAA, 0.1 mmol/l β-mercaptoethanol (Life Technologies), and 50 ng/ml bFGF (PeproTech, Hamburg, Germany). At day 10, cells were trypsinized and maintained on irradiated mouse embryonic fibroblasts (MEFs) in human iPS cell medium. Colonies were selected based on morphology (Okita et al., 2011; Yu et al., 2011), further passaged, and expanded. As quality control for iPS clones, we performed alkaline phosphatase (Vector Blue Alkaline Phosphatase Substrate Kit; Vector Laboratories, SK5300) and pluripotent markers staining.

Endodermal and mesodermal differentiation

Human iPS cells were grown in serum-containing medium for 6 days, detached by accutase treatment, and then replated on collagen type I-coated dishes. Endodermal differentiation was induced by low-serum-containing medium supplemented with HGF, Oncostatin M, and 100 nM dexamethasone for 10 days. Mesodermal differentiation was induced by serum-free medium containing insulin and 10 μM SB431542 for 20 days.

Differentiation and characterization of NPCs

Similar amounts of feeder-free human WT or patient iPS cells were differentiated into neural progenitor cells (NPCs) in STEMdiff Neural Induction Medium (Stem Cell Technologies, Vancouver, Canada). After 5 days, formed neurospheres were collected and cultured on poly-L-ornithine (PLO)/laminin-coated dishes with daily medium change. Neural rosettes were selected after 7 days using neural rosette selection medium and recultured. NPCs were split every 5–7 days and further maintained in STEMdiff Neural Progenitor Medium under non-differentiating conditions.

Generation of organoids

Organoids were generated as described before (Lancaster et al., 2013) with slight modifications. Neurospheres were generated, collected after 5 days, and embedded in Matrigel (Corning, Tewksbury, USA) drops. The embedded droplets were differentiated in suspension culture in 1:1 DMEM/F12 and Neural Basal Medium (v:v), 1:200 N2, 1:100 B27 w/o vitamin A, 1:100 t-glutamine, 0.05 mM MEM non-essential amino acids (NAA), 100 U/ml penicillin, 100 μg/ml streptomycin, 1.6 g/l insulin (Sigma-Aldrich), and 0.05 mM β-mercaptoethanol (all from Life Technologies if not stated). After 4 days, matrigel-embedded neurospheres were transferred to rotary suspension culture in spinner flasks (IBS, Integra biosciences) containing the same differentiation medium with
addition of 0.5 μM dorsomorphin (Sigma-Aldrich). Organoids were collected and analyzed after 14, 28, and 42 days of rotary culture.

To culture single slices of organoid, first organoid slices of about 100 μm were made using previously gelatin-embedded whole cerebral organoids. The slices were grown in wells containing media supplemented with 0.5 μM dorsomorphin, 10 ng/ml BDNF, and 10 ng/ml GDNF for 2 weeks until the extended neuronal structures are visually observed.

Organoid electroporation

Electroporation was carried out using two platinum electrodes and the Intracel TSS20 OVODYNE Electroporator with five square pulses of 80 V within 20 ms width. A total of 1 μg plasmid (CPAP or CPAP’s CC5) was co-electroporated with 2 μg/μl GFP-expressing vector mixed with Fast Green solution (Sigma) at a ratio of 2:1 into 6-day-old organoids at 3°C. The slices were grown in wells containing same media supplemented with 0.5 μM dorsomorphin, 10 ng/ml BDNF, and 10 ng/ml GDNF for 2 weeks until the extended neuronal structures were further grown in suspension cultures for 14 days for further analyses.

Expanded View for this article is available online.

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Author contributions

JG and EG conceived the project, designed experiments, and wrote the manuscript. EG performed experiments and analyzed the data. AW performed biochemical experiments. MLG and AR performed molecular experiments with BACs. NST and TS for reagents and experiments for iPS generation. JH and FSA analyzed the data. JG directed and supervised the experiments. Avidor-Reiss T conceived the project and designed experiments. JH and FSA performed biochemical experiments. JG and EG wrote the manuscript. GC and MG performed ultra-structural analyses. FC and FN supported with manuscript. EG performed experiments and analyzed the data. AW and EG conceived the project, designed experiments, and wrote the manuscript. Elke Gabriel et al. (2014) Defective cilia disassembly in microcephaly. The EMBO Journal 33: 1061–1069

Conflict of interest

The authors declare that they have no conflict of interest.

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