Developmental changes in ciliary composition during gametogenesis in Chlamydomonas

Miho Sakato-Antoku and Stephen M. King*
Department of Molecular Biology and Biophysics, University of Connecticut Health Center, Farmington, CT 06030-3305

ABSTRACT Chlamydomonas reinhardtii transitions from mitotically dividing vegetative cells to sexually competent gametes of two distinct mating types following nutrient deprivation. Gametes of opposite mating type interact via their cilia, initiating an intraciliary signaling cascade and ultimately fuse forming diploid zygotes. The process of gametogenesis is genetically encode, and a previous study revealed numerous significant changes in mRNA abundance during this life-cycle transition. Here we describe a proteomic analysis of cilia derived from vegetative and gametic cells of both mating types in an effort to assess the global changes that occur within the organelle during this process. We identify numerous membrane- and/or matrix-associated proteins in gametic cilia that were not detected in cilia from vegetative cells. This includes the pro-protein from which the GATI-amide gametic chemotactic modulator derives, as well as receptors, a dynamin-related protein, ammonium transporters, two proteins potentially involved in the intraciliary signaling cascade-driven increase in cAMP, and multiple proteins with a variety of interaction domains. These changes in ciliary composition likely directly affect the functional properties of this organelle as the cell transitions between life-cycle stages.

INTRODUCTION
Cilia are highly conserved microtubule-based cellular extensions that function as motile, sensory, and secretory organelles. These structures are of ancient origin, dating to before the divergence from the last eukaryotic common ancestor (Satir et al., 2008), and are found in a very broad array of extant organisms (Carvalho-Santos et al., 2011; Kumar et al., 2019). Cilia are extremely complex, and upward of 1000 different types of proteins are estimated to be involved in their formation and function (van Dam et al., 2019); this represents about 5% of the coding capacity of the human genome. In addition, cilia have a membrane lipid content distinct from that of the plasma membrane (Garcia et al., 2018). In mammals, defects in ciliary formation, motility, and/or signaling lead to numerous developmental defects and complex syndromes (ciliopathies) that may involve multiple organs, and have phenotypes ranging from infertility and polycystic kidneys to skeletal and neurological malformations, epilepsy, and insulin resistance (Reiter and Leroux, 2017).

The unicellular green alga Chlamydomonas reinhardtii has proven to be an exceptional system in which to study the organization and function of cilia, as it is readily amenable to genetic, biochemical, and physiological analysis (see various chapters in Witman, 2009). This organism grows as a haploid, with vegetative cells dividing mitotically. Following nutrient deprivation, these cells undergo a developmental process that results in formation of pregametes of two different genetically determined mating types (termed minus and plus); pregametes become sexually competent following exposure to blue light, which is detected by phototropin (Huang and Beck, 2003). These gametes exhibit opposite chemotactic responses to an amidated peptidergic signal; the peptide attracts minus gametes but repels plus gametes (Luxmi et al., 2019). As chemotaxing gametes retain their cell walls, it is quite possible that the receptor(s) responsible for signaling alterations in dynein-driven motility reside in the ciliary membrane. Gametes of opposite mating type initially interact via their cilia, which leads to complex intraciliary signaling and an increase in cAMP (reviewed in Snell and Goodenough, 2009). Ultimately, the gametes undergo cell fusion and form a quadriricate cell that further develops into a diploid zygote (Figure 1a). Once nutrients are again available,
zygotes pass through meiosis and hatch releasing haploid progeny (Sasso et al., 2018).

Although cilia on both vegetative and gametic cells are motile and power cell locomotion, they exhibit different responses to the chemotactic GATI-amide peptide (Luxmi et al., 2019), differential localization of adhesion molecules (Goodenough and Heuser, 1999), and altered intraciliary signaling (Pan and Snell, 2000). Thus, ciliary composition likely changes at various life-cycle stages. Although a comprehensive whole-cell transcriptomic analysis of mRNA abundance during gametogenesis in Chlamydomonas is available (Ning et al., 2013), a corresponding examination of proteins present specifically in gametic cilia, where release of the amidated ectosome-associated chemotactic modulator (Luxmi et al., 2019) and the initial steps of cell–cell contact and signaling occur during mating (Pan et al., 2003), has not been reported. Here we analyze the composition of cilia isolated from vegetative and gametic cells of both mating types to assess developmentally driven changes in ciliary composition. Our data reveal numerous alterations, especially in detergent-soluble components that likely directly affect ciliary activity, thereby altering organellar functions and responses at these life stages.

RESULTS AND DISCUSSION

Global analysis of vegetative and gametic cilia proteomes

To assess compositional differences that occur during gametogenesis, cilia were isolated from vegetative and gametic Chlamydomonas cells of both mating types and fractionated into a detergent-soluble membrane plus matrix fraction and an axonemal fraction; protein concentrations and sample volumes were carefully adjusted so that the same total cilia mass was analyzed for each preparation. These samples were then electrophoresed, stained with Coomassie blue, trypsinized, and subjected to LC-MS/MS analysis. The protein content of each sample is illustrated in Figure 1b, and the actual triplicate “short gel” samples used for mass spectrometry are shown in Figure 1c. A global analysis of the proteins identified and how many were unique to a particular fraction is provided in Supplemental Table S1. The complete dataset indicating the normalized total spectral counts for every protein identified in each replicate is provided in Supplemental Table S2.

A global comparison of the abundance of every identified protein in vegetative and gametic cilia from both mating types and the Pearson correlation coefficients (r) between vegetative/gametic and mating type minus/plus datasets are shown in Figure 2. Axonemal samples show a very high degree of consistency, with few proteins deviating from an approximate 1:1 ratio. This provides a direct measure of the equivalence of cilia amounts between the samples. In contrast, considerably more variation is evident in the detergent extracts.

To assess intersample consistency of the numeric data, we analyzed the abundance of multiple core axonemal components required for normal ciliary motility that would not be expected to

![FIGURE 1: Cilia samples used for mass spectrometry. a) Schematic illustrating the transition from vegetative to gametic Chlamydomonas cells, and formation of a diploid quadriciliate cell that subsequently develops into a zygote. Mating type is indicated by – and + signs within the nucleus. b) Axoneme and detergent-soluble membrane plus matrix samples from vegetative (Veg) and gametic (Gam) cilia of wild-type Chlamydomonas of both mating types (CC124 and CC125) were electrophoresed in 4–15% gradient gels and stained with Coomassie blue; in combination, the amounts loaded for membrane plus matrix and axoneme fractions of each sample were 50 μg cilia. The number of proteins identified in each sample is indicated at the bottom of each gel. c) Additional aliquots of the samples shown in panel b were run in triplicate using a short gel format and stained with Coomassie blue. Following imaging, the protein-containing gel segments were excised, trypsinized, and subjected to mass spectral analysis.](image-url)
vary either between vegetative and gametic cilia or as a function of mating type. These include components of the inner and outer dynein arms, radial spokes, protofilament ribbons, nexin–dynein regulatory complex, and central pair microtubule complex (Figure 3). These proteins show highly consistent abundance levels between the cilia preparations and their replicates with small standard deviations. One exception is LC8, which is part of multiple axonemal complexes and shows enhanced levels in the mating type plus gametic cilia detergent extract; this correlates well with an increase in the dynein heavy chain that powers retrograde intraflagellar transport in the same fraction (Figure 2). Furthermore, although both α- and β-tubulins (Cre04.g216850 and Cre12.g549550) were present in large and consistent amounts, only two peptides were identified in a single sample for ε-tubulin (Cre03.g172650) and none for either γ- (Cre06.g299300) or δ-tubulin (Cre03.g187350); these three proteins are known to be in the basal body, and in the case of γ-tubulin also the transition zone, but generally absent from the cilium proper e.g. (Dutcher and Trabuco, 1998; Silflow et al., 1999; Dutcher et al., 2002).

Multiple proteins are exclusively present in gametic cilia
We next asked what proteins were exclusively found in gametic and not vegetative cell cilia, as this might highlight key signaling and/or regulatory pathways. The initial cutoff criterion used was stringent, requiring that zero peptides for a protein be found in any of the 12 vegetative cilia samples examined (i.e., membrane/matrix and axonemal samples from both mating types analyzed in triplicate). A total of 33 proteins met this criterion (Table 1), including a dynamin-related protein, which contains the dynamin central and effector domains plus a plekstrin homology region but lacks the GTPase motif, two annotated ammonium transporters, and several putative receptors. Importantly, only gametic samples contained the GATI-amide chemotactic modulator (Cre03.g204500), which is present in gametic cilia (Luxmi et al., 2019) and essentially not expressed in vegetative cells (Ning et al., 2013). Proteins present in both vegetative and gametic cilia but whose abundance changed ≥5-fold between these samples are indicated in Supplemental Tables S3 and S4; these include a dynamin-related GTPase, a RAN GTPase activating protein, multiple subunits of the T-complex chaperonin, ion channels, and putative receptors.

Also identified only in gametic cilia were two proteins with domains suggesting a potential role in the signaling pathway, where ciliary adhesion leads to an intraciliary kinase cascade and ultimately an increase in cellular cAMP that presages
cell fusion and zygote formation (Figure 4 and Table 1). One (Cre03.g199050) contains four N-terminal cNMP binding domains and a C-terminal Ser/Thr kinase module and thus might be a downstream effector following cAMP increase. The second is CYA18 (Cre06.g300500), which consists of a noncanonical K\(^{+}\) channel followed by an adenylyl cyclase domain. A similar protein was previously identified in ciliates and Plasmodium and demonstrated to exhibit both ion channel properties and cyclase activity (Schultz et al., 1992; Weber et al., 2004); BLAST searches also reveal CYA18 orthologs in haptophytes, cryptophytes, and stramenopiles, but not opisthokonts or excavates. CYA18 is the most abundant channel and cyclase in gametic cilia and may represent a cyclase activated during mating-induced signaling. The presence of an N-terminal channel domain in CYA18 indicates a potentially direct connection between cAMP formation and K\(^{+}\) flux that might play an important role in gametic ciliary signaling. This concept can be tested readily, as inward K\(^{+}\) flux is easily controlled by external application of specific channel blocking agents. An additional (putative guanylate) cyclase (CYG38) was also present in low abundance only in gametic cilia.

Generation of 3′-5′ cyclic nucleotides from NTPs releases pyrophosphate, which must be hydrolyzed to generate a sufficiently negative change in ΔG′ so that the back reaction does not occur. Intriguingly, a single pyrophosphatase (IPY3) was present in the membrane/matrix fractions of both vegetative and gametic cilia samples; this implies that there may be multiple pathways (e.g., ubiquitin activation) that lead to ciliary generation of pyrophosphate that must be hydrolyzed. Although not quite making the zero peptide cutoff, abundance of the co-chaperone Sti1p/HOP1 that interacts with and regulates HSP70/HSP90 also increased dramatically in gametic cilia (Figure 4).

Several proteins were mainly or exclusively present in gametic samples of a single mating type, such as the minus agglutinin (Cre06.g254917) that localizes to the ciliary membrane and mediates direct associations with agglutinin molecules on the surface of plus gamete cilia (Ferris et al., 2005). Intriguingly, the plus agglutinin was not identified in any sample, which may reflect low sequence complexity (it is 12% proline, 18% alanine, and 12% glycine) and/or very different abundance levels in resting gametes; indeed, only a few peptides for this protein were identified previously in ectosomes from actively mating gametes (Luxmi et al., 2019). Several other mating type–specific proteins are more enigmatic. Cre02.g079500 encodes a protein with a canonical signal sequence placing it in the secretory pathway and a C-terminal transmembrane segment. It is essentially Chlamydomonas-specific and only weak homology (∼35% identity) was found even in volvocine relatives. Cre06.g278222 consists of seven WD repeats and exhibits 76% identity to the vertebrate receptor for activated protein kinase C 1-like (RACK1) adaptor that is involved in protein kinase C signaling. In addition, a clathrin heavy chain (Cre02.g101400) was identified that is part of a coexpression cluster associated with mating activation induced in minus gametes (Molla-Herman et al., 2010); clathrin has been reported to be present at the ciliary pocket (Clement et al., 2013) and to affect ciliary assembly. In contrast, an E2 ubiquitin-conjugating enzyme UBC21 encoded at Cre12.g510300 was present in plus gamete cilia but apparently absent from vegetative cell cilia.

Previous proteomic studies of Chlamydomonas vegetative cell cilia have defined the core components of these organelles; see for example (Pazour et al., 2005) and http://chlamyfp.org. With a few exceptions, the proteins identified here only in gametic cilia have not
Numerous flagellar-associated proteins exhibit altered abundance in gametic versus vegetative cell cilia

The original *Chlamydomonas* ciliary proteome identified numerous proteins of unknown function that were given the generic designation FAP (flagellar-associated protein; Pazour et al., 2005). We find that several show dramatic decreases in abundance following gametogenesis, while others increase considerably. These proteins were solubilized, either partially or completely, by detergent treatment, indicating that they are components of the membrane and/or the soluble ciliary matrix and not integral axonemal proteins. This supports the concept that the ciliary membrane undergoes multiple alterations in functional specialization as cells transition to mating competency.

Four of these proteins appear to be membrane-associated. FAP24 contains a signal sequence placing it on the external face of the ciliary membrane; levels are reduced by ~50% in gametic cilia. It also has a TroA-like domain that usually function as ferric siderophores and/or are involved in binding transition metals such as Mn²⁺, Cu²⁺, and Zn²⁺. FAP102 exhibits a more complex abundance pattern. Although present and essentially unchanging in vegetative and gametic plus mating type cilia, FAP102 is almost completely absent from mating type minus vegetative cilia and increases more than 40-fold during gametogenesis. FAP102 contains a signal sequence but no obvious structural domains and is redox active (Wakabayashi and King, 2006). Two transmembrane proteins, FAP154 and FAP49, which are almost identical (except at the extreme termini) and encoded by adjacent genes, are abundant in vegetative cell cilia and decrease by ~5-fold following gametogenesis. Both have seven N-terminal transmembrane domains, a central PAS domain, and a C-terminal region containing at least four additional transmembrane segments; PAS domains act as environmental sensors, often of light, oxygen, or other small molecules.

Multiple FAPs that are reduced and/or absent in gametic cilia contain protein–protein interaction motifs including ankyrin repeats (FAP26 and FAP79), armadillo/β-catenin-like repeats (FAP28), tetratricopeptide repeats (FAP185), and a prefoldin-like domain (FAP88). In addition, FAP280 is reduced ~10-fold in gametes. This membrane/matrix-associated protein contains a multiprotein bridging factor domain found in a wide range of eukaryotes and is potentially involved in transcriptional coactivation. Finally, two additional proteins (FAP177 and FAP181) with no obvious domains are both reduced following gametogenesis.

Several proteins were found almost exclusively in gametic axonemes. For example, Cre10.g423600 encodes a protein containing protein–protein interaction motifs including ankyrin repeats (FAP26 and FAP79), armadillo/β-catenin-like repeats (FAP28), tetratricopeptide repeats (FAP185), and a prefoldin-like domain (FAP88). In addition, FAP280 is reduced ~10-fold in gametes. This membrane/matrix-associated protein contains a multiprotein bridging factor domain found in a wide range of eukaryotes and is potentially involved in transcriptional coactivation. Finally, two additional proteins (FAP177 and FAP181) with no obvious domains are both reduced following gametogenesis.
| Gene identifier | Protein name and/or structural/functional attributes | Protein identified in membrane plus matrix, axoneme, or both fractions | Protein identified in minus/plus mating type | Identified in previous cilia proteomic analyses<sup>b</sup> | mRNA abundance (median RPKM %; data from Ning et al., 2013)<sup>c</sup> |
|----------------|-----------------------------------------------------|-----------------------------------------------------------|---------------------------------|---------------------------------|---------------------------------|
| Cre01.g012700  | KCN6. Voltage-dependent K<sup>+</sup> channel       | Both                                                       | Both                            | No                              | 0.05                            |
| Cre02.g079500  | Signal sequence and one transmembrane domain. Mostly in minus gametes | Mainly membrane plus matrix                               | Minus (minimal present in plus) | No                              | 1.34                            |
| Cre02.g079550  | Dynamin-related                                     | Both                                                       | Both (-3× greater in minus)     | No                              | 8.41                            |
| Cre02.g101400  | Contains 7 clathrin heavy chain repeats and present in minus gametes only | Both                                                       | Minus                           | No                              | 26.2                            |
| Cre03.g199050  | cNMP binding domains and C-terminal Ser/Thr kinase domain | Both                                                       | Both (minimal present in plus)  | No                              | 0.70                            |
| Cre03.g204500  | Ectosome-associated amidated chemotactic modulator    | Both                                                       | Both                            | No                              | 0.75                            |
| Cre05.g245950  | Dynamin DRP1                                         | Both                                                       | Minus                           | No                              | 8.41                            |
| Cre06.g254917  | SAD1 minus agglutinin. Transmembrane protein         | Both                                                       | Minus                           | Listed, but no peptides indicated | No data available               |
| Cre06.g278222  | Contains 7 WD repeats and present only in minus gametes | Both                                                       | Minus (minimal present in plus) | Yes                             | No data available               |
| Cre06.g283800  | OB fold protein                                      | Membrane plus matrix                                       | Both                            | No                              | 5.03                            |
| Cre06.g294150  | No obvious domains, almost exclusively axonemal       | Axoneme (1 peptide in Membrane)                            | Both                            | No                              | 0.10                            |
| Cre06.g300500  | CYA18, N-terminal non-canonical K<sup>+</sup> channel with C-terminal adenylyl cyclase domain | Both                                                       | Both                            | No                              | 1.58                            |
| Cre06.g310750  | Vesicle coat COPI complex gamma subunit               | Membrane plus matrix                                       | Both                            | No                              | 14.89                           |
| Cre07.g347150  | No obvious domains                                    | Membrane plus matrix                                       | Minus                           | Yes – one study only            | 4.71                            |
| Cre08.g360050  | Allophanate hydrolase/urea carboxylase                | Membrane plus matrix                                       | Both                            | No                              | 0.27                            |
| Cre08.g364400  | Transcriptional regulator DNA-binding domain          | Membrane plus matrix                                       | Both                            | No                              | 8.89                            |
| Cre08.g364600  | Transmembrane protein                                | Both                                                       | Both                            | No                              | 1.92                            |
| Cre09.g402515  | Copper amine oxidase (tyramine oxidase)               | Both                                                       | Both                            | No                              | 13.25                           |
| Cre09.g414050  | RuvBL1 (Pontin)                                      | Both                                                       | Both                            | Listed, but no peptides indicated | No data available              |

TABLE 1: Proteins identified only in gametic cilia<sup>a</sup>. (Continues)
| Gene identifier | Protein name and/or structural/functional attributes | Protein identified in membrane plus matrix, axoneme, or both fractions | Protein identified in minus/plus mating type | Identified in previous cilia proteomic analyses | mRNA abundance (median RPKM %; data from Ning et al., 2013)* | Synchronous vegetative cells | Minus resting gametes | Plus resting gametes |
|-----------------|------------------------------------------------------|---------------------------------------------------------------|---------------------------------|---------------------------------|-------------------------------------------------|-----------------------------|-------------------------|-------------------------|
| Cre09.g415800   | Programmed cell death protein 4                      | Membrane plus matrix                                          | Both                            | No                              | 24.33                                          | 18.11                       | 12.35                   |
| Cre10.g454450   | Protein phosphatase 2C-like                           | Membrane plus matrix                                          | Both                            | No                              | 8.17                                          | 66.04                       | 63.95                   |
| Cre12.g488850   | Alpha 2 adaptin                                      | Mainly membrane plus matrix                                   | Both                            | No                              | 6.78                                          | 25.48                       | 31.14                   |
| Cre12.g493100   | No obvious domains                                   | Mainly axonemal                                               | Both                            | No                              | 0.45                                          | 16.35                       | 70.88                   |
| Cre12.g496350   | Patched-related sterol sensing protein (single peptide also found in one vegetative cilia sample) | Both                                                          | Yes                             | Yes                             | 11.92                                         | 18.35                       | 24.30                   |
| Cre12.g506400   | No obvious domains                                   | Membrane plus matrix                                          | Both                            | No                              | 1.81                                          | 3.19                        | 2.48                    |
| Cre12.g510300   | Ubiquitin conjugating enzyme E2; mainly plus gametes | Membrane plus matrix                                          | Both                            | No                              | 0.77                                          | 0.97                        | 2.04                    |
| Cre12.g515350   | Transmembrane protein with lectin-binding and polycystin channel domains (single, low-probability peptide also found in one vegetative cilia sample) | Both                                                          | Both                            | No                              | 0.94                                          | 6.03                        | 13.77                   |
| Cre12.g532850   | Ionotopic glutamate receptor family. Periplasmic binding type 2 domain | Both                                                          | Both                            | No                              | 0.54                                          | 55.85                       | 637.5                   |
| Cre12.g537400   | Ser/Thr aurora kinase                                | Both                                                          | Both                            | No                              | 2.42                                          | 77.01                       | 228.54                  |
| Cre13.g569850   | AMT4 NH₄ transporter with 11 transmembrane domains   | Both                                                          | Both                            | No                              | 2.31                                          | 1331.24                     | 971.47                  |
| Cre13.g602400   | KIN8 kinesin (Kif18/19 family member; single peptide also found in one vegetative cilia sample) | Axoneme                                                       | Yes—one study only              | No data available               | 0.95                                          | 8.78                        | 7.37                    |
| Cre14.g629920   | NH₄ transporter with 10 transmembrane domains        | Both                                                          | Both                            | No                              | No data available                             |                             |                         |
| Cre16.g685277   | Adaport complex protein mu subunit homolog          | Membrane plus matrix                                          | Both                            | No                              | No data available                             |                             |                         |
| Cre16.g685650   | Ligand-gated ion channel with N-terminal extracellular solute binding domain | Both                                                          | Both                            | No                              | 0.92                                          | 66.57                       | 269.48                  |
| Cre17.g714300   | No obvious domains                                   | Membrane plus matrix                                          | Both                            | No                              | 0.00                                          | 20.25                       | 18.86                   |
| Cre17.g735450   | SUB14 subtilisin-like protease                       | Membrane plus matrix                                          | Both                            | No                              | 0.81                                          | 12.34                       | 47.75                   |

*aAlso included in this table are three intriguing proteins (Dynamin DRP1, a Patched-related receptor, and the Kin8 kinesin) for which a single peptide was found in one of 12 vegetative cilia samples.

*bThe results from previous proteomic analyses of Chlamydomonas cilia published by Pazour et al. (2005), Wang et al. (2017), Jordan et al. (2018), Picariello et al. (2019), Zhao et al. (2019), and Dai et al. (2020) are tabulated at http://chlamyfp.org/.

*cThese data are from Ning et al. (2013) and were obtained using the 21gr (mating type plus) and 6145C (mating type minus) wild-type strains. The color code is green = > fourfold abundance increase; yellow = abundance increase, but < four-fold; orange = no significant increase.

TABLE 1: Proteins identified only in gametic cilia*. Continued
an N-terminal NTPase domain, 3 EF-hands that are predicted to bind Ca\(^{2+}\), and 11 C-terminal WD repeats. This increased ∼5-fold in minus gamete cilia samples and over 10-fold in plus gamete samples. Similarly, the Cre06.g294150 protein, which has no obvious domains and multiple long regions of low complexity, was not found in any vegetative cilia sample.

**Changes in ciliary receptors following gametogenesis**

*Chlamydomonas* contains almost 150 genes annotated as encoding various receptors including members of the scavenger, ionotropic glutamate, seven-transmembrane domain (listed as G protein–coupled receptor-related), patched-related, Toll-like, and lectin-binding receptor families, as well as the blue light receptor phototropin (Huang et al., 2004; Merchant et al., 2007; Luxmi et al., 2019).

We identified a patched-related receptor (Cre12.g496350) almost exclusively in gametic cilia; however, several previous analyses also found evidence for this protein in vegetative cilia (Jordan et al., 2018; Picariello et al., 2019; Zhao et al., 2019), and its mRNA expression levels only change by ∼2-fold during gametogenesis (Ning et al., 2013; see Table 1). The abundance of a “GPCR-related” protein (Cre13.g604050) with extracellular peptidase lyase repeats likely involved in carbohydrate binding showed a similar gametic-specific pattern with only a single peptide found in one vegetative cilia sample; this putative receptor has not been identified in cilia previously. Importantly, although the topology of this protein is superficially similar to that of GPCRs, *Chlamydomonas* lacks canonical G\(_{q}\) subunits (Merchant et al., 2007; Urano et al., 2012). An additional transmembrane protein (Cre12.g532850) in the ionotropic glutamate receptor family and containing a type 2 periplasmic binding fold often present in small molecule sensors was present only in gametic cilia; this correlates well with the large increase in mRNA expression seen in gametic samples (Ning et al., 2013 and Table 1). In contrast, peptides from a scavenger receptor (Cre05.g240700) and the blue light receptor phototropin required for the acquisition of mating competency decreased in gametic samples.

**mRNA and protein abundance changes following gametogenesis**

Whole-cell transcriptomic analyses are often used as a proxy for proteomic studies; if the mRNA increases the general assumption is that protein levels follow, although the comparative magnitude and timing of that increase is not always clear. Our analysis of the ciliary proteomics of gametogenesis combined with the detailed transcriptomic study of Ning et al. (2013) provides an opportunity to make a broad assessment of the correlation between these measures at the organelle level. The transcriptomic analysis measured whole cell mRNA levels in both asynchronous and synchronized vegetative cells and resting gametes, as well as gametes treated with lysozyme to remove the cell walls or activated with dibutyryl cAMP (Ning et al., 2013). One important caveat for this comparison is that the wild-type strains used in the mRNA analysis are different from those used here for proteomics, which may introduce strain-specific anomalies. Even so, in general there is a strong correlation between increased mRNA levels and the appearance of a protein in cilia following gametogenesis (Table 1). In a few cases, this correlation fails, which might result from experimental issues or incorrect identification of a protein as gamete-specific or be due to vegetative cells maintaining a store of the protein product in the cytoplasm and only relocating it to cilia following a gametogenic signal.

In conclusion, this comparative analysis has identified numerous developmental changes in ciliary composition and found both vegetative cell– and gamete-specific cilia components. Furthermore, it provides a resource for understanding functional specializations that occur in cilia from *Chlamydomonas* cells of different mating types following life-cycle stage transitions.

**METHODS**

**Chlamydomonas culture**

Wild-type *Chlamydomonas reinhardtii* strains CC124 (mating type minus) and CC125 (mating type plus) were grown in 2 × 1 l cultures to a density of ∼5 × 10\(^6\) cells/ml in R medium containing acetate on a 12 h:12 h light/dark cycle and aerated with a 5:1 air to CO\(_2\) mixture. One culture of each strain was processed as described below to provide vegetative cell cilia samples. The second culture was harvested by centrifugation (Fiberte F10 rotor, 1100 × g, 7 mins, 20°C), resuspended in M-N/5 medium and incubated overnight to induce gametogenesis. Successful gamete formation was assessed by mating and formation of quadriciliate cells. This second culture was then processed to provide the gametic cilia samples.

**Cilia isolation and fractionation**

Cilia isolation was performed following (Craigie et al., 2013) with modifications. Cells were harvested by low speed centrifugation (Fiberte F10 rotor, 1,100 × g, 7 min, 20°C), and washed three times for vegetative cells and twice for gametic cells with 10 mM HEPES pH 7.5. Cells were then resuspended in 30 mM HEPES pH 7.5, 5 mM MgSO\(_4\), 4% sucrose (HMS; 10 ml per tube) on ice; all solutions contained protease inhibitor cocktail (Sigma-Aldrich, P9599) and 1 mM DTT hereafter. Subsequently, 100 μl of 5.3% (wt/vol) CaCl\(_2\) was added to each tube followed by 2 ml of 25 mM dibucainc.HCl to induce deciliation, which was assessed by phase contrast microscopy. Deciliated cell bodies were collected by low speed centrifugation (1800 xg, 5 min in a Sorval ST8 centrifuge with a swing-out rotor).

The supernatant containing detached cilia was laid over a 25% sucrose solution made in 30 mM HEPES pH 7.5, 5 mM MgSO\(_4\). The tubes were then spun in a swing-out rotor (2400 × g, 10 min in a Sorval ST8 centrifuge) to pellet any remaining cell bodies through the sucrose under layer. The top sucrose layer and the 4%/25% sucrose interface (the latter contains most of the cilia) were transferred into fresh tubes. Cilia were harvested by centrifugation in a Fiberlite F215 8 × 50y rotor (30,000 × g, 20 min, 4°C). Pellets were resuspended in 0.5 ml of HMEK buffer (30 mM HEPES pH 7.5, 5 mM MgSO\(_4\), 0.5 mM EDTA, 25 mM KCl) and spun in a Fiberlite F215 8 × 50y rotor (30,000 × g, 20 min, 4°C). Isolated cilia were resuspended in appropriate volumes of HMEK to adjust the concentration to 4 mg/ml. Protein concentration was determined with the BCA assay using BSA as standard. IGEPAL-CA630 detergent (Sigma-Aldrich, I3021) was then added to a final concentration of 1% (vol/vol) and extraction allowed to occur on ice for 30 mins with mixing every 10 mins. Following centrifugation (Fiberte F215 8 × 50y rotor, 30,000 × g, 20 mins, 4°C), the supernatant was removed and used as the membrane plus matrix sample, while the extracted axoneme pellet was resuspended in gel sample buffer directly.

Gel samples derived from equal total cilia mass equivalents (50 μg) were denatured at 80°C for 10 min and fractionated in 4–15% SDS gradient MINI-PROTEAN TGX polyacrylamide gels (Biorad, Hercules, CA). For mass spectrometry, samples were separated using a short gel protocol in which the sample entered only about 2 cm into the gel. Each sample was run in triplicate in a different electrophoresis unit to ensure crossover contamination did not occur. Following staining with newly prepared Coomassie blue, each lane was excised, subject to in-gel tryptic digestion and the peptide products analyzed by mass spectrometry (see below).
Mass spectrometry
Trypsinized cilia samples were spiked with 2 pmol trypsin-digested yeast dehydrogenase (ADH) for quality control and normalization. Mass spectral (LC-MS/MS) analysis was performed at the University of Massachusetts medical school mass spectrometry facility. Each sample was injected into a Thermo Scientific Q exactive quadrupole-Orbitrap hybrid mass spectrometer with Waters NanoAcquity ultra-performance liquid chromatography. Six ADH peptides were used to evaluate the acquisitions and yielded a mass error of ~2 ppm. Data were searched against the *Chlamydomonas* genome v.5.5 with Mascot in Proteome Discoverer 2.1.1.21 using a fragment tolerance of 0.050 and a parent ion tolerance of 10 ppm; a maximum of two missed cleavages were allowed. Modifications assessed were carboxamidomethyl on Cys, C-terminal Gly-loss plus amide, N-terminal pyroglutamylation on Gln, oxidation on Met, and N-terminal acetylation and phosphorylation on Ser, Thr, and Tyr. Protein threshold was set to 99.0% minimum with a two-peptide minimum; data were analyzed using Scaffold ver. 5.0.1.

Total normalized spectral count data were plotted using GraphPad Prism ver.7. For bar charts, mean ± SD is shown; for scatterplots of vegetative versus gametic cilia samples, the average spectral counts from *n* = 3 samples were used. Pearson correlation coefficients (*r*) were calculated using GraphPad Prism.

Domain Analysis
Protein domain organization was analyzed using SMART (http://smart.embl-heidelberg.de/), Prosite (https://prosite.expasy.org/), and cDART (https://www.ncbi.nlm.nih.gov/Structure/lexington/docs/cdart_about.html). SignalP was employed to assess the presence of potential signal sequences (https://services.healthtech.dtu.dk/service.php?SignalP-5.0), and TOPCONS used to predict membrane protein topology (https://topcons.net/2; Tiriglos et al., 2015).

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