RNA Interference Silencing of a Major Lipid Droplet Protein Affects Lipid Droplet Size in *Chlamydomonas reinhardtii*\(^{\dagger}\)

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Eukaryotic cells store oils in the chemical form of triacylglycerols in distinct organelles, often called lipid droplets. These dynamic storage compartments have been intensely studied in the context of human health and also in plants as a source of vegetable oils for human consumption and for chemical or biofuel feedstocks. Many microalgae accumulate oils, particularly under conditions limiting to growth, and thus have gained renewed attention as a potentially sustainable feedstock for biofuel production. However, little is currently known at the cellular or molecular levels with regard to oil accumulation in microalgae, and the structural proteins and enzymes involved in the biogenesis, maintenance, and degradation of algal oil storage compartments are not well studied. Focusing on the model green alga *Chlamydomonas reinhardtii*, the accumulation of triacylglycerols and the formation of lipid droplets during nitrogen deprivation were investigated. Mass spectrometry identified 259 proteins in a lipid droplet-enriched fraction, among them a major protein, tentatively designated major lipid droplet protein (MLDP). This protein is specific to the green algal lineage of photosynthetic organisms. Repression of MLDP gene expression using an RNA interference approach led to increased lipid droplet size, but no change in triacylglycerol content or metabolism was observed.

Triacylglycerols (TAGs) are stored in lipid droplets which are subcellular structures in specialized cells ubiquitous to eukaryotes but have more recently also been identified in some prokaryotes (26). In plants and animals, lipid droplets are surrounded by cytosol and are believed to bud off the endoplasmic reticulum (ER) (15, 26). While traditionally considered merely as storage compartments, recent studies suggest that lipid droplets in animals play important additional roles in lipid homeostasis and protein storage (8). In oilseed plants, TAG accumulated in seeds is used as a reservoir of energy and membrane lipid building blocks to support rapid growth after germination (15). Many green algae are capable of accumulating large amounts of TAG in lipid droplets, particularly as a result of abiotic stresses, such as nutrient deprivation or high-light exposure. Although TAG metabolism in algae has not yet been extensively studied at the biochemical or molecular level, it is proposed that TAG turnover contributes primarily to the assembly of membrane lipids to facilitate rapid cell division after the cessation of nutrient limitation (14, 38).

The general structure of lipid droplets is conserved in different species with a globular neutral lipid core enclosed by a membrane lipid monolayer (26). In addition, specific proteins are associated with lipid droplets and play important roles in lipid droplet structure and function. A number of recent proteomic studies of lipid droplets from different animals and tissues (8, 40), *Saccharomyces cerevisiae* (4), and plants (16, 17) have revealed that the lipid droplet-associated proteins of these organisms are quite distinct. For example, the abundant lipid droplet proteins in animals—the so-called “PAT” family of proteins comprised of perilipin, adipose differentiation-related protein (ADRP), and TIP47 (21)—have no apparent orthologs in the desiccating seed plant *Arabidopsis thaliana*; conversely, the oleosins which coat the oil bodies of *Arabidopsis* and many other seed plants are not found in animals (26). Reverse genetic studies of these proteins have helped to elucidate the role of *A. thaliana* oleosins in regulating lipid droplet size and preventing droplet fusion (35, 36) or that of mouse adipocyte perilipin in regulating lipolytic activity at the lipid droplet surface (37). Moreover, recent genomewide RNAi screens in *Drosophila* cells implicated 1.5 to 3.0% of all genes as directly or indirectly involved in lipid droplet formation and/or regulation and resulted in the identification of a new role for the Arf1-COPI vesicular transport machinery in regulating droplet morphology and lipid utilization (7, 12). In contrast, few molecular details are known about algal lipid droplet biogenesis although many TAG-rich algal species have been described (14).

Our efforts to identify proteins related to the PAT protein family or oleosins in the *Chlamydomonas reinhardtii* genome (24) or genomes of other green algal and diatom species, including *Thalassiosira pseudonana*, *Volvox carteri*, and *Chlorella* sp. NC64A, revealed no putative algal orthologs. In order to identify both potentially novel and conserved proteins which function in algal lipid droplet biogenesis, we studied the accumulation of TAG in lipid droplets of nitrogen-limited *C. reinhardtii* cells and identified candidate lipid droplet-associated proteins by mass spectrometry.

**MATERIALS AND METHODS**

**Strains and growth conditions.** *Chlamydomonas reinhardtii dw15* (cw15 nit1 mt\(^+\)) and CC-125 (nit1 nit2 mt\(^-\)) strains were grown in Tris-acetate-phosphate (TAP) medium with NH\(_4\)Cl at 10 mM in liquid cultures or on agar-solidified plates under continuous light (70 to 80 \(\mu\)mol m\(^{-2}\) s\(^{-1}\)) as previously described.
droplets from cells deprived of N for 24 h (data not shown); as such, lipid droplets from cells deprived of N for 24 h were used throughout in this study. Thylakoid membranes were isolated according to Chua and Bennoun (9), and enriched eyespots were prepared as previously described (33).

Mass spectrometric identification of lipid droplet proteins. Proteins were precipitated as previously described (33) and solubilized in sodium dodecyl sulfate (SDS) sample buffer by sonication. Proteins were separated on 4 to 20% SDS-PAGE gels (Bio-Rad, Hercules, CA) and stained with E2Blue gel staining reagent (Sigma-Aldrich, St. Louis, MO). The lane containing lipid droplet proteins was sliced into 15 bands, which were processed for in-gel trypsin digestion as previously described (34) and analyzed by liquid chromatography-tandem mass spectrometry (LC-MS/MS) by the Michigan State University Research and Technology Facility. The extracted peptides were resuspended in a solution of 2% acetonitrile-0.1% trifluoroacetic acid to 20 μL. From this, 10 μL were automatically injected by a Waters nanoAcquity sample manager (Waters, Milford, MA) and loaded for 5 min onto a Waters Symmetry C18 peptide trap (5 μm, 180 μm by 20 mm) at 4 μL/min in 2% acetonitrile-0.1% formic acid. The bound peptides were then eluted onto a Waters BEH C18 nanoAcquity column (1.7 μm, 100 μm by 100 mm) and eluted over 35 min with a gradient of 2% B to 35% B for 21 min, 90% B from 21 to 24 min and 5% B after 24.1 min at a flow rate of 0.6 μL/min. Separations were done using a Waters nanoAcquity UPLC (buffer A, 99.9% water-0.1% formic acid; buffer B, 99.9% acetonitrile-0.1% formic acid), and the separated peptides were sprayed into a Thermo Fisher Q Exactive orbitrap (Waltham, MA) LTQ mass spectrometer by using a Michrom Advance nanospray source. Survey scans were taken in the ion trap, and the top five ions from each survey scan were automatically selected for high-resolution zoom scans followed by low-energy collision-induced dissociation (CID). The resulting MS/MS spectra were converted to peak lists by the use of BioWorks browser version 3.3.1 (Thermo Fisher Scientific) using the default parameters and searched against the Chlamydomonas reinhardtii database, version 3.0, downloaded from the DOE Joint Genome Institute (http://www.jgi.doe.gov), using the Mascot searching algorithm, version 2.1 (Matrix Science). The Mascot output was then analyzed using Scaffold (Proteome Software) to probabilistically validate protein identifications by using the ProteinProphet computer algorithm (27). Assignments validated above the Scaffold 95% confidence filter were considered true. Mascot parameters for database searching allowed up to two missed tryptic sites, variable modification of oxidation of methionine and carbamidomethyl cysteine, a peptide tolerance of ±0.1 ppm, an MS/MS tolerance of 0.8 Da, and a peptide charge state limited to +2/+3.

Fatty acid and lipid analysis. Various cellular and subcellular fractions were extracted into methanol-chloroform-formic acid (2:1:0.1 [vol/vol/vol]) followed by phase separation by the addition of 1 M KCl and 0.2 M H3PO4. The organic phase was spotted onto amonium sulfate-imregnated Si250 silica plates, and TAG was separated with petroleum ether-diethyl ether-acetic acid (80:20:1 [vol/vol/vol]) and visualized by a potassium permanganate spray. Lipid droplets were delipidated and reconstituted with the same enzyme; a clone containing the desired orientation of the promoter region was amplified from the vector pMN24 (11) and P3 (5’-GGTTGTAAACGACGGCAGTGCC-3’) and ligated into Km-digested pBlueScript II KS+ (Stratagene, Agilent Technologies, Santa Clara, CA) after digestion with the same enzyme; a clone containing the desired orientation of the insert was identified by PCR amplification using P2 (5’-GGTTGTAACGACGGCAGTGCC-3’) and P3 (5’-TTAATACGACTCACTATAGGG-3’), the T7 sequencing primer, and the resulting plasmid was designated pEM1. The Chlamydomonas NIT1 promoter region was amplified from genomic DNA with primers P1 (5’-GTTGTTGATGATATTTTATTTTAAAAC-3’) and P2 (5’-GTTGTTGATGATATTTTATTTTAAAAC-3’), which were digested with the same enzyme; a clone containing the desired orientation of the insert was identified by PCR amplification using P2 (5’-CTGAGGAAACCCGACGCGTCC-3’) and P3 (5’-TTAATACGACTCACTATAGGG-3’), the T7 sequencing primer, and the resulting plasmid was designated pEM1. The Chlamydomonas NIT1 promoter region was amplified from genomic DNA with primers P1 (5’-GTTGTTGATGATATTTTATTTTAAAAC-3’) and P2 (5’-GTTGTTGATGATATTTTATTTTAAAAC-3’), which were digested with the same enzyme; a clone containing the desired orientation of the insert was identified by PCR amplification using P2 (5’-CTGAGGAAACCCGACGCGTCC-3’) and P3 (5’-TTAATACGACTCACTATAGGG-3’), the T7 sequencing primer, and the resulting plasmid was designated pEM1. 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digested pSP124S—a plasmid containing the ble selectable marker gene for
*Chlamydomonas* transformation.

To generate an MLDP RNAi construct, primers P8 (5′-GGGACTAGTCT
ggaaagcccTGAACAC-3′) and P9 (5′-CTTCCATGGGTTCGTTAGT
accttgccg-3′) were used to amplify a sense amplicon from genomic DNA,
while P10 (5′-CCCTCCTGACGAAACCTGAAACAC-3′) and
P11 (5′-GAAACATGGTGGTCCGTTGGAGAATCTT-3′) were used to
amplify the antisense amplicon from cDNA. The sense and antisense amplicons
were double digested with SpeI and PstI, respectively, and ligated into
SpeI/PstI-digested pBuilderII KS*. The formation of the triple ligation prod-
uct was confirmed by DNA sequencing, and the resulting hairpin insert was
excised by SpeI/PstI digestion and ligated into pNTPRO1 digested with the
same enzymes. The resulting plasmid was designated pMLDP-RNAi. The
*C. reinhardtii* dw15

Quantitative real-time reverse transcription-PCR (RT-PCR). Total RNA was
purified using the RNeasy plant minikit (Qiagen, Germantown, MA) including
the on-column DNase digestion. Five hundred nanograms of RNA was reverse
transcribed with the RETROscript kit (Ambion, Austin, TX) in a 10-
µl reaction

Melting curves were performed after PCR to confirm that the amplification
product was unique.

**Fluorescent, confocal, and electron microscopy analyses.** For confocal micros-
copy analysis, cells were resuspended in phosphate-buffered saline (PBS). Cells
were stained with Nile red at a final concentration of 2.5 µg ml⁻¹ (from a stock
of 50 µg ml⁻¹ in methanol), followed by a 10-min incubation in the dark. Stained
cells were then mixed 1:1 (vol/vol) with 2% low-temperature melting point
agarose kept at 38°C and applied to the microscope slide. After the solidification
of the agarose, images were captured using a Zeiss 510 Meta Confoco3 LSM
microscope (Zeiss, Maple Grove, MN). For neutral-lipid-specific detection of
Nile red fluorescence, the 488-nm argon laser was used in combination with a
560- to 615-nm filter; for combined chlorophyll autofluorescence and Nile red
fluorescence of polar lipids, a 647- to 753-nm filter was used. Samples were
viewed with a 63× oil immersion lens objective. Postacquisition image handling
was done with Zeiss AIB software. For fluorescence microscopy analysis, a Zeiss
Axio Imager M1 microscope was used. Nile red fluorescence in neutral lipids was
detected using a UV light source with a 500 ± 24 nm/542 ± 27 nm excitation/
emission filter set. For the measurement of apparent lipid droplet diameter,
micrographs were analyzed using Image J software (1). For electron microscopy
analysis, centrifuged cell pellets were processed as previously described (13), and
transmission electron micrographs were captured using a JEOL100 CXII instru-
ment (Japan Electron Optics Laboratories, Tokyo, Japan).

Quantitative real-time reverse transcription-PCR (RT-PCR). Total RNA was
purified using the RNeasy plant minikit (Qiagen, Germantown, MA) including
the on-column DNase digestion. Five hundred nanograms of RNA was reverse
transcribed with the RETROscript kit (Ambion, Austin, TX) in a 10-µl reaction
mix. Real-time PCR was performed on an ABI Prism 7000 (Applied Bio-
systems, Foster City, CA) using the SYBR green PCR master mix with an
equivalent input of 50 ng RNA and 0.3 µM of each primer (MLDP-F [5′-GGA
TGCCCTGACCAAGTTC-3′] and MLDP-R [5′-GGATGGACCAAGGAGTC
GT-3′], RACK1-F [5′-GACACACACACCCCATCATC-3′] and
RACK1-R [5′-A GACGGTACAGTTGACGAGC-3′]). In all experiments, expression of
MLDP was normalized to RACK1 (CBLP) expression, a gene commonly used for nor-
malization in *C. reinhardtii*. Analysis of RACK1 expression by the 2⁻₅₇C
method (20) resulted in the following fold changes (averages and standard
deviations are indicated) in expression when using the average threshold cycle
(Ct) value of cells grown in 10 mM NH₄Cl, and 1.14 ± 0.11, 1.79 ± 0.72, and 1.62 ± 0.28 for cells
switched to 0 mM N for 1, 2, and 3 days, respectively. The standard curve method
for quantification of mRNA abundance was used as previously described (19).

**RESULTS**

TAG accumulation following shift to N-limited medium. While *C. reinhardtii*
does not accumulate TAG under nutrient-
TABLE 1. Total fatty acid content of whole cells compared to fatty acids bound in TAGs isolated from +N cells and −N cells.

| Fatty acid | Whole cells | TAG |
|------------|-------------|-----|
|             | +N          | −N  | +N    | −N  |
| 16:0        | 22.1 ± 0.14 | 26.2 ± 0.70 | 51.3 ± 1.30 | 37.6 ± 3.99 |
| 16:1        | 3.4 ± 0.02  | 2.9 ± 0.01  | 5.4 ± 2.37  | 3.7 ± 0.54  |
| 16:2        | 1.2 ± 0.00  | 1.4 ± 0.06  | 1.6 ± 0.96  | 1.4 ± 0.17  |
| 16:3        | 4.5 ± 0.15  | 3.4 ± 0.25  | 13.9 ± 6.88 | 3.2 ± 0.27  |
| 18:0        | 2.4 ± 0.02  | 2.5 ± 0.06  | 0.8 ± 0.84  | 1.4 ± 0.17  |
| 18:1, 16:4  | 21.9 ± 0.13 | 21.2 ± 0.14 | 7.9 ± 2.46  | 22.1 ± 8.45 |
| 18:2        | 8.0 ± 0.02  | 9.9 ± 0.06  | 6.1 ± 3.33  | 13.6 ± 1.51 |
| 18:3<sup>a</sup>, 12, 15 | 6.7 ± 0.01  | 6.5 ± 0.12  | 4.6 ± 1.62  | 5.7 ± 0.63  |
| 18:4        | 27.9 ± 0.10 | 24.7 ± 0.13 | 7.3 ± 3.69  | 9.9 ± 1.20  |

* Averages of three replicates and the standard deviation are shown.

** +N indicates cells grown in medium containing 10 mM NH<sub>4</sub><sup>+</sup>; −N indicates cells switched to 0 mM NH<sub>4</sub><sup>+</sup> and grown for 1 day.

replete growth conditions, TAG has been reported to accumulate in this genus after N or phosphorus limitation (39). In order to optimize conditions for the isolation of lipid droplets from C. reinhardtii cells, the time course of N deprivation and its effect on TAG accumulation and lipid droplet formation was analyzed. The amount of fatty acid esterified in TAG steadily increased over time to approximately 12%, 45%, and then 65% of total cellular fatty acids 24, 48, and 72 h, respectively, after resuspending the cells in TAP medium lacking a source of N, as shown in Fig. 1A. Total fatty acid profiles of N-limited cells showed moderate changes in comparison to N-replete cells, with a slight increase in 16:0 (carbon number: number of double bonds) and a concomitant decrease in 18:3<sup>a</sup>, 12, 15 (superscript indicates the position of the double bond counting from the carboxyl end); TAG fatty acid profiles of N-replete and N-limited cells were more distinct, with a decrease in 16:0 and an increase in 18:1/16:4 in N-limited cells (Table 1). Total cellular fatty acids increased on a per-cell basis 48 and 72 h after imposing N-limiting conditions (Fig. 1A), indicating that de novo fatty acid synthesis contributes at least in part to TAG accumulation.

A time course of lipid droplet formation in N-deprived C. reinhardtii cells is shown in Fig. 1B. Using the lipid-specific fluorescent dye, Nile red, lipid droplet formation was observed by confocal microscopy, adjusting excitation and emission filters to predominately detect fluorescence from the neutral lipid fraction (10). The first lipid droplets were observed after 12 h of N deprivation in distinct locations within the cell, and prolonged exposure of the cells to 24 or 48 h of N deprivation caused an increase in size and number of lipid droplets. Similarly, transmission electron micrographs revealed several ultrastructural changes that occur over a 72-h time course of N limitation, including a reduction of stacked thylakoid membranes, accumulation of starch granules, and the appearance of lipid droplets (Fig. 2).

Properties of lipid droplets. Lipid droplets were prepared from cells that were deprived of N for 24 h and compared to isolated thylakoid membranes and a fraction highly enriched in eyespots—a chloroplast-localized, carotenoid- and lipid-rich organelle (33). Results of spectral analyses of the various subcellular fractions indicated the presence of carotenoids (Fig. 3A). These results were similar to what was previously observed for an enriched eyespot fraction (33), although when normalized on a per-fatty-acid basis, carotenoids were much less abundant in lipid droplets. The near absence of chlorophylls in isolated lipid droplets represented by the absorbance maximum at approximately 660 nm visible in the thylakoid fraction (Fig. 3A) suggested low contamination of the lipid droplets with thylakoid membranes. The ratios of TAG-esterified fatty acids to total fatty acids of lipid droplets and eye-
TABLE 2. Fatty acid content of whole organelles (eyespot and lipid droplet) compared to fatty acids bound in TAGs isolated from indicated organelles

| Fatty acid | Eyespot | Lipid droplet |
|------------|---------|---------------|
| Whole organelle | TAG | Whole organelle | TAG |
| 16:0 | 35.5 ± 0.67 | 67.0 ± 7.55 | 32.2 ± 1.24 | 41.8 ± 4.33 |
| 16:1 | 3.0 ± 0.01 | 4.5 ± 1.08 | 3.4 ± 0.37 | 3.1 ± 0.58 |
| 16:2 | 3.4 ± 0.02 | 3.0 ± 0.30 | 1.2 ± 0.15 | 0.9 ± 0.06 |
| 16:3 | 15.7 ± 0.18 | 7.8 ± 0.45 | 6.4 ± 3.03 | 5.1 ± 1.70 |
| 18:0 | 3.0 ± 0.01 | 1.9 ± 0.50 | 2.1 ± 0.87 | 1.5 ± 0.77 |
| 18:1, 16:4 | 18.4 ± 0.39 | 7.9 ± 1.99 | 27.2 ± 1.74 | 27.3 ± 2.86 |
| 18:2 | 10.9 ± 0.71 | 4.7 ± 1.77 | 10.7 ± 0.67 | 10.0 ± 1.67 |
| 18:3 | 5.6 ± 0.09 | 1.4 ± 0.57 | 7.7 ± 1.59 | 5.9 ± 2.62 |
| 18:4 | 3.3 ± 0.47 | 0.8 ± 0.41 | 6.8 ± 4.69 | 2.9 ± 0.34 |
| 18:4 | 1.1 ± 0.02 | 1.1 ± 0.47 | 2.3 ± 0.92 | 1.4 ± 0.38 |

* Organelles were isolated from cells that were switched to medium containing 0 mM NH₄⁺ and grown for 1 day. Two replicates were averaged for eyespot samples and three for lipid droplet samples. The averages and the standard deviation are shown.

spots were approximately 88% and 25%, respectively (Fig. 3B), corresponding to the higher relative membrane abundance in the eyespots. The TAG fatty acid profile of isolated lipid droplets was very similar to the profile of TAGs isolated from cells deprived of N for 1 day, whereas the TAG fatty acid profile of eyespots was quite distinct, being much more enriched with palmitic acid on a mole percentage basis (Table 2). Taken together, these results indicated that the lipid droplets were distinct from the lipid globules found in the eyespot fraction, with minimal contamination of thylakoid membranes, the predominant membrane fraction in *C. reinhardtii*.

Analysis of proteins from whole cells, thylakoids, the eyespots, and lipid droplets by SDS-PAGE followed by total protein staining (Fig. 3C) revealed a pattern for lipid droplets that was distinct from the other protein fractions, with one major protein of approximately 27 kDa, as well as several less-abundant proteins, which were specifically enriched in lipid droplets compared to thylakoids and the eyespots. The lipid droplet proteins were excised from the gel (as indicated in Fig. 3C) and subjected to MS for protein identification.

**Proteins identified in the lipid droplet fraction.** As summarized in Table S1 in the supplemental material, 259 proteins with two or more unique peptides were identified and fell into a broad range of protein functional groups (see Fig. S1A in the supplemental material). Sixteen proteins predicted to be involved in lipid metabolism, including three presumed acyl coenzyme A (acyl-CoA) synthetases and a predicted lipoygenase—proteins previously identified as associated with lipid droplets by studies of other organisms (6, 8)—were among those proteins. BT1A, the enzyme responsible for the synthesis of a major extraplastidic membrane lipid, diacylglycerol-β-trimethylhomoserine (30), was also present. Several predicted components of vesicular trafficking pathways were identified, including subunits of the coat protein complex I and its putative regulator, ARF1a, as well as other small Rab-type GTPases. Orthologs of these proteins have also been identified in lipid droplet proteomics studies done on different animal species (6, 8), and recent *Drosophila* cell RNAi inactivation studi
ties targeting Arf79a (ARF1a ortholog) and the coat protein complex I-encoding genes revealed a previously unknown role for these proteins in lipid droplet formation and utilization (7, 12). The presence of presumed orthologs in *C. reinhardtii* lipid droplets suggests the possibility of a similar role in algal lipid droplet biogenesis.

Other functional groups which were represented by many proteins included 54 predicted ribosomal and translation-related proteins, 26 predicted mitochondrial proteins, and 24 predicted proteins of the photosynthetic apparatus. Ribosomal and mitochondrial proteins have been identified in lipid droplet proteomic studies of different animal species (8, 40) and plant oil bodies (in plants from mitochondria only) (17), and could be common contaminating proteins in lipid droplet preparations or, alternatively, could indicate the presence of ribosomal/mitochondrial-lipid droplet physical interactions, as has been previously postulated (8). The identification of 24 photosynthetic apparatus proteins was surprising, given the near absence of chlorophyll in purified lipid droplets, and it is likely that these hydrophobic apoproteins are contaminating artifacts that become associated during the isolation procedure; however, the most abundant stromal enzyme, ribulose-1,5-bisphosphate carboxylase/oxygenase, was not detected. A more stringent analysis of 93 proteins (identified by more than 12 spectral counts) resulted in an ~38% decrease in the number of lipid metabolism proteins retained in the “identified” group (from 16 to 10), whereas relative decreases in the more predominant contaminating groups were generally higher (~78% for ribosome/DNA/translation-related proteins, ~69% for mitochondrial proteins, and ~50% for photosynthetic apparatus proteins [see Fig. S1A and S1B in the supplemental material]). In addition, 45 proteins of unknown function were identified. Based on spectral counts for all unique peptides for a given protein (see Table S2 in the supplemental material), one of these was, by a factor of 10, the most abundant protein in gel slice 9 (Fig. 3C), which also contained the most abundant protein in the lipid droplet fraction based on staining. Its calculated molecular mass is 28 kDa, which approximately corresponded to its position in the gel (Fig. 3C). Therefore, this protein was presumed to be the most abundant protein in the lipid droplet fraction and was tentatively designated the major lipid droplet protein (MLDP; protein identification no. 192823 in the *C. reinhardtii* v.3.0 database [http://genome.jgi-psf.org/Chlr3/Chlr3.home.html]; NCBI accession number XP_001697668). Notably, the second-most abundant protein based on spectral counts was BT1A, the enzyme catalyzing the biosynthesis of betaine lipid. Both proteins were also identified in studies of *C. reinhardtii* lipid droplets conducted in parallel but were dismissed as contamination (41).

**MLDP suppression affects lipid droplet size.** Steady-state mRNA abundance of MLDP was determined by quantitative real-time RT-PCR in cells grown in 10 mM N and cells deprived of N for 24, 48, and 72 h as shown in Fig. 4A. The widely used *C. reinhardtii* control gene *RACK1* (*BLP*) was used as an internal control. Analysis of apparent fold change in *RACK1* expression by the 2⁻ΔΔCt method (see Materials and Methods) indicated that changes in its expression based on total RNA input to the reverse transcriptase reaction were minor if there were any (~1.5-fold increases at 48 and 72 h with no N).
Previous studies have shown that during N deprivation in *C. reinhardtii*, the amount of chloroplast and cytoplasmic ribosomes and total RNA content decrease (24). While this can potentially add an additional level of difficulty in identifying a suitable control gene in these experiments, the degree to which this contributes to the apparent minor fold changes in expression for *RACK1* is unknown. With the above in mind, *RACK1* was deemed a suitable control gene for these experiments, with the caveat that it may slightly underestimate the fold changes for genes induced by 48 or 72 h of N deprivation, and vice versa for downregulated transcripts. When normalized to *RACK1*, *MLDP* mRNA showed a 16-fold increase in abundance after 24 h of N deprivation and continued to be abundant even after 72 h of N deprivation. Moreover, *MLDP* mRNA abundance qualitatively corresponded during a time course of N deprivation with the accumulation of TAG (Fig. 1A). The large increase in *MLDP* transcript abundance in N-deprived cells was further corroborated by Northern blotting (Fig. 5).

The *MLDP*-RNAi lines were analyzed for possible lipid droplet morphological phenotypes by fluorescence microscopy analysis of Nile red-stained cells deprived of N for 3 days. An apparent increase in the average size of lipid droplets in the *MLDP*-RNAi lines was observed (Fig. 4D and E).

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ent individual diameters of a population of >200 random and in-focus lipid droplets per RNAi or control line were measured after 3 days of N deprivation (Fig. 4D). An approximately 40% increase in the average lipid droplet diameter was determined in the MLDP-RNAi lines.

MLDP has orthologs only in green algae. The newly discovered MLDP protein has few orthologs, none of which have been functionally characterized. When searched against the National Center for Biological Information database of non-redundant proteins by the use of BLAST (3), no sequence with significant similarity is identified. However, several genome projects of green algal species are currently under way, and by searching against green algal genomes at the respective websites, potential orthologs with high sequence identity were identified in Chlorella sp. NC64A, Chlorella vulgaris, and Volvox carteri (Fig. 6). No discernible ortholog was present in available diatom or red algal genomes or the genomes of Ostreococcus species, which belong to a green algal lineage distinct from C. reinhardtii. Apparently, MLDP is a rare protein in nature as we currently know it at the molecular level and presumably arose during the evolution of green algae. As such, it also has no discernible domains to which functions have been assigned.

Grand average of hydrophathy (GRAVY) indices and Kyte-Doolittle (18) hydroPATHY plots were generated for MLDP, as well as for mouse ADRP and perilipin, and Arabidopsis oleosin 1 (OLE1) (Fig. 6B to E). All four proteins were distinct with respect to extent and region of hydrophobicity, with the most hydrophobic region for MLDP occurring approximately between amino acid residues 160 and 215. The proteins ranked according to their GRAVY index (highest to lowest) are MLDP (0.11), OLE1 (−0.17), ADRP (−0.28), and perilipin (−0.40). Thus, MLDP is a very hydrophobic protein although it lacks a specific hydrophobic core characteristic for oleosin.

DISCUSSION

The potential of microalgae to provide sustainable feedstock for biofuel production has led to renewed interest in the biology of this diverse group of organisms (14). While C. reinhardtii is unlikely to become the microalga of choice for the production of biofuels, it is a well-established genetic and genomic model for the study of unicellular green algae (25). However, very little is known about lipid metabolism or the cell biology of lipid droplets and its regulation in this organism. The time course and conditions for the induction of oil accumulation described here (Fig. 1 and 2) and the detailed analysis of lipid droplets provide the basis for a molecular analysis of the mechanisms of TAG accumulation in C. reinhardtii and possibly unicellular green algae in general. As a logical first step, we focused on the TAG-storing compartment itself, the lipid droplet. Its fatty acid and pigment composition is clearly distinct from that of other organelles (Fig. 3). This distinct composition and the very low chlorophyll content in view of the predominance of thylakoid membranes in the cell suggested that the obtained oil droplet preparation was only slightly contaminated with thylakoid membranes, the largest membrane system in the cell. However, we cannot rule out some level of contamination with ER-derived microsomes especially since lipid droplets are generally thought to be derived from the ER. An increase in the total amount of fatty acids in N-deprived cells indicated that the biosynthesis of TAGs is not merely due to conversion of membrane lipids but the result of de novo synthesis. The specific fatty acid composition of lipid droplet lipids tended toward shorter and more saturated fatty acids, suggesting that newly synthesized fatty acids were quickly incorporated into TAGs before fatty acid modification by desaturases or elongases could occur; alternatively, reduced activities of these fatty acid metabolic enzymes could account for this observation.

Lipid droplets in different organisms are increasingly recognized as dynamic organelles that provide important nodes of cellular metabolic networks and participate in intracellular lipid trafficking (e.g., see reference 12). In algae, the induction of lipid droplet formation is intricately regulated by nutrient deprivation and other abiotic stresses that lead to a cessation of cell division or an overreduction of the photosynthetic electron transport chain (14). In this study, we sought to identify the proteins associated with lipid droplets to determine factors involved in the biogenesis or turnover of this organelle in C. reinhardtii. Aside from those of unknown function, of particular interest are lipid metabolic enzymes and proteins involved in vesicular transport. Intriguingly, the betaine lipid (diaclylglycerol-<i>N</i>,<i>N</i>,<i>N</i>-trimethylhomoserine) biosynthetic enzyme, BTA1 (30), had the second highest spectral count of proteins in the lipid droplet fraction. This protein was also identified as one of few in the lipid droplet fraction in a parallel study but was considered a contamination (41). This parallel study concluded that no protein was specifically associated with the respective lipid droplet fraction prepared and treated differently than in this study.

Here, a number of predicted acyl-CoA synthetases and acyltransferases were present as well that could be involved in the biosynthesis of TAGs or the transfer of acyl groups between membrane lipid and neutral lipid pools. In animal cells, enzymes involved in phosphatidylcholine biosynthesis have been found to be critical for lipid droplet formation and turnover. Lipid droplets in animal cells are surrounded by a monolayer of phospholipids, and the availability of those phospholipids affects lipid droplet size. When the rate-liming enzyme of phosphatidylcholine biosynthesis was inactivated in Drosophila cells, lipid droplets increased in size presumably because the larger droplets have a lower surface area-to-volume ratio and require less phospholipids to form the phospholipid monolayer (12). As C. reinhardtii lacks phosphatidylcholine, it seems plausible that its role in the formation of lipid droplets is taken by betaine lipid, which has similar properties and is thought to functionally replace phosphatidylcholine in this green alga (32). A rich complement of proteins predicted to be involved in vesicular trafficking, including COP1 homologs, Rab GTPase, or ARF-related GTPase, were also identified in the proteomics data set. Repression of the respective orthologs in Drosophila cells was shown to affect lipid droplet formation and dynamics and TAG utilization (12). Our understanding of the interaction of lipid droplets with the vesicular transport machinery is only rudimentary at this time. The presence of the respective proteins in C. reinhardtii follows a common theme also observed for animal cells and provides an opportunity to study the role of the vesicular transport machinery in lipid droplet formation in a unicellular model organism. However, the specific associ-
ation of these proteins with lipid droplets in *C. reinhardtii* remains to be confirmed.

The identification of proteins in the ribosome-/translation-/DNA-related, mitochondrial, and photosynthetic apparatus functional groups suggests a basic level of contamination in our lipid droplet protein fraction. While more-stringent filtering of the proteomics data set to include proteins with more than 12 spectral counts resulted in a larger decrease in the relative proportion of these apparent contaminating functional groups, compared to lipid metabolism proteins (see Fig. S1B and Table 104 MOELLERING AND BENNING EUKARYOT. CELL

**FIG. 6.** Multiple sequence alignment of MLDP with other green algal orthologs and hydropathy plots of MLDP and other known lipid droplet binding proteins. (A) MLDP and putative orthologs identified in the *Chlorella* sp. NC64A (http://genome.jgi-psf.org/ChlNC64A_1/ChlNC64A_1.home.html), *Chlorella vulgaris* C-109 (http://genome.jgi-psf.org/Chlvu1/Chlvu1.home.html), and *Volvox carteri* (http://genome.jgi-psf.org/Volca1/Volca1.home.html) genomes. Conserved residues are indicated with asterisks, and the coloring indicates the CORE index, where red and blue indicate the highest and lowest probability of the alignment being correct at a given residue, respectively, with colors in between the spectrum of red and blue representing a range of intermediate probabilities. Hydropathy plots for the proteins (followed by NCBI accession numbers in parentheses) MLDP (XP_001697668) (B), mouse ADRP (NP_031434) (C), mouse perilipin (NP_783571) (D), and *Arabidopsis* OLE1 (NP_194244) (E).
S2 in the supplemental material), the significance of such analyses should be regarded as circumstantial, given the potential for protein size bias in the method. Comparing the results of the analysis of the lipid droplet proteins presented here with proteomic studies of the eyespot (33), flagella (28), and mitochondria (5) (see Table S1 in the supplemental material) reveals that some degree of contamination with proteins from specific functional groups is common (e.g., ribosome-/translational-/DNA-related, mitochondrial, and photosynthetic apparatus proteins found in the eyespot study, or ribosome contamination in the mitochondrial and flagellar studies). As such, apparent contamination of subcellular preparations in *C. reinhardtii* seems to be a common challenge.

To begin the functional analysis of lipid droplet-associated proteins in *C. reinhardtii*, we focused on the most abundant protein in the lipid droplet fraction based on Coomassie brilliant blue staining (Fig. 3C) and spectral counts (see Table S2 in the supplemental material). The identity of this protein was deduced by MS analysis of all proteins in the respective gel slice, and MLDP was identified based on 10-fold overabundance of protein in the lipid droplet fraction based on Coomassie brilliant blue staining. Detection of MLDP was based on 10-fold overabundance of specific functional groups is common (e.g., ribosome-/translational-/DNA-related, mitochondrial, and photosynthetic apparatus proteins found in the eyespot study, or ribosome contamination in the mitochondrial and flagellar studies). As such, apparent contamination of subcellular preparations in *C. reinhardtii* seems to be a common challenge.

The newly discovered MLDP is a rare protein in nature unique to green algae. The expression of its gene is strictly limited to conditions favoring TAG biosynthesis, providing indirect evidence for a role of MLDP in lipid droplet formation or maintenance. These properties potentially make MLDP a marker for lipid droplets and TAG accumulation.

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