Fbxo41 Promotes Disassembly of Neuronal Primary Cilia

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Neuronal primary cilia are signaling organelles with crucial roles in brain development and disease. Cilia structure is decisive for their signaling capacities but the mechanisms regulating it are poorly understood. We identify Fbxo41 as a novel Skp1/Cullin1/F-box (SCF) E3-ligase complex subunit that targets to neuronal centrioles where its accumulation promotes disassembly of primary cilia, and affects sonic hedgehog signaling, a canonical ciliary pathway. Fbxo41 targeting to centrioles requires its Coiled-coil and F-box domains. Levels of Fbxo41 at the centrioles inversely correlate with neuronal cilia length, and mutations that disrupt Fbxo41 targeting or assembly into SCF-complexes also disturb its function in cilia disassembly and signaling. Fbxo41 dependent cilia disassembly in mitotic and post-mitotic cells requires rearrangements of the actin-cytoskeleton, but requires Aurora A kinase activation only in mitotic cells, highlighting important mechanistical differences controlling cilia size between mitotic and post-mitotic cells. Phorbol esters induce recruitment of overexpressed Fbxo41 to centrioles and cilia disassembly in neurons, but disassembly can also occur in absence of Fbxo41. We propose that Fbxo41 targeting to centrosomes regulates neuronal cilia structure and signaling capacity in addition to Fbxo41-independent pathways controlling cilia size.

Primary cilia are antennae-like sensory organelles that project from the plasma membrane of a wide variety of cells, including neurons. They are composed of a microtubule-based core structure that elongates from a membrane-anchored modified centriole, the basal body1. Both primary cilia membrane and cytosol content are isolated from the rest of the cell. This allows cilia to concentrate several signaling pathways, and act as a sensory organelle for extracellular and intracellular cues2. For example, the Sonic hedgehog (Shh) cascade, a key signaling pathway during brain development, has been shown to depend on ciliary integrity3,4. Mutations disrupting ciliary structure can lead to debilitating disorders, referred to as ciliopathies, which have pleiotropic clinical features5–7. Among the 87 genes associated with ciliopathies, 77 genes have been linked to neurological deficits in humans8, such as altered brain anatomy, obesity, and intellectual disability9.

Cilia structure is dynamically modulated in both mitotic cells and neurons, with important functional consequences. As such, cilia assembly and disassembly are highly regulated processes, and can be induced by several signaling cues10 in mitotic cells. The most well-studied cue is cell cycle entry where cilia disassemble before cell division and re-assemble after the end of the cell cycle11–13. One canonical mechanism of ciliary disassembly in mitotic cells, is through the centrosomal kinase Aurora A: activation of Aurora A leads to activation of HDAC6, which deacetylates and destabilizes axonemal microtubules14. Primary cilia are also present in post-mitotic cells, such as neurons. Neuronal cilia structure is dynamic, and depends on developmental stage and final layer position of the cell15,16. In migrating cortical interneurons, cilia length is highly dynamic17 and cilia of olfactory neurons in C. elegans remodel in a sensory signaling-dependent manner18. This implies the presence of machinery that senses extracellular cues and modulates ciliary architecture. So far, however, the mechanisms through which neuronal primary cilia length is regulated remain elusive.

One powerful modulatory system to control cilia structure and function is the ubiquitin proteasome system (UPS). The UPS selectively modulates the cellular protein pool to temporally and spatially control cellular activities. UPS components have been shown to accumulate at the centrosome19–21, and are able to control ciliary length19,23. F-box proteins are substrate binding adaptors of a Skp1/Cullin1/F-box (SCF) E3-ligase complex22 that
confer selectivity to the UPS by selecting the target of ubiquitination. Fbxo41 is a brain enriched F-box protein, with high expression in hippocampal neurons, where it accumulates in centrioles from which primary cilia emanate, making it a prime target for regulating neuronal primary cilia.

In this study we show that Fbxo41 assembles into an SCF complex, targets to neuronal centrioles, and its accumulation promotes disassembly of primary cilia. Fbxo41 requires its Coiled-coil and F-box domains for targeting to centrioles. Centriolar Fbxo41 levels show strong inverse correlation with cilia length, but not in mutants with disrupted Fbxo41-Skp1 interaction. We show, for the first time, that neurons treated with the phorbol ester PDBU, but not canonical network-activity modulators (Gabazine, APV, DNQX or TTX) have shorter cilia and increased centrosomal Fbxo41 expression. However, ciliary disassembly induced by PDBU also occurs in absence of Fbxo41. The effect of Fbxo41 in cilia disassembly in mitotic cells can be rescued by inhibiting the canonical Aurora A pathway, or perturbing actin dynamics by cytochalasin D. The latter compound also prevents Fbxo41-dependent cilia shortening in neurons. Finally, we show that Fbxo41 disturbs Shh signaling, a prominent ciliary pathway. We propose a mechanism where neurons can shorten their cilia by regulating centriolar levels of Fbxo41, which affects ciliary signalling capacity.

**Results**

**Fbxo41 is an SCF-complex subunit that targets to neuronal centrioles.** Generally, F-box proteins are modular substrate binding adaptors of a Skp1/Cullin1/F-box (SCF) E3-ligase complex. Since this has not been established for every F-box protein family member, we tested whether Fbxo41 assembles into a SCF-complex by expressing EGFP- or FLAG-tagged Fbxo41, Skp1 and Cullin1 in HEK293T cells, and performing immunoprecipitation experiments. Indeed, Fbxo41 associated with Skp1 and Cullin1, albeit less efficiently than Fbx23, which was included as a positive control. Deleting Fbxo41 AF-box or mutating Fbxo41 W577A (the F-box domain) abolished these interactions, confirming that, like other F-box proteins, Fbxo41 assembles into SCF-complexes via an essential F-box domain (Fig. 1a and Supplementary Fig. S1).

In the brain Fbxo41 is exclusively expressed in neurons and Fbxo41 levels increase with age in the cerebellum. We confirmed that Fbxo41 expression increases in whole-brain lysates throughout development (Fig. 1b), and show that Fbxo41 is expressed in cultured neurons, but not in astrocytes (Fig. 1c). Skp1 and Cullin1 are enriched at centrosomes of dividing cells where their ubiquitin-ligase activity is essential for many centrosomal functions controlling cell division. Although our antibodies did not allow immunoprecipitation of sufficient amounts of Fbxo41 from brain lysate (Supplementary Fig. S1), we could assess the presence of Fbxo41 at centrosomes in primary hippocampal neurons. Endogenous Fbxo41 located throughout the cytosol with distinct enrichments in the soma, which co-localized with the pericentriolar protein Pericentrin (Fig. 1d), but Fbxo41-EGFP did not colocalize with several other somatic organelles (Supplementary Fig. S2). Super-resolution stochastic optical reconstruction microscopy (STORM) revealed that Fbxo41-EGFP resides adjacent to centrosomal protein 135 (Cep135), a centriole proximal-end protein.

**Centrosomal targeting of Fbxo41 requires its F-box and coiled-coil domains.** The closest homologue of Fbxo41, ZNF365, is a brain-enriched zinc finger protein that targets to centrosomes via its Coiled-coil domain. We further characterized the Fbxo41 domains required for centrosomal targeting by quantifying centrosomal enrichment of several other Fbxo41 mutants in neurons and HEK293T cells (Fig. 2 and Supplementary Fig. S3). Indeed, in HEK293T cells mutants with a deleted Coiled-coil domain (Fbxo41 ΔCC, Fbxo41 ΔF, Fbxo41 C-term and Fbxo41 ΔN-term) did not target to centrosomes, regardless of an intact F-box domain, indicating that the Coiled-coil domain is indeed necessary for Fbxo41 centrosomal targeting (Fig. 2g and Supplementary Fig. S3). However, mutants with a deleted (Fbxo41 AF-box) or mutated (Fbxo41 W577A) F-box domain were also diffusely located throughout the cytoplasm (Supplementary Fig. S3). Together, this suggests that both the Coiled-coil and the F-box domains are required for centrosome targeting. A truncation containing only the right half of the protein (Fbxo41 BF1), which includes the F-box domain but lacks the Coiled-coil domain, failed to target to centrosomes (Supplementary Fig. S3) showing that the F-box domain is not sufficient for centrosomal targeting. Unexpectedly, a mutant that lacked the F-box domain and the C-terminus (Fbxo41 ΔAF) did target to centrosomes. Finally, deletion of the zinc-finger domain (Fbxo4132Af) did not affect centrosomal targeting.

To verify Fbxo41 centrosome-targeting domains in neurons, we assessed the co-localization of several Fbxo41 mutants with pericentrin, an endogenous centrosomal protein (Fig. 2). We expressed Fbxo41 W77 (Fig. 2a), GFP (Fig. 2b), and Fbxo41 mutants (Fig. 2c,d) and quantified their centrosomal enrichment in primary hippocampal neurons (Fig. 2e). Both Fbxo41 W77, and Fbxo41 lacking the F-box domain and the C-terminus of the protein (Fbxo4132Af) were enriched at the centrosome; in contrast, Fbxo41 with a mutated F-box domain (Fbxo41 W577A) was diffusely located throughout the cytoplasm, corroborating our findings in HEK293T cells. The neuronal protein MAP2 did not accumulate at the centrosome in neurons expressing GFP or Fbxo41 constructs (Fig. 2f), indicating that Fbxo41 accumulation at the centrosome is specific.

Collectively, these results are compatible with a model in which both Fbxo41’s Coiled-coil and F-box domains are required for centrosomal targeting. The C-terminal prevents centrosomal targeting of Fbxo41, possibly through steric hindrance of the Coiled-coil domain, that can be released by the F-box domain.

**Fbxo41 promotes disassembly of primary cilia.** Given the presence of Fbxo41 at neuronal centrioles (Figs 1d and 2e), we tested whether Fbxo41 plays a role in the regulation of cilia structure. Primary neurons were infected with shRNAs or EGFP-tagged Fbxo41 constructs at day in vitro 1 (DIV1, Fig. 3a), and cilia length was measured at DIV15 (Fig. 3b) using the endogenous ciliary marker Type III adenyl cyclase (ACIII). Fbxo41 silencing was effective (Fig. 3c) but had no effect on cilia length or the percentage of ciliated neurons, indicating...
that Fbxo41 is not essential for neuronal ciliogenesis (Fig. 3d–f). However, Fbxo41WT expression in wild type neurons drastically reduced cilia length as well as the percentage of ciliated neurons (Fig. 3d–f). To discriminate whether elevated levels of Fbxo41 impair ciliogenesis or promote cilia disassembly (Fig. 3b), we also performed these experiments at a later time point. Neurons infected after ciliogenesis (DIV13) and fixed at DIV21 showed the same reduction in cilia length (Fig. 3f), suggesting Fbxo41 expression promotes cilia disassembly rather than inhibiting ciliogenesis (Fig. 3b). Similar results were obtained in human retinal pigment epithelial cells (hTERT-RPE1). Using either Arl13b or acetylated tubulin as cilia marker, we observed a significant decrease in ciliation upon expression of Fbxo41WT (Supplementary Fig. S4). Fbxo41 expression or silencing in high-density neuronal cultures did not affect total protein levels of ACIII or Arl13b, verifying that the decreased length measured with these ciliary markers was not due to a reduction in their expression levels (Fig. 3c). Taken together, using two independent model systems and three different cilia markers, we demonstrate that increased Fbxo41 expression robustly disassembles primary cilia.

Centrosomal levels of Fbxo41 correlate with cilia length. To further investigate the role of Fbxo41 in ciliary disassembly, we tested whether Fbxo41 levels at the centrosome correlate with cilia length (Fig. 3a–c). Indeed, neurons with low Fbxo41WT expression at centrosomes had normal cilia length (Fig. 3a), whilst neurons with high levels of Fbxo41WT at centrosomes generally had no visible cilia (Fig. 3b). We quantified Fbxo41WT

**Figure 1.** Fbxo41 assembles into SCF-complexes and targets to centrioles. (a) Fbxo41 assembles into SCF-complexes. HEK293T cells were transfected with the indicated constructs and subjected to immunoprecipitation with empty beads (EB), Fbxo41 or Fbxo21 antibody. Deleting (ΔF-box) or mutating (W577A) Fbxo41’s F-box domain prevented SCF-complex assembly. Fbxo21 was included as a positive control. Gel was cropped for clarity (full length blot available in Supplementary Fig. S5). (b) Fbxo41 is increasingly expressed in brain throughout development. Mouse brains were extracted at the indicated ages and immunoblotted with the indicated antibodies. Syntaxin1a was included as a positive control62. Gel was cropped for clarity (full length blot available in Supplementary Fig. S5). (c) Fbxo41 is expressed in neurons and not in astrocytes. High-density astrocyte or neuronal cultures were lysed at DIV11 and subjected to immunoblotting. βIII-tubulin was included to demonstrate the presence of primary cortical neurons. Gel was cropped for clarity (full length blot available in Supplementary Fig. S5). (d) Example of a primary hippocampal neuron fixed at DIV15 and immunostained with Fbxo41 (red), Pericentrin (green) and MAP2 (blue) antibodies. Endogenous Fbxo41 was observed in enrichments in the soma (open arrowhead). In a subset of neurons (such as this example), Fbxo41 was enriched at the pericentriolar region as indicated by Pericentrin (arrowhead). Scale bars, 5 and 2 μm. (e) Widefield fluorescence image of a primary DIV15 hippocampal rat neuron expressing Fbxo41WT-FLAG and immunostained with antibodies for FLAG (green) and Cep135 (red). Imaging was performed with FLAG-tagged Fbxo41WT due to antibody incompatibility. Scale bar, 1 μm. (f) Super-resolution dSTORM image of boxed region in (e). Scale bar, 500 nm.
Figure 2. Fbxo41 is enriched in neuronal centrioles. (a–d) Primary hippocampal neurons were infected at DIV9 with GFp or the indicated Fbxo41 mutants, and fixed at DIV14. Centrioles were marked with Pericentrin antibody (red), neurons with MAP2 antibody (blue) and EGFP-Fbxo41 mutants with GFp (green). Fbxo41WT (a), and Fbxo41ΔCΔF (d) were enriched at pericentrin-positive centrioles, whereas GFp (b), and Fbxo41W577A (c) were not. Scale bars, 5 μm. (e) Ratio of centrosomal over cytoplasmatic intensity was quantified for each Fbxo41 mutant and GFp. Fbxo41WT and Fbxo41ΔCΔF have a significantly higher enrichment at centrosomes, compared to GFp and Fbxo41W577A. (f) As a control, the ratio of centrosomal over cytoplasmatic intensity of MAP2 was quantified in the same neurons. MAP2 does not enrich in the centrosomes, indicating that the centrosomal enrichment of Fbxo41 is specific to this protein. (g) Scheme of EGFP-fused Fbxo41 mutants used for domain mapping. Colored boxes depict predicted domains: Zinc-finger (ZnF, dark blue), Coiled-coil (orange) and F-box domain (red). White asterisk represents W577A mutation. Centrosomal enrichment of the first three mutants (Fbxo41WT, Fbxo41W577A and Fbxo41ΔCΔF, inside the green box) was tested in neurons and HEK293T cells, whereas the other mutants were studied in HEK293T cells (see Supplementary Fig. S3). Data from 4 experimental weeks are represented as mean ± SEM. ****p < 0.0001 Kruskal-Wallis Test, followed by a post-hoc Dunn’s multiple comparisons test.
levels at the centrosome and detected a significant inverse correlation between cilia length and centrosomal Fbxo41WT expression (Fig. 4d). The fact that cilia length gradually declines with Fbxo41WT intensity is compatible with a model in which cilia are disassembled, instead of a binary model such as cilia shedding. This observation raised the possibility that Fbxo41 promotes cilia disassembly by sequestering, hindering or competing for proteins locally at centrioles. For this reason, we also tested the effect of Fbxo41ΔCΔF (Fig. 4c), which does target to centrioles but lacks the F-box domain essential for SCF-complex assembly (Fig. 2g). No correlation between cilia length and centrosomal expression was observed for Fbxo41ΔCΔF (Fig. 4e), despite the fact that Fbxo41ΔCΔF targets to centrosomes (Figs 2e and 4f) and showed similar expression levels compared to Fbxo41WT (Figs 2e and 4e,f). The fact that expression of Fbxo41ΔCΔF had no effect on cilium length, shows that a functional F-box domain is required for the dosage-dependent relation between centrosomal Fbxo41 levels and neuronal cilia length. To further explore the mechanism by which Fbxo41 reduces cilia length, we infected primary neurons with various Fbxo41 mutants lacking a functional F-box domain (Fig. 4g). Unlike Fbxo41WT, expression of Fbxo41ΔF-box or Fbxo41W577A had no effect on cilia length (Fig. 4g). Collectively, these results demonstrate that Fbxo41 requires a functional F-box domain for centriole targeting and cilia disassembly.

Phorbol esters, but not neuronal activity, affects neuronal cilia length. As the mechanisms that modulate cilia length in neurons remain elusive, we set out to identify physiological stimuli that can induce cilia disassembly. As such, we tested if altering neuronal activity or modulating synaptic pathways in high
Figure 4. Cilia length inversely correlates with centriolar Fbxo41 levels and requires a functional F-box domain. (a–c) Primary hippocampal neurons were infected with the indicated constructs at DIV1, fixed at DIV15, and immunostained with GFP (green), ACIII (red) and MAP2 (blue) antibodies. Fbxo41WT, but not Fbxo41ΔCΔF mutant levels at centrosome correlate with cilia length: (a) neuron with low Fbxo41WT expression (green) and long cilia (red), (b) neuron with high Fbxo41WT expression and small/no cilia, and (c) neuron with high expression of Fbxo41ΔCΔF mutant and a long cilium. The bottom panel is a pseudo color image of EGFP-Fbxo41 to emphasize the difference in Fbxo41 expression. Calibration bar illustrates color-coded signal intensity. Scale bars, 2 μm. (d) Cilia length inversely correlates with centriolar Fbxo41 levels. Scatterplot displays the relationship between Fbxo41WT intensity at the centrioles and cilia length. Each dot represents a single measurement from an individual neuron (n = 55 cells). The black line shows a linear fit through the data. Inset displays the Spearman correlation coefficient (rs) and significance (p-value). (e) Cilia length does not correlate with centriolar Fbxo41ΔCΔF levels. Scatterplot displays the lack of relationship between Fbxo41ΔCΔF intensity...
density cultures induces cilia re-modeling. The stimuli we used to modulate neuronal activity had no effect on cilia length: inhibiting post-synaptic activity by blocking NMDA and AMPA receptors (using APV and DNQX, respectively), inhibiting neuronal activity (using Tetrodotoxin) or increasing network activity (using the GABA-A receptor antagonist Gabazine) for 48 hours had no effect on cilia length (Fig. 5a). Hence, changes in neuronal network activity do not affect cilia morphology. However, treating neurons with the diacylglycerol analog PDBU, an activator of protein kinase C (PKC), a well-known modulator of synaptic plasticity26, decreased cilia length (Fig. 5a). Importantly, the PDBU-induced reduction in cilia length correlated with increased centrosomal expression of Fbxo41-EGFP (Fig. 5b,c). The effect of PDBU and Fbxo41 on cilia length were not additive (Fig. 5c), indicating that PDBU and Fbxo41 may act on a similar pathway. In order to test if PDBU is required for PDBU-dependent cilia disassembly, we tested if PDBU also reduced cilia length in the absence of Fbxo41 (Fig. 5d). DIV11 neurons treated with PDBU showed a decrease in cilia length in the presence or absence of Fbxo41 (Fig. 5d). PDBU induces cilia disassembly and increased Fbxo41 centrosomal expression in cultured neurons, but PDBU-dependent cilia disassembly is not dependent on Fbxo41.

Fbxo41-dependent cilia disassembly mechanisms differ between mitotic cells and neurons.

Cilia have been extensively studied in mitotic cells, where both Aurora A kinase and actin have emerged as key regulators of cilia structure14,38. Aurora A is a centrosomal kinase that plays a pivotal role in cilia disassembly, whereas the absence of actin polymerization increases cilia length and numbers (14,38 and Fig. 6a). We tested whether Fbxo41 promotes cilia disassembly through Aurora A kinase activation by expressing Fbxo41WT in RPE1 cells in the presence of Aurora A inhibitor PHA-680632 (Fig. 6b,c). As described above, cilia length and number were reduced by Fbxo41WT expression (Fig. 6b,c). However, in mitotic cells Fbxo41-mediated cilia disassembly was rescued by inhibiting Aurora A kinase activity (Fig. 6b,c). Drugs that perturb rearrangements of the actin-cytoskeleton also have a strong effect on cilia structure38,39, and these drugs have been shown to rescue disrupted cilia structure in several null mutants38-42. We assessed the importance of the actin-network in Fbxo41-mediated cilia disassembly by treating RPE1 cells and neurons with Cytochalasin D (CytoD), an actin filament polymerization inhibitor. In control RPE cells expressing only GFP, CytoD increased ciliary length, without affecting the percentage of ciliated cells. In Fbxo41 expressing RPE cells, CytoD rescued cilia length and percentage of ciliated cells (Fig. 6b,c). In neurons, the effect of CytoD and Aurora A Kinase on cilia length has not been studied. In contrast to its effect in mitotic cells, CytoD does not increase cilia length in neurons (Fig. 6f).

However, CytoD treatment of Fbxo41-expressing neurons rescued cilia length, but not percentage of ciliated neurons, to DMSO control neurons (Fig. 6e,f). Inhibiting Aurora A kinase failed to rescue the percentage of ciliated neurons and cilia length in Fbxo41-expressing neurons (Fig. 6e,f). Neither of the drugs did affect Fbxo41 targeting to centrosomes (Fig. 6d), indicating that the rescued cilia structure is not due to displacement of Fbxo41 from centrosomes. Moreover, these data show that centrosomal targeting of Fbxo41 occurs independently of Aurora A kinase activity and the actin-cytoskeleton. Overall, our results indicate that rearrangements of the actin-cytoskeleton are required for Fbxo41 mediated cilia disassembly, and that ciliary disassembly occurs via different mechanisms in mitotic cells and neurons.

Fbxo41-dependent cilia shortening impairs sonic hedgehog signaling. After establishing the role of Fbxo41 on cilia structure, we investigated its functional consequences, by studying Sonic hedgehog (Shh) signaling, a ciliary pathway3,4. At the basal state, the Shh receptor Ptch1 is in the cilium. Upon Shh binding, Ptch1 releases its inhibition of Smo, allowing it to target to the cilium. There Smo allows Gli proteins to go from a repressor to an activated form, promoting the expression of target genes such as Gli1 and Gli1 expression (15, and Fig. 7a). To study the effect of Fbxo41 on the Shh cascade, we expressed GFP or Fbxo41 constructs in RPE1 cells and neurons, activated the Shh cascade, and assessed Ptch1 and Gli1 expression. Fbxo41WT, but not Fbxo41WT, and Fbxo41ΔCΔF, reduced the percentage of ciliated RPE1 cells (Fig. 7b and Supplementary Fig. S4), and cilia length (Fig. 7c). As observed before, RPE1 cells were treated with conditioned media (CM) from HEK293T cells expressing N-terminal Shh (Shh-N) to activate the Shh cascade, or an empty vector (EV) as control. Shh cascade activation resulted in a significant increase in the levels of Gli1 mRNA levels in control cells and cells expressing Fbxo41WT or Fbxo41ΔCΔF, but not in cells expressing Fbxo41WT (Fig. 7d). In addition, the ratio of Gli1 expression between cells treated with Shh-N and EV was significantly lower in Fbxo41 expressing cells (Fig. 7c). To assess if the Shh cascade is also affected in neurons in the presence of Fbxo41, we activated the cascade using the Smoothened agonist SAG, as neurons did not tolerate HEK293T CM, and measured mRNA levels of Ptch1 and Gli1 (Fig. 7f). Shh pathway activation with SAG in neurons was much less efficient than Shh-N in RPE cells and Gli1 mRNA levels were not significantly upregulated upon SAG treatment in all groups (Fig. 7f). The other major downstream Shh gene Ptch1 was significantly upregulated upon SAG treatment in GFP expressing neurons, but not in those expressing Fbxo41 (Fig. 7g). Ptch1 response in neurons expressing Fbxo41WT did not reach statistical significance. Collectively, these data indicate that Fbxo41-induced cilia disassembly affects their signaling capacity in mitotic cells and, albeit to a lesser extent, in neurons.
Discussion

Although the presence of primary cilia in most neuronal cell types is now well established, and it is widely recognized that defective cilia signaling contributes to brain dysfunction, many questions about the structural regulation of these dynamic organelles remain. We identified a neuron-specific F-box protein, Fbxo41, which assembles into SCF-complexes and targets to centrosomes. Its accumulation in centrosomes promotes disassembly of neuronal primary cilia and affects their signaling capacity.

Like other F-box proteins, Fbxo41 assembles into SCF-complexes by interacting with Skp1 and Cul1 through its F-box domain (Fig. 1a and Supplementary Fig. S1). Interestingly, deleting (Fbxo41ΔF) or mutating (Fbxo41W577A) the F-box domain abolishes SCF-complex formation as well as centrosomal targeting. Skp1 and Cul1 are highly enriched at centrosomes20, but interaction with these proteins is insufficient for centrosomal targeting of Fbxo41, since truncation mutants containing an intact F-box domain but lacking the Coiled-coil domain (Fbxo41ΔCC and Fbxo41RH) do not enrich at centrosomes (Fig. 2 and Supplementary Fig. S3). Interestingly, the F-box domain is not required for centrosomal targeting in a mutant lacking the C-terminal domain (Fbxo41ΔΔF).

Therefore, our data support a mechanism whereby the C-terminal domain of Fbxo41 functions to inhibit centrosome targeting, probably through steric hindrance of the Coiled-coil domain, and the F-box domain releases this inhibition, likely through Skp1-dependent mechanisms.

Centrosomal expression of Fbxo41 robustly reduced primary cilia length (Figs 3, 4, 7 and Supplementary Fig. S4). Mutants impaired in SCF-complex formation, Fbxo41ΔF-box, Fbxo41W577A and Fbxo41ΔΔF, fail to promote cilia disassembly (Fig. 4), indicating that Fbxo41 requires engagement into- and possibly ubiquitin ligase activity of- SCF-complexes to promote cilia disassembly. Importantly, Fbxo41ΔΔF does target to centrioles, but increased centrosomal levels of Fbxo41ΔΔF do not reduce cilia length, which indicates that the effect of Fbxo41 on cilia length is not a result of build-up of protein at the centrosome but requires functional Fbxo41. Similar mechanisms have been shown for other centrosomal proteins, as overexpression of active Plk1 reduces the percentage of ciliated cells, but not overexpression of kinase dead Plk1Δ44. Similarly, overexpressed wildtype and catalytically dead Nek2 target to centrosomes, but only wildtype Nek2 reduces cilia lengthΔ45. Hence, our data shows that the engagement of Fbxo41 in SCF complexes, and not its centriolar targeting per se, affects cilia size. This is in line with reports that implicate the ubiquitin system in disassembly of ciliaΔ35,46. Although Fbxo41 expression in the brain is restricted to neurons (Fig. 1c and25), we also observed cilia disassembly in non-neuronal cells upon

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**Figure 5.** Phorbol esters, but not neuronal activity, promote ciliary disassembly. (a) In neurons at DIV 16 cilia length is unchanged by decreasing (using APV & DNQX or TTX) or increasing neuronal activity (using Gbz), but it is significantly lower in neurons treated with Phorbol ester (PDBU). Data from five independent experiments. (b,c) At DIV 16 the decrease in cilia length caused by 6 h treatment with PDBU correlates with an increase in centrosomal accumulation of Fbxo41. Data from two independent experiments. (d) Neurons treated at DIV 11 with PDBU also show a decrease in cilia length. This decrease is also observed in neurons lacking Fbxo41 (expressing shRNA against Fbxo41 since DIV1). **p < 0.01 ****p < 0.0001 Kruskal-Wallis Test, followed by a post-hoc Dunn’s multiple comparisons test. Data are represented as mean ± SEM.
overexpression of Fbxo41 (Fig. 7 and Supplementary Fig. S4), suggesting Fbxo41-mediated cilia disassembly in neurons may occur through ubiquitination of conserved cilia-associated proteins.

Recent work in RPE1 cells has revealed proteins involved in cilia disassembly. These include AuroraA, Plk1, HDAC6, Tctex1, and Nek2. None of these proteins are required for ciliogenesis but overexpression induces robust disassembly of primary cilia. We found similar phenotypes for Fbxo41 in neurons, where Fbxo41 silencing (Figs 3 and 5) or expression of dominant-negative mutants (Fig. 4g) does not disrupt ciliogenesis or ciliary maintenance, but overexpression promotes cilia disassembly. Clearly, maintaining cilium size (and hence its signaling capacity) is essential for cellular viability, which may explain why the structure of this organelle is maintained upon depletion of a single centrosomal protein.

Accumulation of catalytically-active Fbxo41 at centrioles is sufficient to disassemble neuronal primary cilia, but can ciliary disassembly in neurons occur in the absence of Fbxo41? In contrast to well-studied mechanisms of cilia disassembly in mitotic cells, mechanistic insight in cilia disassembly in post-mitotic neurons is generally lacking. We therefore tested several stimuli that could promote ciliary disassembly in neurons and observed that PKC activation by phorbol esters reduces cilia length, while modulation of neuronal network activity did not. The latter implies that neuronal cilia maintain their signaling capacity independent of network activity status. This robustness may be indicative of the importance of cilia signaling for neuronal function. In fact, to our knowledge in addition to the effects of PDBU reported here only brain ischemia results in cilia length reductions. Although there was no additive effect of PDBU on Fbxo41-mediated cilia shortening, PDBU-dependent cilia length reduction in neurons was independent of Fbxo41. Hence, although increased centriolar Fbxo41 levels leads to disassembly of cilia, Fbxo41 is not required during PDBU induced ciliary disassembly in neurons. This indicates that PDBU also triggers Fbxo41-independent pathways to reduce cilia size.

How does Fbxo41 promote disassembly of primary cilia? In dividing cells, timely disassembly of cilia is required for liberation of the centrioles and mitotic spindle formation. The molecular mechanisms underlying cilia disassembly are not completely understood but in mitotic cells involve Aurora A kinase activation.
intraflagellar transport, rearrangements of the actin cytoskeleton and/or modulation of (axonemal) microtubule stability. CytoD blocks rearrangements of the actin cytoskeleton and increases cilia length in several cell types, but not in neurons (Fig. 6). CytoD increased cilia length in RPE1 cells in both control and Fbxo41-GFP expressing cells, indicating that this mechanism is not affected by Fbxo41. In neurons, CytoD rescued Fbxo41-mediated ciliary shortening without affecting cilia length in control neurons. Together these data

**Figure 7.** Fbxo41 regulates sonic hedgehog signaling. (a) Scheme depicting sonic hedgehog cascade, which in mammals concentrates in the cilium. In the basal state, the Ptc1 receptor is in the cilium, and inhibits Smo targeting to cilium. Upon binding of sonic hedgehog (Shh) to Ptc1, Smo can target to the cilium and induces the conversion of the Gli proteins from a repressed (GliRep) to an activated (GliActv) form. Once activated, Gli proteins target to nucleus and promote transcription of Gli1 and Ptc1. (b–e) RPE1 cells were infected with the indicated constructs for three days and then fixed for quantification of percentage of ciliated cells (b) and cilia length (c), or treated with Shh-conditioned media to study sonic hedgehog cascade activation (d,e). (b) Cells expressing Fbxo41 had significantly less cilia than the cells in other conditions. Data from 2 experimental weeks. (c) Cilia length determined in the same cells as in (b), showed significantly smaller cilia in cells expressing Fbxo41WT but not in cells expressing Fbxo41WT or Fbxo41ΔCΔF. (d) Fbxo41 overexpression prevents the increase in Gli1 expression in cells stimulated with Shh-N condition media. Data from 4 or 5 experimental weeks. (e) The ratio of Gli1 mRNA levels between cells stimulated with Shh-N CM and control condition media is significantly lower in cells overexpressing Fbxo41, compared to cells expressing GFP, other Fbxo41 mutants and uninfected cells (same cells as in (d)). (f) Gli1 mRNA level increase in response to SAG treatment is not statistically significant in any condition. (g) Neurons expressing GFP but not those expressing Fbxo41, show increased Ptc1 mRNA levels in response to SAG treatment. Data are represented as mean ± SEM. *p < 0.05, **p < 0.01, ***p < 0.001 ****p < 0.0001 Kruskal-Wallis test followed by a Dunn’s multiple comparison post-hoc test.
argue for a role of actin polymerization in the mechanism of Fbxo41-dependent ciliary disassembly. Aurora A is a centrosomal kinase that plays a pivotal role in cilia disassembly during the cell cycle. Aurora A, which itself is activated through many different pathways, phosphorylates HDAC6, which in turn deacetylates and destabilizes axonemal microtubules. We found that Fbxo41 promotes cilia disassembly through activation of the Aurora A pathway in RPE cells, but not in neurons, and that Fbxo41 expression leads to a decrease in acetylated axonemal microtubules (Supplementary Fig. S4). This suggests that there are important differences between mechanisms regulating ciliary structure in mitotic cells and post-mitotic neurons. Further elucidation of these differences is pivotal for understanding ciliopathies and their effects on the brain.

Importantly, we found that Fbxo41 regulation of ciliary length impacts Shh signaling, a canonical ciliary pathway in mitotic cells. RPE1 cells and, albeit to a lesser extent, neurons respond to Shh cascade activation by increasing mRNA levels of Gli1 and Ptc1 respectively, but not in the presence of Fbxo41. This indicates that cilia length impacts ciliary signaling. In line with the notion that defective cilia signaling in neurons contributes to brain dysfunction, a recent study showed that deletion of Fbxo41 affects neuronal migration of granule neurons in the developing cerebellum. Future studies will address the question whether defective cilia signaling in the absence of Fbxo41 underlies these defects in cerebellar development.

Overall, our study sheds light on a molecular mechanism that controls primary cilia structure and signaling in neurons and highlights potential differences in cilia size modulation in mitotic and post-mitotic cells. We describe a role for Fbxo41 in this process and show that this requires rearrangements of the actin cytoskeleton in mitotic and post-mitotic cells. Primary cilia and Shh signaling have been shown to be important for crucial processes such as brain development and regulation of neural stem cell populations. By unraveling a new process through which neural cilia can be regulated, this study contributes to a better understanding of these processes, and its implications in disease.

Materials and Methods

**Laboratory animals.** Animal experiments were approved by the animal ethical committee of the VU University/VU University Medical Centre (license number: FGA 11-03) and are in accordance with Dutch governmental guidelines and regulations.

**Cell culture and transfection.** Primary neurons were prepared from embryonic day 18 mice as previously described. Hippocampi were dissected in Hanks Buffered Salt Solution (Sigma-Aldrich) buffered with 7 mM HEPES (Invitrogen) and digested with 0.25% trypsin (Invitrogen) for 20 minutes at 37 °C. Hippocampi were washed and triturated with fire-polished Pasteur pipettes, counted, and plated in Neurobasal medium (Invitrogen) supplemented with 2% B-27 (Invitrogen), 1.8% Hepes, 0.25% GlutaMAX (Invitrogen), and 0.1% penicillin-streptomycin (GE Healthcare). High-density cultures (25,000 or 600,000 neurons/well) were plated on pregrown cultures of rat glia cells (25,000 or 50,000 cells/well) on 18 mm glass coverslips or directly onto Poly-L-Ornithine coated plastic for 12-well plates (for imaging experiments) or 6-well plates (for biochemical experiments), respectively.

HEK293T and hTERT-RPE1 (gift from Rob Wolthuis, VUmc, Amsterdam) cells were cultured in DMEM (Gibco) supplemented with 10% fetal bovine serum (Gibco), penicillin-streptomycin and non-essential amino acids (Gibco). HEK293T cells were transfected at ~70% confluency in serum free DMEM using the calcium phosphate method. RPE1 cells were infected with lentivirus in serum free medium. All DNA vectors used were from mouse origin. HEK293T cells were cultured on Poly-D-Lysine (Sigma) coated 18 mm glass coverslips (for imaging experiments) or directly on plastic (for biochemical experiments).

**Immunoprecipitation experiments.** Forty hours post-transfection, HEK293T cells were washed in ice-cold PBS, harvested in Lysis buffer (50 mM Tris pH 7.5, 1% Triton X-100, 1.5 mM MgCl2, 5.0 mM EDTA, 100 mM NaCl, and phosphatase/protease inhibitor cocktails (Santa Cruz and Sigma)) and tumbled for 1 hour to ensure complete lysis. The mouse brain was collected and immediately lysed in Lysis buffer. Lysates were then centrifuged at 10,000xg for 10 minutes and pellets were discarded. Bradford assays were performed to determine protein concentrations. Antibodies were added to lysates and gently tumbled overnight. Agarose beads (Vector Laboratories) were washed twice in Lysis buffer, added to the samples, and tumbled for 1 hour. Samples were spun down at 4000xg for 5 minutes and washed in Wash buffer (50 mM Tris, 0.1% Triton X-100, 1.5 mM MgCl2, 5.0 mM EDTA, 250 mM NaCl, and phosphatase/protease inhibitor cocktails) three times. Lysates were kept on ice or at 4 °C throughout the entire experiment. Beads were boiled for 10 minutes in 2xLSB and samples were run on SDS-PAGE.

**Generation of shRNA lentivirus particles.** Fbxo41 shRNA target sequences were selected using iRNAi (Mekentosj, v2.0) and cloned into the PLL3.7 backbone with a Synapsin promoter driving EGFP expression. Oligonucleotides were designed with target sequences separated by a seven-nucleotide loop and a XhoI digestion site overhang. Fbxo41 shRNA#1:

5′-TGGCCTCTCTGTATGTTTCTGAGAGGAGCTGGCCTTTTTC

Fbxo41 shRNA#2:

5′-TAGGGCTCACTGTTAACATTCTTTGCAGAGAAATGTTAACGAGTGAGCCTGGTTTTTTC (shRNA target sequences are underlined).

**Immunocytochemistry.** Cultured cells were fixed at indicated ages with 3.7% formaldehyde in PBS, pH 7.4, for 10 minutes at room temperature. For some of the experiments with centriolar antibodies, cells were fixed with an ice-cold Methanol and Acetone solution (1:1) at −20 °C for 10 minutes. Cells were washed three times in PBS,
percentage of ciliated cells was determined by counting the total number of cells in each image, and determining subtraction. Percentage of cells with centriolar enrichment of Fbxo41 mutants was determined by visual scoring.

10% w/v glucose, 700 ng of Alexa Fluor 647 and A488 was performed by continuous oblique laser illumination with 640 nm diode laser Optovar (to achieve an effective pixel size of 64 nm) and an Andor DU-897D EMCCD camera. Sequential imaging on a Nikon Ti microscope equipped with a 100x Apo TIRF objective (NA. 1.49), a Perfect Focus System, a 2.5x 5000 and 15000 frames were recorded per acquisition with exposure times of 30 ms. Single molecule localization was performed using 5 mM MEA, and were maximally projected for analysis and display.

**Two color dSTORM imaging.** Prior to staining and imaging, DIV11 hippocampal neurons were infected for 4 to 7 days with Fbox 41^{WT}-FLAG. Neurons were fixed in MeOH for 10 minutes, permeabilized for 7 minutes using 0.25% Triton X-100 in PBS, washed and incubated for 30 minutes in blocking solution (2% BSA + 0.2% gelatin, 10 mM glycine + 50 mM NH₄Cl in PBS, pH 7.4). Neurons were incubated with primary and secondary antibodies in the blocking solution. For Fbox 41-FLAG, antibodies against Flag tag (Sigma, clone M2, 1:400 dilution) was used followed by anti-mouse coupled to A488 (Invitrogen, A-11036). For Cep 135, rabbit anti Cep135 (Sigma, clone SAB4503685, 1:250 dilution) was used followed by anti-rabbit coupled to A488 (Invitrogen, A-11034). Stained neurons were post-fixed for 10 minutes using 2% PFA in PBS. Imaging was performed using 5 mM MEA, 10% w/v glucose, 700µg/ml glucose oxidase, 40µg/ml catalase in PBS. dSTORM microscopy was performed on a Nikon Ti microscope equipped with a 100x Apo TIRF objective (NA. 1.49), a Perfect Focus System, a 2.5x Optovar (to achieve an effective pixel size of 64 nm) and an Andor DU-897D EMCCD camera. Sequential imaging of Alexa Fluor 647 and A488 was performed by continuous oblique laser illumination with 640 nm diode laser and 491 nm DPSS laser, respectively. For A488 the sample was also illuminated with 405 nm diode laser. Between 5000 and 15000 frames were recorded per acquisition with exposure times of 30 ms. Single molecule localization was performed as previously described[27]. A particle table with molecule coordinates and errors was used to reconstruct a super resolution image. 10 nm pixel size was used for super-resolution image display. For two-color imaging, chromatic corrections obtained from images with multichromatic 100 nm-beads (Tetraspeck, Invitrogen) were applied to the A488 particle table. For sample drift during acquisition, a correction algorithm was applied[28].

**Drug treatments.** For modulating neuronal activity high density neuronal cultures (40K neurons per well) were treated for 48 h with compounds known to increase or decrease neuronal activity and then fixed at DIV 16 for analysis: Gabazine (Gbz, 10µM), APV (50µM) and 6,7-dinitroquinoxaline-2,3-dione (DNQX, 20µM), tetrodotoxin (TTX, 1µM). For experiments with Phorbol 12,13-dibutyrate (PDBU; 1µM) neurons were treated for 6 hours and fixed at the indicated time points. For experiments testing the effect of AuroraA Kinase and actin in ciliary disassembly RPE1 cells were seeded in DMEM (Gibco) supplemented with 10% fetal bovine serum, and one day later the medium was replaced for DMEM without serum. After 2 days cells were infected with the indicated constructs, and 2h after treated for 18h with either DMSO, 0.5µM of PHA-680632 (PHA) or 0.5µM CytoD. Neurons were infected at DIV 9 with the indicated constructs and treated one day later with either DMSO, 0.5µM of PHA or 0.5µM CytoD, and fixed at DIV 12 for analysis. Neurons were infected for one day before treatment with PHA and CytoD.

**Shh Conditioned Media.** ShhN and EV condition media were generated as previously described[29]. Briefly, HEK293T cells grown to 70% confluency were transfected (calcium transfection using 1µg DNA, 125 mM CaCl₂ and HEPES buffered saline) with either empty vector or the N-terminus of rat Shh (gift from Lotte Bang Pedersen, University of Copenhagen). The following day cells were changed to serum free medium, and after 1 day the
medium was collected, centrifuged (5 min at 1200 rpm) and filtered (0.2μm filter). The medium was kept at −80°C and used at 1:1 for 14 to 16 h, as previously described60.

RT-qPCR. RPE1 cells were seeded in DMEM (Gibco) supplemented with 10% fetal bovine serum, and one day after the medium was replaced for DMEM without serum. After 2 days cells were infected with the indicated constructs, and 3 days afterwards treated with conditioned media for 15 hours. Neurons were infected at DIV1 with the indicated constructs and treated with 300–400 nM SAG (Enzo Lifesciences) for 24 h, as done before61. Total RNA was isolated using TRIzol reagent (Life Technologies), phase lock gel tubes and ISOLATE II RNA Mini Kit (Bioline) according to manufacturer’s instructions. The purity and quantity of RNA was assessed on a NanoDrop spectrophotometer, and RNA was reverse-transcribed into cDNA with SensiFAST™ cDNA Synthesis Kit (Bioline). The resulting cDNA was quantified using the SensiFAST™ SYBR No-ROX Kit (Bioline) in the LightCycler 480 (Roche Life Sciences). Analysis was performed using the Advanced Relative Quantification method. The cDNA was quantified in triplicates, and quantification was only considered valid when the Ct values determined did not differ for more than 1 unit among the triplicates. The relative mRNA quantity was normalized to 18S in RPE1 cells and EEF in neurons. The primers used are provided in Supplementary Table S1.

Data representation and statistics. In all graphs, data is presented as mean values ± SEM. Differences between two groups were tested for significance using a Student’s t test for unpaired data when data passed normality test (Kolmogorov-Smirnov) or a Mann-Whitney test when it did not. For multiple group comparisons, one-way ANOVA was used if allowed otherwise the non-parametric Kruskal-Wallis test was used. P-values below 0.05 are considered significant.

Data Availability All the data generated during this work is presented in the datasets. Papers are available from corresponding author upon reasonable request.

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Author Contributions
C.R.K., A.R.A.A.Q. and R.F.T. designed research, C.R.K. and A.R.A.A.Q. performed most of the imaging experiments and biochemical experiments, I.S. performed biochemical experiments and most of the cloning, A.C. performed and analyzed the STORM microscopy, A.J.G. performed imaging experiments, C.R.K. and A.R.A.A.Q. analyzed the data, and C.R.K., A.R.A.A.Q., M.V. and R.F.T. wrote the manuscript.

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