Regulation of *Chlamydomonas* flagella and ependymal cell motile cilia by ceramide-mediated translocation of GSK3

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Running head: Ceramide-mediated translocation of GSK in cilia

Abbreviations: aPKC, atypical protein kinase C; DHS, dihydrosphingosine; FB1, fumonisin B1; GSK, glycogen synthase kinase; HDAC, histone deacetylase, myr, myriocin; nSMase2, neutral sphingomyelinase 2, SPT, serine palmitoyl transferase

Supplemental Material can be found at:
http://www.molbiolcell.org/content/suppl/2015/10/05/mbc.E15-06-0371v1.DC1.html
Abstract

Cilia are important organelles formed by cell membrane protrusions, however, little is known about their regulation by membrane lipids. We characterize a novel activation mechanism for glycogen synthase kinase-3 (GSK3) by the sphingolipids phytoceramide and ceramide, which is critical for ciliogenesis in Chlamydomonas and murine ependymal cells, respectively. We show for the first time that Chlamydomonas expresses serine palmitoyl transferase (SPT), the first enzyme in (phyto)ceramide biosynthesis. Inhibition of SPT in Chlamydomonas by myriocin led to loss of flagella and reduced tubulin acetylation, which was prevented by supplementation with the precursor dihydrosphingosine. Immunocytochemistry showed that (phyto)ceramide was colocalized with phospho-tyr216-GSK3 (pYGSK3) at the base and tip of Chlamydomonas flagella and motile cilia in ependymal cells. The (phyto)ceramide distribution was consistent with that of a bifunctional ceramide analog UV-crosslinked and visualized by click-chemistry mediated fluorescent labeling. Ceramide depletion, by myriocin or neutral sphingomyelinase deficiency (fro/fro mouse), led to GSK3 dephosphorylation and defective flagella and cilia. Motile cilia were rescued and pYGSK3 localization restored by incubation of fro/fro ependymal cells with exogenous C24:1 ceramide, which directly bound to pYGSK3. Our findings suggest that (phyto)ceramide-mediated translocation of pYGSK into flagella and cilia is an evolutionarily conserved mechanism fundamental to the regulation of ciliogenesis.
Introduction

Flagella and cilia are slender organellar protrusions of the eukaryotic cell membrane with sensory and motor function. Flagella and cilia are similar in that they share a microtubule-based cytoskeleton called the axoneme (Snell et al., 2004; Sung and Leroux, 2013). Primary, non-motile cilia have a ring of nine outer microtubule doublets and are studded with growth factor receptors to function as sensory organelles regulating cell migration and differentiation. Motile cilia and flagella contain two central microtubules and the motor protein dynein attached to the nine peripheral microtubule doublets (see (Silflow and Lefebvre, 2001; Snell et al., 2004; Satir et al., 2010) for comprehensive reviews on the structure of primary and motile cilia/flagella). Motile cilia and flagella move extracellular fluids and particles, or propel cells such as sperm or unicellular organisms such as the green algae Chlamydomonas reinhardtii (Vincensini et al., 2011). Flagella dysfunction leads to reduced or absent motility, while aberrant motile cilia lead to ependymal cell malfunction and hydrocephalus (Young et al., 2013). It is vital for the signaling and motility functions of cilia that length, number, and intraciliary or intraflagellar transport (IFT) of cargo are dynamically regulated. The protein machinery regulating assembly, length, movement, and transport within these organelles is conserved from Chlamydomonas to mammals (Snell et al., 2004; Vincensini et al., 2011; Sung and Leroux, 2013). Although cilia and flagella are localized extensions of the plasma membrane, little is known about the role of lipids in this regulation of ciliogenesis and cilium function.

Sphingolipids are essential components of eukaryotic membranes and regulate vital cell signaling pathways (Bartke and Hannun, 2009; Milhas et al., 2010; Bieberich, 2012; Markham et al., 2013). The common long chain base precursor for de novo biosynthesis of animal, plant, fungus, and algae sphingolipids is dihydrosphingosine (DHS, Figure 1). DHS is converted to
ceramide (animals) and phytoceramide (plants, algae, and yeast). In animals, ceramide and its metabolite, sphingomyelin are unique in that their levels can be rapidly regulated by enzymatic conversion, the so-called sphingomyelin cycle (Figure 1) (Hannun, 1996). Ceramide is converted to sphingomyelin by sphingomyelin synthases, while sphingomyelin is hydrolyzed to ceramide by acid or neutral sphingomyelinases (aSMase, nSMase) (Nikolova-Karakashian et al., 2008; Milhas et al., 2010; Shamseddine et al., 2015). We previously reported that aSMase and nSMase-mediated generation of ceramide is critical for ciliogenesis in MDCK cells and neural progenitors, respectively (Wang et al., 2009a; He et al., 2012; He et al., 2014). In contrast to mammalian cells, Chlamydomonas and plants do not synthesize sphingomyelin, which therefore is excluded as a source for ceramide. However, in all eukaryotic cells, ceramides can be generated by de novo biosynthesis (Figure 1) (Sperling and Heinz, 2003; Gault et al., 2010; Pata et al., 2010; Hannun and Obeid, 2011; Markham et al., 2013). The first enzyme in this pathway, serine palmitoyltransferase (SPT) can be selectively inhibited by the fungal toxin myriocin, allowing for loss-of-function experiments to understand the significance of ceramide for ciliogenesis in animal cells and algae (Miyake et al., 1995; Hanada et al., 2000; Wadsworth et al., 2013). The long chain base in algae and plant sphingolipids is more complex than that in animals. Hydroxylation of DHS at C4 results in biosynthesis of phytosphingosine, which can be acylated to give rise to phytoceramides (Figure 1) (Spassieva et al., 2002; Sperling and Heinz, 2003; Pata et al., 2010; Markham et al., 2013). While plants and algae sphingolipids are relatively well known, the sphingolipid pathway in Chlamydomonas has not been investigated.

Physical interaction of (phyto)ceramide with proteins that are critical for ciliogenesis is likely to underlie the regulation of flagella and motile cilia, respectively. Previously, we found that in MDCK cells and human ES cell-derived neuroprogenitors, atypical PKCζ (aPKCζ)
directly interacts with ceramide and promotes primary cilium extension ultimately through inhibition of histone deacetylase 6 (HDAC6) (Bieberich et al., 2000; Wang et al., 2005; Wang et al., 2009b; He et al., 2012; He et al., 2014). HDAC6 inhibition prevents deacetylation of lysine 40 of α-tubulin, a reaction that destabilizes microtubules and promotes cilium disassembly (Pugacheva et al., 2007; Loktev et al., 2008). A Chlamydomonas tubulin deacetylase has been described, but it is not clear whether it has the same function for flagella regulation as HDAC6 for cilia in animals (Maruta et al., 1986). It is likely that ceramide interacts with additional proteins to regulate HDAC6 and/or cilium length. One candidate protein, conserved in Chlamydomonas and mammalian cilia is GSK-3β (Wilson and Lefebvre, 2004; Thoma et al., 2007). GSK-3β (GSK3) is critical for ciliogenesis, regulates HDAC6, and phosphorylates kinesin light chains that control cargo binding to microtubules (Szebenyi et al., 2002; Thoma et al., 2007; Song et al., 2014).

To identify ceramide-protein complexes in cilia, we used chemoproteomics technology for in vitro and in vivo UV cross-linking of a bifunctional ceramide analog (pacFACer) to interacting proteins and functionalizing this covalent complex for visualization using “Click chemistry” (azide-alkyne cycloaddition, Figure S1 for structure). In addition, we have developed a novel anti-ceramide antibody that recognizes ceramide and phytoceramide. Using pacFACer cross-linking and anti-ceramide antibody, we demonstrate that Chlamydomonas flagella and ependymal cell cilia show a similar distribution of ceramides that bind to GSK3, which is critical for the regulation of ciliogenesis by an evolutionarily conserved mechanism of ceramide-mediated translocation of GSK3 into flagella and cilia.
Results

*Chlamydomonas* expresses serine palmitoyltransferase and generates plant sphingolipids

There is no prior information on the presence of sphingolipids or enzymes required for sphingolipid biosynthesis in *Chlamydomonas*. Using reverse transcriptase (RT)-PCR and oligonucleotide primers based on sequence similarities between mouse serine palmitoyltransferase (SPT), the first enzyme in the sphingolipid biosynthetic pathway, and *Chlamydomonas* expressed sequence tagged (EST) clones, we found that *Chlamydomonas* expresses mRNAs for the subunits SPT1 and SPT2 (Figure 2A). mRNAs for these SPT subunits are similar to those expressed in mouse ependymal cells enriched from primary cultured glial cells (Weibel *et al.*, 1986). In addition to SPT, we also detected mRNAs for lag1 and lag2, two ceramide synthases expressed in plants, suggesting that *Chlamydomonas* expresses several enzymes in the phytoceramide biosynthesis pathway (Figure S1B). Expression of SPT was confirmed by immunoblot. Using an antibody raised against mammalian SPTs, we were able to detect SPT2 at the predicted molecular weight (72 kD) but not SPT1 in *Chlamydomonas* (Figure 2B). The expression of SPTs and ceramide synthases in *Chlamydomonas* suggested that sphingolipids are synthesized. Using LC-MS/MS, we found that *Chlamydomonas* generates very long chain C26:0 (11±3 pmol/mg cells) and C28:0 (8±2 pmol/mg cells) phytoceramides, which are common plant sphingolipids (Figure 2C). In comparison, glial cells contain relatively small proportions of phytoceramide (<10% of total ceramides) (Dasgupta *et al.*, 2013) but larger amounts of ceramide (800±150 pmol/mg cells), with C18:0 ceramide (35%) and C24:1 ceramide (25%) being the most abundant (Wang *et al.*, 2012). Based on the observation that SPT2 is expressed in *Chlamydomonas* as well as mammalian cells, we tested the effect of myriocin, a fungal toxin and highly selective inhibitor of SPT by covalent binding to SPT2 (Wadsworth *et
al., 2013), on (phyto)ceramide biosynthesis. Figure 2D shows that myriocin is an extremely potent inhibitor of phytoceramide biosynthesis (>90% at 5 nM) in *Chlamydomonas*, suggesting that myriocin can be effectively used to test the function of phytoceramide in *Chlamydomonas*.

**Phytoceramide and ceramide are critical for motility and ciliary length regulation**

Based on our previous studies showing that ceramide is critical for primary ciliogenesis (Wang *et al.*, 2009a; He *et al.*, 2012; He *et al.*, 2014), we tested if a similar function exists for the regulation of flagella in *Chlamydomonas* and motile cilia in ependymal cells. To determine the effect of inhibition of *de novo* sphingolipid biosynthesis by myriocin on flagella, we first performed a phototaxis motility assay (Lechtreck *et al.*, 2009). Following incubation of *Chlamydomonas* in a 6-well dish with 0-50 nM myriocin, half of each well was shielded from light. At 7 hours, most of the algae were still motile and swimming towards light. By 15 hours, only the algae not exposed to myriocin were fully motile (Figure 3A). This was confirmed both by cell counting with a hemocytometer and by measuring optical absorbance of chlorophyll at 450 nm. At a concentration of myriocin as low as 5 nM, approximately half of the cells were immotile (Figure 3B). At 10 nM myriocin, phase contrast microscopy showed that >70% of cells had extremely shortened (<2 µm length) flagella (Figure 3C) and often formed multicellular clusters (Figure S2A). Only a small population of cells (<20%) did not show significant flagella shortening. The inhibitory effect of myriocin on cilium length and motility was reversible since myriocin-treated cells regained motility within 48 h when resuspended in myriocin-free medium (Figure S2B). This result was consistent with a trypan blue exclusion assay showing that myriocin-treated cells did not undergo cell death (Figure S2C), further suggesting that the effect of myriocin on motility was specific for ciliogenesis and not due to general toxicity. To test if
the effect of myriocin was caused by reduced synthesis of sphingolipids, we performed motility
rescue experiments by supplementing the medium with various sphingolipid precursors 2 h prior
to myriocin exposure. Among the sphingolipids tested, dihydrosphingosine (DHS), a metabolic
precursor for (phyto)ceramide biosynthesis (Figure 1), and phytoceramide itself were the most
effective (60±10% and 42±12%, respectively) in rescuing flagella length and motility (Figure 3D
and E). Sphingosine and various ceramides were not effective for restoration of cilia (not
shown). Fumonisin B1, a ceramide synthase inhibitor, failed to reduce motility in
Chlamydomonas. This inhibitor has a higher dissociation constant than myriocin (100 nM vs.
0.28 nM) and in the case of lower uptake, the intracellular concentration of fumonisin B1 may
not allow for effective inhibition of phytoceramide biosynthesis (Wang et al., 1991; Miyake et
al., 1995). In ependymal cells, both myriocin and fumonisin B1 (FB1; Figure 1) reduced the
length of motile cilia (Figure 3G). Addition of C24:1 ceramide to the culture medium rescued
cilia in cells treated with FB1 and induced elongation of cilia in cells not treated with ceramide
biosynthesis inhibitors (Figure 3F and G). In addition to FB1-treated wild type ependymal cells,
we used cells from the fragilitis ossium (fro/fro) mouse, a mouse model for genetic deficiency of
nSMase2 (Guenet et al., 1981; Poirier et al., 2012). In these cells, lack of nSMase2-catalyzed
ceramide generation led to shortened cilia, which was rescued by exogenous C24:1ceramide
(Figure S3). Other ceramides such as C18:0 ceramide were not effective in rescuing cilia in
ependymal cells (not shown). These results suggest that phytoceramides in Chlamydomonas and
C24:1 ceramide in ependymal cells are critical for cillum length regulation, which is likely to
rely on a conserved downstream mechanism.
Phytoceramide and ceramide are enriched in compartments that regulate ciliogenesis

Any effect of (phyto)ceramide on cilium regulation is likely to require the physical interaction of (phyto)ceramide with proteins in compartments that are known to regulate cilia, in particular vesicles and membranous compartments at the cilium base and the ciliary membrane. We previously generated a ceramide-specific antibody that detected an apical ceramide-enriched compartment (ACEC) at the base of primary cilia (Wang et al., 2009a; Bieberich, 2011; He et al., 2012; He et al., 2014). To test (phyto)ceramide distribution in *Chlamydomonas* and ependymal cells, we have generated a new antibody that reacts with both ceramide and phytoceramide as shown by lipid ELISAs (Figure 4A; data shown are background subtracted; (He et al., 2014; Dinkins et al., 2015). This antibody does not react with fatty acid (nervonic acid was tested), but it recognizes sphingosine and phytosphingosine, which are only minor cellular sphingolipids (≤10% of (phyto)ceramide) under physiological conditions. Immunocytochemistry using this antibody showed punctate labeling for (phyto)ceramide at tips of flagella and motile cilia (Figure 4B). In addition, (phyto)ceramide was enriched in vesicles or a compartment at the bases of flagella and motile cilia (Figure 4B).

We next tested if this compartment supplies (phyto)ceramide for the ciliary membrane. We used *Chlamydomonas* because it can be subjected to several cycles of de- and reflagellation where functionality is noted by return of motility (Wilson and Lefebvre, 2004). *Chlamydomonas* was 4-times deflagellated at pH 4.5 followed by rapid neutralization and reflagellation in the absence or presence of myriocin. With each de/reflagellation cycle, fewer cells retained motility. After 4 cycles, myriocin-treated cells completely lost motility while control cells were not affected (Figure 4C). In contrast to overnight incubation (Figure 3A), continuous myriocin incubation for 5 h of cells not undergoing de/reflagellation did not show loss of motility.
Immunocytochemistry using the anti-(phyto)ceramide antibody showed that loss of motility in myriocin-treated cells undergoing repeated de/reflagellation was concurrent with the inability to regrow flagella and disappearance of the phytoceramide-enriched compartment at the flagella base (Figure 4D). This result suggests that the phytoceramide-enriched compartment in the cell body is dependent on de novo phytoceramide biosynthesis and supplies phytoceramide for the flagellar membrane. In the presence of myriocin, phytoceramide transported into the flagellar membrane and lost by repeated deflagellation can no longer be replenished from the cell body and flagella are no longer formed.

To test if phytoceramide is cotransported with proteins from the base into flagella, we used a bifunctional ceramide analog pacFA Ceramide (pacFACer; Figure S1A) to covalently label protein interaction partners of phytoceramide in the compartment at the flagella base and the flagellar membrane. After UV cross-linking, the pacFACer-protein complex was visualized by covalent linking to Alexa Fluor 594 azide using click chemistry. Alexa Fluor 594-labeled pacFACer was enriched in the base and tip of flagella (Figure 4E), consistent with the results obtained with anti-(phyto)ceramide antibody (Figure 4B). Next, we incubated *Chlamydomonas* with pacFACer followed by UV cross-linking and one cycle of de/reflagellation in the presence of myriocin before labeling of the pacFACer-protein complex with Alexa Fluor 594 azide. A proportion of the cells (15±5%) formed single very long (20±3 µm) flagella approximately double the length of normal flagella, which was not observed in the absence of UV cross-linking (Figure 4F). In these cells, the ceramide-enriched compartment at the flagella base was absent, consistent with the results obtained from repeated de/reflagellation (Figure 4D). The formation of a single, elongated flagellum labeled with pacFACer suggests that the pacFACer-protein complex is cross-linked at the flagella base and is then transported into the newly formed
flagellum. Failure to dissociate after initial binding and transport due to covalent cross-linking leads to irreversible membrane association and excess cilium extension, similar to the phenotype of *Chlamydomonas* mutants with defective protein kinases involved in cilium length regulation (Berman *et al.*, 2003; Wilson and Lefebvre, 2004; Hilton *et al.*, 2013). These results suggest that (phyto)ceramide at the cilium base transiently interacts with proteins, in particular protein kinases that are critical for cilium length regulation.

**Motile ciliogenesis in Chlamydomonas and ependymal cells relies on active GSK-3β**

One of the protein kinases that is highly conserved from *Chlamydomonas* to mammals and critical for ciliogenesis is GSK-3β. In *Chlamydomonas*, whose genome encodes for a single GSK3 isoform, activated pYGSK3 has been shown to be enriched in flagella and to regulate flagellar length (Wilson and Lefebvre, 2004). We performed immunoblotting of proteins from myriocin-treated *Chlamydomonas* and found a reduction of pYGSK3 levels, which was prevented by exogenous DHS, the phytoceramide precursor (Figure 5A). Consistently, acetylated tubulin was also reduced by myriocin and rescued by DHS, supporting the hypothesis that pYGSK3 levels are correlated with tubulin acetylation in flagella. In addition to pYGSK3, total GSK3 levels were reduced by myriocin and rescued by DHS, suggesting that ceramide depletion leads to decreased expression or increased degradation of *Chlamydomonas* GSK3.

To show the direct effect of GSK3 activity on flagella and cilia we incubated *Chlamydomonas* and ependymal cells with BIO ((2′Z,3′E)-6-bromoindirubin-3′-oxime), a highly selective GSK3 inhibitor (Sato *et al.*, 2004), for 60 min. GSK3 inhibition led to immotility and flagellar length reduction (Figure 5B) similar to that observed with myriocin (Figure 3C), which is consistent with the hypothesis that phytoceramide and ceramide are critical for activation of
GSK3 to regulate cilium length. Lithium chloride (LiCl), a non-competitive GSK3 inhibitor also led to immotility, although flagella were first elongated (after 2h incubation, Figure 5B) and then lost (after 24 h, not shown) consistent with previously published studies (Wilson and Lefebvre, 2004).

The enrichment of pYGSK3 in flagella indicates that it may be colocalized with phytoceramide and ceramide in *Chlamydomonas* and ependymal cells, respectively. We found that pYGSK3 was colocalized with phytoceramide in the flagellar membrane, particularly in the flagella tips (Figure 5C). In ependymal cells, cilia were colabeled for pYGSK3 and ceramide (Figure 5D). Interestingly, we found that while pYGSK3 colocalized with acetylated tubulin in cilia (Figure 5E, left panel) inactive GSK3 phosphorylated at ser9 (pSGSK) was not found in cilia (Figure 5E, right panel). This result suggested that pYGSK3 may bind directly to ceramide in the ciliary membrane. A lipid ELISA-based binding assay using recombinant human GSK-3β expressed in Sf9 cells confirmed that GSK3 binds to different ceramide species (Figure 5F). Affinity of human GSK3 was highest to C24:1 ceramide, which was able to rescue cilia in ceramide-depleted ependymal cells (Figure 3F and G). Immunoblots showed that human GSK3 was phosphorylated at tyr216, consistent with the hypothesis that pYGSK3 binds directly to ceramide (Figure 5G).

Immunoblots of ependymal cells from wild type and *fro/fro* mice showed reduced pYGSK3 and increased pSGSK3 in *fro/fro* cells, suggesting that reduced cellular ceramide leads to less activation of GSK3 (Figure 6A), concurrent with reduced cilium length and diminished labeling of pYGSK3 in motile cilia (Figure 6B and Figure S3). Immunohistochemistry on cryosections of *fro/fro* brains showed that the length of motile cilia in the ependyma was also reduced by more than 50% (Figure 6C and D). Motile cilia in the ependyma showed colocalization of ceramide
with pYGSK3, predominantly in punctate structures along the membrane and cilium tip (Figure 6E and F), which was consistent with the results obtained with primary cultured ependymal cells (Figure 6B and C). Taken together, these results suggest that phytoceramide and ceramide may induce activation and translocation of pYGSK3 into flagella and cilia, which is instrumental for the regulation of their length.

**Ceramide interaction with aPKCζ regulates ciliogenesis in ependymal cells**

Our previous studies showed that atypical PKCζ (aPKCζ), another protein kinase that directly binds to ceramide regulates primary ciliogenesis in mammalian cells (Lozano et al., 1994; Muller et al., 1995; Wang et al., 1999; Bieberich et al., 2000; Bourbon et al., 2000; Fan et al., 2004; Ossipova et al., 2007; Pruliere et al., 2011; He et al., 2012; He et al., 2014). GSK-3β can be inactivated by aPKCζ-mediated phosphorylation of Ser9 (pSGSK3), suggesting that mammalian ciliogenesis can be regulated by binding of ceramide to both, GSK-3β and aPKCζ (Etienne-Manneville and Hall, 2003; Kim et al., 2007; Krishnamurthy et al., 2007b). Genome analysis shows that plants and *Chlamydomonas* do not express a homolog of aPKCζ and GSK3 does not contain an equivalent to the Ser9 phosphorylation site found in mammalian GSK-3β (Kruse et al., 1996; Wilson and Lefebvre, 2004). Therefore, we limited the analysis motile cilia regulation by ceramide and aPKCζ to ependymal cells. An orthogonal view of a confocal z-scan shows that UV cross-linked pacFACer colocalizes with aPKCζ in the ceramide-enriched compartment at the cilium base (Figure 7A). The physical interaction of pacFACer with aPKCζ was demonstrated by a lipid ELISA-based assay using a surface coat of pacFACer for binding to recombinant aPKCζ. Binding was enhanced by UV cross-linking, suggesting that aPKCζ could be covalently linked to pacFACer and is indicative of high affinity for ceramide. The reaction did
not occur in the absence of pacFACer (pCer), aPKCζ, or aPKCζ-specific antibody (Figure 7B). The ceramide binding affinity of aPKCζ was tested using lipid-ELISA with surface coats of ceramides differing in the chain length and degree of saturation of the fatty acid moiety. Affinity of aPKCζ was higher for ceramides with very long chain and unsaturated fatty acids, with highest affinity for C24:1 ceramide (Figure 7C), similar to the affinity found with GSK-3β (Fig. 5F). This result is consistent with rescue of cilia length in myriocin-treated ependymal cells and elongation of cilia in untreated cells by C24:1 ceramide.

We tested whether aPKCζ activation or inhibition is critical for ceramide-regulated motile ciliogenesis. Incubation of ependymal cells with fumonisin B1 (FB1), a ceramide synthase inhibitor, led to reduction of motile cilium length (Figure 7D). The cell-permeable myristoylated aPKCζ pseudosubstrate inhibitor (PZI) prevented reduction of motile cilium length by FB1, suggesting that ceramide-mediated inhibition of aPKCζ is critical for motile ciliogenesis (Figure 7D). Consistent with this hypothesis, addition of PZI to cells not treated with FB1 resulted in increase of cilium length by 50-60% (Figure 7D). Taken together, these results suggest that in ependymal cells, ceramide binds and inhibits aPKCζ, which leads to increase of cilium length. However, since co-localization of aPKCζ with ceramide was only detected at the cilium base, but not in the ciliary membrane (Figure 7A), we concluded that binding of aPKCζ to ceramide at the cilium base affects a secondary target that regulates cilium length by its translocation into cilia. Consistent with the observation that GSK3 is critical for flagella formation and a target for mammalian aPKCζ, we found that inhibition of GSK3 with BIO obliterated motile cilia in ependymal cells (Figure 7D). Therefore, we hypothesize that GSK3 is this secondary target by being phosphorylated at Ser9 and inactivated by aPKCζ, unless aPKCζ is bound to ceramide at the cilium base and sequestered from GSK3 that is translocated to the
ciliary membrane.

**Discussion**

Length regulation of primary and motile cilia and flagella is essential for their biological function. To date, there are more than 20 proteins known to regulate cilium length, among which many are protein kinases (e.g., Aurora A kinase (AurA), MAPK/MAK/MRK overlapping kinase or MOK/RAGE1), IFT proteins, and enzymes involved in tubulin modification (e.g., HDAC6) (Pan *et al.*, 2004; Pugacheva *et al.*, 2007; Cao *et al.*, 2013; Broekhuis *et al.*, 2014). Little is known about the dynamic role of lipids in ciliogenesis. Since the cell membrane cannot be stretched any cilium length extension is inevitably accompanied by net synthesis and transport of membrane lipids, by either vesicle traffic or lateral movement of lipids within the cell membrane toward the cilium base. While new membrane is added at the cilium base, tubulin and other proteins for cilium extension are transported along the cilium and then added at the tip. It is not clear how membrane lipid and protein transport are coregulated to ensure that elongation of the axoneme is adjusted to the adequate expansion of the ciliary membrane. Our study presents new evidence that the sphingolipid (phyto)ceramide is instrumental for length regulation of flagella and cilia based on data from the green algae *Chlamydomonas* flagella and murine ependymal cells.

*Chlamydomonas* is a well-established model to study regulation of motile cilium length by testing mechanisms that affect flagella growth and regeneration. It has been shown that the flagellar membrane is enriched with phosphatidylethanolamine, which can be hydrolyzed by phospholipase D (PLD) to phosphatidic acid (PA), a lipid known to induce deflagellation (Quarmby *et al.*, 1992; Goedhart and Gadella, 2004; Lechtreck *et al.*, 2013). However, it is not
clear if PLD-mediated generation of PA is part of a process regulating flagella length. It is also
not known whether *Chlamydomonas* expresses enzymes that generate sphingolipids and whether
these lipids, if present, are essential for regulation of flagella. Using RT-PCR and
immunoblotting we showed that SPT, the first enzyme in de novo sphingolipid biosynthesis, in
particular its subunit SPT2, is conserved and expressed in *Chlamydomonas*. LC-MS/MS
confirmed that C26:0 and C28:0 phytoceramide are generated, suggesting that *Chlamydomonas*
expresses a complete set of enzymes for de novo sphingolipid biosynthesis. While the analysis of
the complete set of enzymes in (phyto)ceramide biosynthesis is part of our future studies, we
could show that *Chlamydomonas* expresses the mRNAs of two putative ceramide synthases, lag1
and lag2. Among the enzyme inhibitors tested, the SPT inhibitor myriocin blocked
phytoceramide biosynthesis and led to flagella length reduction and immotility. The specificity
of this effect was demonstrated by the recovery of flagella growth and motility after removal of
myriocin and by rescuing flagella with exogenous DHS, a downstream product of SPT and
precursor for phytoceramide. Flagella were also partially rescued by exogenous phytoceramide,
suggesting that flagella length is regulated by DHS that is metabolically converted to
phytoceramide. Exogenous ceramide did not rescue flagella in *Chlamydomonas*, but it prevented
loss of motile cilia in ependymal cells treated with inhibitors of ceramide biosynthesis. These
results demonstrate that phytoceramide and ceramide are ciliogenic sphingolipids that regulate
cilium length in *Chlamydomonas* and ependymal cells, respectively.

In animals, ceramide is synthesized in the endoplasmic reticulum (ER), transported to the
Golgi, where it can be converted into glycosphingolipids or sphingomyelin, which are further
transported to the cell membrane. Sphingomyelin can be internalized by endocytosis and
hydrolyzed to ceramide by sphingomyelinases (SMases). Our previous studies have shown that
SMases in mammalian cells generate ciliogenic ceramide that is accumulated in an apical ceramide-enriched compartment (ACEC) at the base of primary cilia (Wang et al., 2009a; He et al., 2012; He et al., 2014). In plants and yeast, phytoceramide is also synthesized in the ER and derivatized to complex phytosphingolipids in the Golgi. However, plants and algae do not contain sphingomyelin, but instead they generate inositolphosphorylsphingolipids (Lester and Dickson, 1993; Sperling and Heinz, 2003; Markham et al., 2013). Therefore, phytoceramide in *Chlamydomonas* is likely to be generated at the flagella base, either by *de novo* biosynthesis or degradation of complex phytosphingolipids, and then further transported into the ciliary membrane. This transport was tested using two techniques: labeling of phytoceramide and ceramide with an antibody and direct fluorescent labeling of the bifunctional ceramide analog pacFACer after UV cross-linking to interacting protein(s). Anti-ceramide IgG was originally generated in our laboratory and has been extensively tested in independent laboratories using immunocytochemistry (Krishnamurthy et al., 2007a; Wang et al., 2009a; Muscoli et al., 2010; He et al., 2012; He et al., 2014). Anti-ceramide IgG was found to detect ceramide and phytoceramide in lipid ELISAs (He et al., 2014; Dinkins et al., 2015). Using this antibody, we showed that phytoceramide and ceramide are enriched at the base and the tip of flagella and motile cilia in *Chlamydomonas* and ependymal cells, respectively. These results suggest that (phyto)ceramide is transported from the ciliary base to the tip.

Figure 8A shows a model consistent with a by co-transport of (phyto)ceramide and flagella/cilia length-regulating proteins. In this “flux equilibrium model”, lipid vesicles from the ceramide compartment are incorporated into the ciliary membrane at the cilium base. Lipid-cargo protein co-transport ensures the stoichiometry required for simultaneous cilium elongation and membrane expansion. (Phyto)ceramide may serve as a membrane anchor for co-transported
cargo proteins, or alternatively, it may activate a loading/unloading mechanism at the base or tip of the cilium. The cilium length is regulated by the size or ceramide content of the compartment at the base and its lipid flux to the cilium: more ceramide favors cilium assembly, less favors disassembly until flux rates in both directions are equal and the cilium length is maintained. Consistent with this model, repeated de- and reflagellation in the presence of myriocin led to the inability to regenerate flagella, indicating that phytoceramide is continuously incorporated into the ciliary membrane and replenished by de novo biosynthesis. Appearance of fluorescently labeled pacFACer in newly formed flagella was concurrent with disappearance of labeling at the base, further supporting the hypothesis that (phyto)ceramide is transported from the base into the ciliary membrane.

Any specific regulation of cilia by (phyto)ceramide requires its interaction with proteins involved in ciliogenesis. It has been shown that several evolutionary conserved kinases with mammalian homologs such as *Chlamydomonas* AurA-like kinase (CALK), LF4p (MOK in mammals), and GSK3 are important for flagella length regulation (Silflow and Lefebvre, 2001; Berman et al., 2003; Pan et al., 2004; Wilson and Lefebvre, 2004; Thoma et al., 2007; Cao et al., 2009; Ou et al., 2009; Cao et al., 2013; Hilton et al., 2013; Broekhuis et al., 2014). GSK3 is of particular importance since its role in ciliogenesis has been confirmed in numerous studies, but it is unclear if cilium length extension is induced by activation or inhibition of GSK3, or both. Lithium chloride (LiCl), a non-competitive inhibitor of GSK3, has been shown to induce cilium extension in *Chlamydomonas* and mammalian cells (Berman et al., 2003; Nakakura et al., 2014). However, LiCl is not specific and requires mM concentrations for effectiveness, and other studies have shown that its effect on ciliogenesis is independent of GSK3 (Ou et al., 2009). Specific GSK3 inhibitors have not been tested in *Chlamydomonas*, and they have given
inconsistent results when used to test their effect on mammalian ciliogenesis (Ou et al., 2009; Wang et al., 2009a). We administered two specific GSK3 inhibitors, BIO and indirubin-3-monoxime, to Chlamydomonas and found that they lead to complete shortening of flagella within 60 min, consistent with their effect on ependymal cells. Therefore, we conclude that active GSK3 is critical for maintenance or extension of cilia. This conclusion is consistent with previous studies showing that active pYGSK3 is translocated into flagella and that GSK3 knockdown leads to flagella shortening (Wilson and Lefebvre, 2004). It is also consistent with studies showing that specific inhibition or knockdown of GSK3 impairs ciliogenesis in mammalian cells (Ou et al., 2009). The critical role of pYGSK3 for ciliogenesis was further substantiated by our observation that pYGSK3 is colocalized with phytoceramide and ceramide in Chlamydomonas flagella and ependymal cell motile cilia, respectively. In ependymal cells, nSMase2 deficiency led to decreased levels of pYGSK and increased levels of pSGSK, clearly showing that nSMase2-mediated generation of ceramide is instrumental for increasing pYGSK3 and reducing pSGSK levels.

Activation of GSK3 by autophosphorylation of tyr216 is a cotranslational process, while inactivation by desphosphorylation at this residue and phosphorylation of ser9 is post-translational and regulated by phosphatases and kinases, respectively (Beurel et al., 2015). Since nSMase2-deficient ependymal cells show reduction of tyr216 and increase of ser9 phosphorylation, ceramide is likely to affect phosphatases and kinases that regulate GSK3 phosphorylation. We and other laboratories have found that ceramide binds and sequesters aPKCζ (Lozano et al., 1994; Muller et al., 1995; Wang et al., 1999; Bieberich et al., 2000; Bourbon et al., 2000; Wang et al., 2005; Fox et al., 2007; Wang et al., 2009b), which was confirmed by lipid-ELISAs and UV cross-linking to pacFACer. Ceramide-induced sequestration
may prevent aPKCζ-mediated phosphorylation of GSK3 at ser9 and lead to enrichment of pYGSK3 in the cillum tip (Figure 8B). The hypothesis that ceramide-mediated sequestration of aPKCζ is critical for cilium length extension is supported by the observation that inhibition of aPKCζ with PZI promotes ciliogenesis in ependymal cells. Ceramide depletion prevents sequestration of aPKCζ and enrichment of pYGSK3 in cilia, while it promotes phosphorylation of GSK3 at ser9 and elevation of pSGSK3 levels (Figure 6A). Neither aPKCζ nor pSGSK3 are transported into cilia, suggesting that ciliary pYGSK3 is critical for cilium length regulation.

In *Chlamydomonas*, pYGSK3 levels and transport into flagella are elevated by phytoceramide, but the interaction partner is currently unknown as plants and algae do not possess aPKC homologs (Kruse et al., 1996). Moreover, *Chlamydomonas* GSK3 does not have a phosphorylation site homologous to ser9 in mammalian GSK-3β (Wilson and Lefebvre, 2004). Therefore, phytoceramide may sustain pYGSK3 levels and its enrichment into flagella by a different mechanism. Our data showing that phytoceramide is colocalized with pYGSK3 in flagella strongly suggests that phytoceramide may regulate IFT of pYGSK3 (Figure 8B). Likewise, colocalization of ceramide with pYGSK3 in motile cilia suggests that ceramide may interact directly with pYGSK3 within the ciliary membrane and promote its transport to the tip. Support for this hypothesis comes from lipid-based ELISAs showing that recombinant human GSK-3β binds to C24:1 ceramide, the ceramide species rescuing motile cilia in ceramide-depleted ependymal cells. Potential binding to pYGSK3 may be the evolutionarily older mechanism by which (phyto)ceramide regulates flagella/cilia, while ceramide-mediated sequestration of aPKC may have developed more recently as complementary mechanism of GSK3 activation in animals.

It is currently unknown how the interaction of phytoceramide or ceramide with pYGSK3 in
the cilium tip may regulate cilium length extension. It is possible that pYGSK downregulates HDAC6, thereby preserving tubulin acetylation (Figure 8B). Support for this hypothesis comes from a previous study showing that dysfunctional GSK3 leads to upregulation of HDAC6 and ciliary defects (Song et al., 2014). It is also possible that pYGSK does not directly regulate tubulin acetylation but instead cargo release at the cilium tip. It has been shown that GSK3-mediated phosphorylation of kinesin light chain leads to release of transport vesicles in neuronal axons (Szebenyi et al., 2002). It is likely that a similar mechanism may induce phosphorylation of kinesin 2 in cilia, which then allows for release of proteins that are critical for cilium assembly at the tip. The interaction of phytoceramide and ceramide with pYGSK at the cilium tip may trigger this cargo release and promote ciliogenesis.

In summary, our study shows that (phyto)ceramide is critical for the regulation of flagella in *Chlamydomonas* and motile cilia in ependymal cells *in vitro* and *in vivo*. We present new evidence that nSMase2-deficiency results in shortening of motile cilia in the ependyma, suggesting that reduction of ceramide generation leads to ciliary dysregulation *in vivo*. While it has not been described that mutations of enzymes in ceramide metabolism are linked to ciliopathies, our study shows that (phyto)ceramide levels and composition are important for adjusting cilium length. Therefore, (phyto)ceramide is the first signaling sphingolipid demonstrated to metabolically regulate ciliogenesis by an evolutionarily conserved mechanism in *Chlamydomonas* and mammalian cells.
Materials and Methods

Cultivation of *Chlamydomonas* and ependymal cells

All experiments were carried out according to an Animal Use Protocol approved by the IACUC at GRU. *Chlamydomonas reinhardtii* (wild type strain CC-125 obtained from the *Chlamydomonas* Resource Center at the University of Minnesota) were grown in TAP growth medium (Cellgro, Manassas, VA) in 6-well plates or 250 ml Erlenmeyer flasks with a light/dark cycle of 14:10 h at room temperature. Primary cell culture of mixed glial cells was isolated from brains of 1-day-old C57BL/6 wild type mouse pups. Brains were dissociated in PBS containing 0.1 M glucose, passed through a 40 µm filter, and plated in T-25 flasks in DMEM (Gibco/Life technologies, Grand Island, NY ) supplemented with 10% FBS and 1% penicillin/streptomycin solution at 37°C in a humidified atmosphere containing 5% CO₂. After 7 days, cells were passed to 24-well plates with poly-D-lysine-coated glass coverslips and medium was changed to MEMα (Gibco) supplemented with bovine serum albumin (0.5 mg/ml), insulin (10 µg/ml; bovine), transferrin (5.5 µg/ml), sodium selenite (6.7 ng/ml), and human thrombin (1 unit/mL) to enrich for ependymal cells (Weibel et al., 1986). Cultures of >70% proportion of ependymal cells were visible after another 7 days cultivation in the thrombin-supplemented medium (purity was checked by immunocytochemistry for multiple vs. primary cilia and glial markers such as GFAP). Ependymal cells were treated with myriocin (Cayman, Ann Arbor, MI), GW4869 (Cayman), Fumonisin B1(FB1) (Enzo Life Science, Farmingdale, NY), BIO ((2’Z,3’E)-6-bromoindirubin-3’-oxime) (Sigma-Aldrich, St. Louis, MO), Lithium chloride (LiCl) (Sigma-Aldrich), myristoylated pseudosubstrate (aa113-125) inhibitor of PKCζ (PZI; Biomol, Plymouth Meeting, PA), and dihydrosphingosine (Avanti Polar Lipids, Alabaster, AL) at various concentrations and time periods in TAP or ependymal cell medium as indicated in Results.
**Lipid analysis**

ESI/MS/MS analysis of endogenous (phyto)ceramide species were performed on a Thermo Fisher Quantum triple quadrupole mass spectrometer, operating in a Multiple Reaction Monitoring (MRM) positive ionization mode, using modified version (Bielawski *et al.*, 2006). Briefly, all biological materials were fortified with the internal standards (ISs: C$_{17}$ base D-erythro-sphingosine (17CSph), C$_{17}$ sphingosine-1-phosphate (17CSph-1P), N-palmitoyl-D-erythro-C$_{13}$ sphingosine (13C16-Cer) and heptadecanoyl-D-erythro-sphingosine (C17-Cer) and C6-Phyto-ceramide), then extracted with ethyl acetate/isopropanol/water (60/30/10 %v/v) solvent system. After evaporation and reconstitution in 150 μl of methanol samples were injected on the HP1100/TSQ Quantum LC/MS system and gradient eluted from the BDS Hypersil C8, 150 x 3.2 mm, 3 μm particle size column, with 1.0 mM methanolic ammonium formate/2 mM aqueous ammonium formate mobile phase system. Peaks corresponding to the target analytes and internal standards were collected and processed using the Xcalibur software system.

Quantitative analysis was based on the calibration curves generated by spiking an artificial matrix with the known amounts of the target analyte synthetic standards and an equal amount of the internal standards (ISs). The target analyte/IS peak areas ratios were plotted against analyte concentration. The target analyte/IS peak area ratios from the samples were similarly normalized to their respective ISs and compared to the calibration curves, using a linear regression model. Introduction of the internal standards to the samples prior to extraction, yields results already “recovery corrected”, therefore, no further data manipulation is necessary.
RT-PCR

Total RNA was prepared from each experimental group of cells using TRIzol reagent following the manufacturer’s protocol (Invitrogen/ Life technologies). First-strand cDNA was synthesized using an iScript cDNA Synthesis kit according to the manufacturer’s instructions (Bio-Rad, Hercules, CA). GI:158275604 and GI:158278434 were used to design *Chlamydomonas* serine palmitoyltransferase 1 and 2 primers, respectively. GI:159490045 and GI:159465562 were used to design *Chlamydomonas* lag 1 and 2 primers, respectively. The following primers were used for RT-PCR:

mSPT1 - (sense): 5’- TCCCTCCAGTCTCCAAGAAC -3’

mSPT1 - (antisense): 5’- CAGGCTCTCCTCCAGGAATA -3’

mSPT2 - (sense): 5’- CCACCATGCAACAGAAAGAG -3’

mSPT2 - (antisense): 5’- AGGTTTCCAATTTCTCTGACG -3’

ChlamySPT1 - (sense): 5’- CCTGGCTTCCTACAACTACCTG -3’

ChlamySPT1 - (antisense): 5’- CAAACGACTTGGTGAACGTG -3’

ChlamySPT2 - (sense) 5’- CTGGAGGCGGAGGAGAAG -3’

ChlamySPT2 - (antisense) 5’- CGAGGCGGAGAAGCAGTAG -3’

Chlamylag1 – (sense) 5'- GTCGCCACACCACATCGTCACC -3’

Chlamylag1 – (antisense) 5’- GCAATCTCCACAATCTTGTAG -3’

Chlamylag2 – (sense) 5’- TCTGCGGGGTCTGCTGGACAC -3’

Chlamylag2 – (antisense) 5’- ATTGACGATGCGGAATG TGAG -3’
Deflagellation and reflagellation

*Chlamydomonas* cells were deflagellated in 10 mM Hepes by lowering pH to 4.5 with hydrochloric acid for 60 sec. pH was rapidly neutralized by the addition of sodium hydroxide, and cells were transferred to TAP medium for further incubation in the presence or absence of myriocin (Wilson and Lefebvre, 2004).

Phototaxis motility assay

To analyze phototaxis behavior, *Chlamydomonas* were cultivated at a density of logarithmic growth rate and then transferred to a 6-well dish. The wells were shielded from light in such a way that cells were only exposed to light coming from one defined direction. Cells with intact motility move toward and accumulate at the side exposed to light for 15-20 min. Phototaxis was quantified both by counting cells with a hemocytometer and by measuring optical absorbance of chlorophyll at 450 nm. All phototaxis experiments were performed 4 h after the beginning of the light phase (Lechtreck *et al.*, 2009).

Immunocytochemistry

*Chlamydomonas* or ependymal cells were grown in suspension or adherent on glass cover slips, respectively. Cells were fixed with 4% PFA/PBS for 15 min and then permeabilized by incubation with 0.2% Triton X-100 in PBS for 10 min at room temperature. Nonspecific binding sites were blocked with 3% ovalbumin/10% donkey serum/PBS for 1 h at 37 °C. Primary antibodies used were: anti-acetylated tubulin mouse IgG (1:3000, Sigma-Aldrich, clone 6-1113-1, Cat#. T6793), anti-ceramide rabbit IgG (1:100, our laboratory) (Krishnamurthy *et al.*, 2007a; He *et al.*, 2014), anti-ceramide mouse IgM (1:100, MAS0014, Glycobiotech, Kuekels, Germany), anti-pY216-GSK-3β rabbit IgG, anti-pY16-GSK-3β goat IgG, and anti-total GSK-
3α/β mouse IgG (1:100, Santa Cruz, cat#. sc-135653, sc-11758, sc-7291, Dallas, TX), anti-pSer9-GSK-3β rabbit IgG (1:100, Cell Signaling, cat#. 9336S), and anti-SPT2 antibody (Cayman cat. No. 10005260, 1:1000). Secondary antibodies (Alexa Fluor 546-conjugated donkey anti-rabbit IgG, Cy5-conjugated donkey anti-mouse IgM, μ-chain specific, Alexa Fluor 647-conjugated goat anti-mouse IgG γ-chain specific (all Jackson ImmunoResearch, West Grove, PA) were diluted 1:300 in 0.1% ovalbumin/PBS and samples incubated for 2 h at 37 °C. After washing, cover slips were mounted using Fluoroshield supplemented with DAPI (Sigma-Aldrich) to visualize the nuclei. Confocal fluorescence microscopy was performed using a Zeiss LSM780 upright confocal laser scanning microscope (Zeiss, Jena, Germany) equipped with a two-photon argon laser at 488 nm, 543 nm, or 633 nm (Alexa Fluor 647), respectively. LSM 510 Meta 3.2 software was used for image acquisition. Images obtained with secondary antibody only were used as negative controls representing the background intensity in a particular laser channel.

**Bifunctional ceramide analog (pacFACer) cross-linking and conjugation with fluorophore**

*Chlamydomonas* or ependymal cells were incubated under protection from light for 1 h in TAP medium or serum-free MEMα supplemented with vehicle (DMSO) or 5 μM of the bifunctional ceramide analog N-(9-(3-pent-4-ynyl-3-H-diazirine-3-yl)-nonanoyl)-D-erythro-sphingosine (pacFACer, Avanti Polar Lipids) diluted 1:1000 from a stock in ethanol/2% dodecane. Cells were UV irradiated at 365 nm for 5 min at room temperature and fixed with 4% p-formaldehyde/PBS for 15 min. Unbound pacFACer was removed by washing with in methanol, methanol/chloroform (1:1), and methanol again for 5 min each. Cells were equilibrated in PBS and the click reaction performed using the Click-iT Cell Reaction buffer Kit with Alexa Fluor
Ceramide (lipid) ELISA and binding to aPKCζ and GSK-3β

Binding assays were performed using a modified lipid (ceramide) ELISA as described previously (Krishnamurthy et al., 2007a; He et al., 2014; Dinkins et al., 2015). In brief, ceramides or pacFACer (500 ng in 50 µl ethanol) were coated on 96-well Immulon-1B ELISA plates (Thermo Scientific/Life Technologies) and washed with PBS. Non-specific binding sites were blocked by incubation for 1 h at 37 °C with 200 µl of 1% Fraction V BSA in PBS. Recombinant human aPKCζ (50 ng/50 µl, Enzo Life Science, Farmingdale, NY) or recombinant human GSK-3β (50 ng/50 µl, SignalChem, Richmond, Canada) in 0.1% BSA/PBS was incubated for 2 h at 37°C. Wells were washed 3 times with PBS and then incubated with 50 µl of 1:1000 α-PKCζ rabbit IgG (C20, Santa Cruz, cat# sc-216) or 1:500 α-GSK-3α/β mouse IgG (Santa Cruz, cat# sc-7291) overnight at 4 °C. Wells were washed 4 times with PBS, incubated with secondary antibody (50 µl of 1:2000 α-rabbit IgG HRP-conjugated in 0.1% BSA/PBS, Jackson ImmunoResearch) for 2 h at 37 °C, washed 4 times with PBS, and then developed using 0.8 mg/mL o-phenylenediamine and 0.03% hydrogen peroxide. After stopping the reaction with 1 M sulfuric acid, optical absorbance was monitored at 492 nm.

Miscellaneous

For immunoblot analysis, protein concentrations were determined using the RC/DC protein assay, in accordance with the manufacturer's (Bio-Rad, Hercules, CA) instructions. Equal amounts of protein were loaded onto a 4-20% gradient gel, and SDS-PAGE was performed using the Laemmli method. Ponceau S staining was performed to confirm equal loading. For immunoblotting, membranes were first blocked with 5% dry milk (or 3% BSA for
phosphorylated proteins) in PBST (PBS containing 0.1% Tween-20) and incubated with primary antibodies: SPT1/LCB1 antibody (BD Transduction Lab, 1:1000), SPT2 antibody (Cayman, 1:1000), Acetylated tubulin (Sigma-Aldrich, 1:10,000), total GSK-3α/β (Santa Cruz, 1:500), pY216-GSK-3β (Santa Cruz, 1:500), pS9-GSK-3β (Cell Signaling, 1:500) diluted in the blocking buffer overnight at 4°C. Membranes were then washed three times with PBST and incubated with the appropriate horseradish peroxidase-conjugated secondary antibodies (Jackson ImmunoResearch, 1:10,000) for 1 h at room temperature. After washing, bands were detected using chemiluminescence (ECL) reagent and exposure to X-ray film. Membranes were then stripped and re-probed, as described above, with anti-β-actin (1:2,000) to confirm equal loading.

Statistics
The mean, SEM, and statistical tests of control and treatment samples were calculated using GraphPad Prism. We used Student’s t-test to compare two groups, one-way ANOVA with Bonferroni post-hoc test for three or more groups, and non-linear regression for motility assay (Figure 3B). Values of at least p<0.05 were considered significant.

Acknowledgments
This study was supported by grants to E.B. (NSF: 1121579 and NIH/NIA: 1R01AG03438), and M.D. (NIH/NIA: 5 F32 AG044954-02). The authors are thankful to Dr. Christopher Poirier for providing the fro/fro mouse. They are also thankful to Dr. Matt Laudon, curator at the Chlamydomonas Resource Center at the University of Minnesota, St. Paul, MN. The authors thank for support by the Imaging Core Facility at GRU (under supervision by Drs. Paul and Ana McNeil). The authors are grateful to the Department of Neuroscience and Regenerative Medicine (Chair, Dr. Lin Mei) at Georgia Regents University for support.
References


Sphingolipid metabolism in animals, plants, algae, and fungi.

In animals, plants, and algae, *de novo* biosynthesis of ceramide and phytoceramide is initiated by conjugation of serine with palmitoyl-CoA (other activated fatty acids are possible but less frequent), a reaction catalyzed by serine palmitoyl transferase (SPT) and inhibited by myriocin. After another reaction, the intermediate dihydrosphingosine (DHS) is synthesized, which is then converted to ceramide via ceramide synthases (inhibited by fumonisin B1) in animals and
phytoceramide in plants, algae, and fungi. In animals, ceramide can be converted to sphingomyelin (SM). In the cell membrane or endocytic vesicles, ceramide can be rederived from SM by hydrolysis catalyzed by sphingomyelinases (SMases). Plants, algae, and fungi do not synthesize sphingomyelin but inositolphosphorylsphingolipids (not shown).
*Chlamydomonas* synthesizes phytoceramide, which can be blocked by myriocin.

(A) RT-PCR using SPT1 and SPT2 primers based on the mouse genome and *Chlamydomonas* expressed sequence tag (EST) clones. (B) Immunoblots for SPT1 and SPT2 in mouse ependymal cells and *Chlamydomonas*. (C, D) Sphingolipidomics LC-MS/MS analysis of phytoceramides (pmol/mg *Chlamydomonas* cells) in control and myriocin-treated (5 or 10 nM) cells (15 h). Data presented show mean ± SEM, n=3, ***p<0.001 (One-way ANOVA with Bonferroni post-hoc test).
test).
Figure 3

Ceramide depletion shortens cilia and leads to *Chlamydomonas* immotility, while exogenous DHS and C24:1 ceramide rescue *Chlamydomonas* flagella and ependymal cell cilia.

All graphical data presented show mean ± SEM. (A and B) Phototaxis motility assay (15 h; n=5; non-linear regression) with *Chlamydomonas* shows prolonged incubation with myriocin (myr), which leads to (A) time- and (B) dose-dependent immotility. (C) Phase contrast of *Chlamydomonas* showing flagella shortening in myriocin-treated cells. Scale bar, 10 µm. (D) Addition of DHS (1 µM) prevents immotility in myriocin-treated cells. (E) Motility assays using DHS and phytoceramide (Phyc) to rescue flagella in myriocin-treated *Chlamydomonas* (n=10, **p<0.01, ***p<0.001, One-way ANOVA with Bonferroni post-hoc test). (F)
Immunocytochemistry using control and C24:1 ceramide (1 µM)-treated ependymal cells (red, acetylated tubulin; blue, DAPI). Scale bar, 10 µm.(G) Quantitation of motile cilium length in ependymal cells treated for 48 h with ceramide biosynthesis inhibitors (5 µM myriocin, 10 µM fumonisin B1, FB1) with or without exogenous C24:1 ceramide (1 µM) (n=5, *p<0.05, **p<0.01, One-way ANOVA with Bonferroni post-hoc test).
Figure 4

Phytoceramide and ceramide are distributed to flagella and motile cilia, respectively.

(A) Lipid ELISA testing the specificity of anti-(phyto)ceramide rabbit IgG for various sphingolipids, mean ± SEM, n=5. (B) Immunocytochemistry using anti-(phyto)ceramide rabbit
IgG (green) and anti-acetylated tubulin mouse IgG (red) with *Chlamydomonas* and ependymal cells. In *Chlamydomonas*, arrows point at the base and tip of cilia. (C) *Chlamydomonas* was repeatedly (3x or 4x) de/reflagellated in the presence or absence of 10 nM myriocin. Incubation with myriocin without de/reflagellation for the duration of the experiment (5 h) was used as control. (D) Immunocytochemistry of de/reflaggelated *Chlamydomonas* in the presence or absence of myriocin using anti-(phyto)ceramide rabbit IgG (green) and anti-acetylated tubulin mouse IgG (red). (E) pacFACer (red) cross-linked in flagella tips and bases (arrowheads) co-labeled with anti-acetylated tubulin (green). Visualization was performed by labeling the cross-linked ceramide analog with Alexa594 using click chemistry. (F) UV cross-linking of pacFACer (red) followed by de/reflagellation led to formation of single, long flagella, which was absent when cells were incubated with pacFACer without UV cross-linking (anti-acetylated tubulin, green). Scale bars, (B) 10 µm left, 5 µm right, (D) 5 µm, (E) 2 µm, (F) 5 µm top, 20 µm bottom.
Figure 5

(Phyto)ceramide binds to pYGSK3 and sustains its phosphorylation level and localization in flagella and cilia.

(A) Immunoblot of pYGSK, total GSK3, and acetylated tubulin from Chlamydomonas treated for 24 h with myriocin (Myr, 5 or 10 nM) ± 1 µM dihydrosphingosine (DHS). Bottom panel shows Ponceau S staining for loading control. (B) Immunocytochemistry using antibodies against (phyto)ceramide rabbit (green), pYGSK (red), and acetylated tubulin (blue) with Chlamydomonas incubated for 60 min with GSK3β inhibitors BIO (2 µM) or lithium chloride (25 mM). (C) Higher magnification of flagella in left panel in (B). (D) Same as in (B) and (C).
with ependymal cells (Z-scan projection). (E) Z-scan orthogonal view of ependymal cells labeled with DAPI (blue) and antibodies against acetylated tubulin (green), pYGSK (green, left) and pSGSK (green, right). (F) C18 sphingomyelin and indicated ceramides were coated on ELISA plates, bound to recombinant GSK-3β, and binding quantified by development with anti-GSK-3α/β IgG and anti-mouse IgG-HRP. Mean ± SEM, n=3. (G) Immunoblot showing reactivity of recombinant human GSK-3β with anti-pYGSK3 IgG. Scale bars, (B) 5 μm, (C) 1 μm, (D) 2 μm, (E) 5 μm.
Figure 6

nSMase2-deficiency leads to shortening of motile cilia in vivo

(A) Immunoblot for pYGSK3, pSGSK3, and total GSK-3α/β using lysates from wildtype and tro/tro.
*fro/fro* ependymal cells. (B) Immunocytochemistry shows that pYGSK (green) is transported to cilia tips (arrows) in wildtype but not in *fro/fro* ependymal cells. (C) Immunocytochemistry for acetylated tubulin (red) comparing cryosections of wildtype and *fro/fro* brain (3 months old mice) shows that cilia are shorter in nSMase2-deficient ependymal cells. (D) Quantitation of A. Mean ± SEM, n=3, 10 sections/mouse. Students t-test, p<0.0001. (E) Immunocytochemistry for ceramide (green) shows punctate labeling along the ciliary membrane and tip. (F) Ceramide (green) is colocalized with pYGSK (blue). White punctae indicate colocalization of ceramide, pYGSK, and acetylated tubulin (red). Scale bars, (B) 5 µm, (C) 10 µm, (E, F) 1 µm.
Figure 7

Ceramide binds to aPKCζ, whose inhibition leads to cilium length extension.

(A) pacFACer was UV cross-linked to protein in primary cultured ependymal cells, linked to Alexa Fluor 594 azide (red), and labeled with anti-aPKCζ (magenta) and anti-acetylated tubulin (green) antibodies. Scale bar, 2 μm. (B) pacFACer was coated on ELISA plates and recombinant aPKCζ cross-linked by UV (365 nm) radiation. Presence/absence of pacFACer (pCer) and
recombinant aPKCζ along with primary antibody used are indicated. (C) Indicated ceramides were coated on ELISA plates, bound to recombinant aPKCζ, and binding quantified by development with anti-aPKCζ IgG and anti-rabbit IgG-HRP. Mean ± SEM, n=3. (D) Incubation of ependymal cells (24 h) with Fumonisin B1 (FB1, 10 µM) reduces cilium length, while addition of aPKCζ inhibitor PZI (10 µM) leads to extension of untreated or (partial) restoration of FB1-treated cilia. The GSK-3β inhibitor BIO (2 µM) obliterates cilia. Data presented show mean ± SEM, n=5, *p<0.05, ***p<0.001 (One-way ANOVA with Bonferroni post-hoc test).
Figure 8

Hypothetical model for flagella/cilia length regulation by (phyto)ceramide

(A) Flux equilibrium model. Lipid vesicles from the ceramide compartment (red) are
incorporated into the ciliary membrane at the cilium base. Lipid-cargo protein (green) co-transport ensures the stoichiometry required for simultaneous cilium elongation and membrane expansion. (Phyto)ceramide may serve as a membrane anchor for co-transported cargo proteins, or alternatively, it may activate a loading/unloading mechanism at the base or tip of the cilium. The cilium length is regulated by the size or ceramide content of the compartment at the base and its lipid flux to the cilium: more ceramide favors cilium assembly, less favors disassembly until flux rates in both directions are equal and the cilium length is maintained. (B) Regulation of GSK3 by (phyto)ceramide. In *Chlamydomonas* and ependymal cells, (phyto)ceramide binds to pYGSK3 at the ceramide-enriched compartment (ACEC) and transports it to flagella/cilia. In ependymal cells, ACEC-resident ceramide (red) binding and sequestration of aPKCζ prevents inactivation of GSK-3β by aPKCζ-mediated phosphorylation of ser9, which complements activation and transport of pYGSK3. In the cilium tip, pYGSK3 down-regulates HDAC6 or induces cargo release from the motor protein complex, most likely regulated by its interaction with (phyto)ceramide.