A complex system of small RNAs in the unicellular green alga *Chlamydomonas reinhardtii*

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Endogenous small RNAs function in RNA interference (RNAi) pathways to control gene expression through mRNA cleavage, translational repression, or chromatin modification. Plants and animals contain many microRNAs (miRNAs) that play vital roles in development, including helping to specify cell type and tissue identity. To date, no miRNAs have been reported in unicellular organisms. Here we show that *Chlamydomonas reinhardtii*, a unicellular green alga, encodes many miRNAs. We also show that a *Chlamydomonas* miRNA can direct the cleavage of its target mRNA in vivo and in vitro. We further show that the expression of some miRNAs/Candidates increases or decreases during *Chlamydomonas* gametogenesis. In addition to miRNAs, *Chlamydomonas* harbors other types of small RNAs including phased small interfering RNAs (siRNAs) that are reminiscent of plant *trans*-acting siRNAs, as well as siRNAs originating from protein-coding genes and transposons. Our findings suggest that the miRNA pathway and some siRNA pathways are ancient mechanisms of gene regulation that evolved prior to the emergence of multicellularity.

[Keywords: *Chlamydomonas*, miRNA, mRNA cleavage, small RNA, RNAi, RISC]

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In eukaryotes, small RNAs of ~20–30 nucleotides (nt) function as guides for effector complexes [RNA-induced silencing complexes, RISCs] that can regulate gene expression at the transcriptional or post-transcriptional level. Small RNAs are produced by the RNase III enzyme Dicer or Dicer-like proteins (DCLs) from their precursors, which can be stem–loop RNA transcripts or long double-stranded RNAs (dsRNAs). These small RNAs directly interact with proteins in the Piwi/Argonaute (AGO) family to form the core of RISC (Hannon 2002; Meister and Tuschl 2004).

In multicellular systems, three major classes of endogenous small RNAs have been discovered: microRNAs (miRNAs), Piwi-associated small RNAs [piRNAs], and small interfering RNAs [siRNAs]. miRNAs are processed from RNA transcripts that form stem–loop structures. They regulate gene expression through mRNA cleavage or translational repression based on the degree of sequence complementarity between miRNAs and their target mRNAs, many of which have demonstrated regulatory roles in development and other physiological processes (Carrington and Ambros 2003; Bartel 2004). piRNAs are a newly discovered class of small RNAs that are associated with Piwi family proteins (Aravin et al. 2006; Girard et al. 2006; Grivna et al. 2006; Lau et al. 2006). The biogenesis machinery and biological roles of piRNAs remain largely unknown. siRNAs are produced from long dsRNAs that originate from various sources including products of bidirectional transcription or RNA-dependent RNA polymerases (RDRs). siRNAs play various roles in post-transcriptional regulation of gene expression in *cis* or in *trans* (Vaucheret 2006), suppression of transposable elements, and the formation of heterochromatin (Lippman and Martienssen 2004).

Based on their origins and functions, siRNAs can be further classified into several subclasses: heterochromatic siRNAs [hc-siRNAs], repeat-associated siRNAs [ra-siRNAs], *trans*-acting siRNAs [ta-siRNAs], and natural antisense transcript-derived siRNAs [nat-siRNAs]. In *Arabidopsis*, there is an abundant class of DCL3 and
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RDR2-dependent ~24-nt hc-siRNAs that act through AGO4 to maintain DNA/histone methylation of transposons and repeats (Hamilton et al. 2002; Zilberman et al. 2003; Xie et al. 2004; Qi et al. 2006). Similarly, ra-siRNAs silence retrotransposons and repetitive sequences in the Drosophila germ line, but their biogenesis does not seem to require Dicer (Aravin et al. 2003; Vagin et al. 2006). ta-siRNAs have been found only in plants (Peragine et al. 2004; Vazquez et al. 2004; Allen et al. 2005; Axtell et al. 2006; Talmor-Neiman et al. 2006). Their production is initiated by the miRNA-mediated cleavage of their precursor transcripts (Peragine et al. 2004; Vazquez et al. 2004; Allen et al. 2005). The cleaved transcripts serve as substrates for RDR6 to make dsRNAs that are in turn processed by DCL4 (Gasciolli et al. 2005; Xie et al. 2005; Yoshikawa et al. 2005). It was recently shown that double miRNA complementary sites are required for efficient ta-siRNA production (Axtell et al. 2006). Like miRNAs, ta-siRNAs function in trans to guide the cleavage of endogenous mRNAs, and have been shown to regulate vegetative phase changes in Arabidopsis (Peragine et al. 2004; Vazquez et al. 2004; Allen et al. 2005; Gasciolli et al. 2005; Xie et al. 2005; Yoshikawa et al. 2005) and in moss (Talmor-Neiman et al. 2006). nat-siRNAs have also been found in Arabidopsis, where they are derived from natural antisense transcripts. dsRNAs formed by the transcription of antisense overlapping gene pairs are processed by DCL2 into 24-nt nat-siRNAs, which direct the further production of 21-nt nat-siRNAs in a DCL1-dependent fashion (Borsani et al. 2005).

In contrast to the complex small RNA pathways found in multicellular systems, small RNA pathways in unicellular organisms appear, so far, to be relatively simple. In fission yeast, a single class of siRNAs exists in the RNA-induced transcriptional silencing complex [RITS]. These siRNAs have demonstrated roles in epigenetic silencing at centromeres and the initiation of heterochromatin assembly at the mat locus (Lippman and Martienssen 2004). The ciliated protozoan Tetrahymena thermophila has ~26- to 31-nt scan RNAs [scnRNAs] that are involved in developmentally regulated DNA elimination (Mochizuki and Gorovsky 2004a,b; Yao and Chao 2005), and ~23- to 24-nt small RNAs with suggested roles in gene regulation at the post-transcriptional level (Lee and Collins 2006). Thus far, no miRNAs have been identified in unicellular systems, raising the question whether miRNAs exist only in multicellular systems, where regulation of cell differentiation, developmental patterning, and timing is important. Indeed, it has been suggested that miRNAs helped to drive the evolution of a multicellular state (Bartel 2004).

With the goal of probing the diversity of small RNA pathways in unicellular eukaryotes, we turned to Chlamydomonas reinhardtii. Chlamydomonas is a unicellular green alga with well-understood haploid genetics. It also occupies a unique niche in evolution. It has served as a model system to understand various aspects of cell biology including the assembly and functions of the basal body, chloroplast, and flagellum (Lefebvre and Silflow 1999). The completion of the Chlamydomonas genome sequence and substantial expression sequence tag (EST) projects has greatly increased its utility as a model system (Grossman et al. 2003; Shroger et al. 2003).

Chlamydomonas possesses a functional RNA interference [RNAi] machinery, as evident from the observation that the target gene expression can be specifically knocked down in transgenic lines expressing corresponding antisense RNAs or dsRNAs (Schroda 2006). In accord with these observations, bioinformatics analysis predicted that the Chlamydomonas genome encodes key RNAi components including three Dicer-like proteins and at least two Argonautes [Supplementary Fig. 1; Schroda 2006]. The complexity of RNAi components in Chlamydomonas strongly suggested the existence of multiple small RNA pathways. However, it remains unknown whether Chlamydomonas encodes endogenous small RNAs and whether such small RNAs have regulatory roles in gene expression.

To address the aforementioned questions, we determined a large number of small RNA sequences from Chlamydomonas using a highly parallel pyrosequencing technology (Margulies et al. 2005). We found that Chlamydomonas encodes miRNAs, as well as other small RNAs derived from various genomic regions that include protein-coding genes and transposable and repetitive sequences. We also found that, like plant miRNAs, Chlamydomonas miRNAs can guide the cleavage of their target mRNAs. We further show that the accumulation of some miRNAs was either elevated or decreased in gametic cells. Our findings not only reveal the existence of multiple RNAi pathways in a unicellular eukaryotic organism, but also provide a rich resource for studying the roles of small RNAs in basic cell biology and the evolutionary transition from unicellularity to multicellularity.

**Results**

**Identification of Chlamydomonas small RNAs**

To examine whether Chlamydomonas encodes small RNAs, we prepared total RNA from Chlamydomonas cultures under vegetative growth conditions. After removal of larger RNAs by polyethylene glycol [PEG] precipitation, RNAs were separated by denaturing polyacrylamide gel electrophoresis and visualized by SYBR-gold staining. We observed an abundant class of small RNAs, which we cloned and analyzed further as described below. The sequences matched the chloroplast and mitochondrial genomes, respectively (Table 1). These sequences were analyzed further as described below. The sequences
without perfect genomic matches might arise from unsequenced regions of the genomes or from sequencing errors and were not pursued further.

The 82,350 small RNAs with perfect nuclear genomic matches (the redundant data set) represent 4182 unique sequences (the unique data set) [Table 1]. Of these, 2812 small RNAs were cloned only once, indicating that the small RNA population in *Chlamydomonas* is complex.

**Figure 1.** A catalog of *Chlamydomonas* endogenous small RNAs. (A) SYBR-gold was used to visualize small RNAs in total RNA extracted from *Chlamydomonas* cultures. A 21-nt synthetic RNA oligo was used as a size reference. (B) Size distribution of *Chlamydomonas* small RNAs. The sets of redundant [red] and unique [blue] small RNAs were used to generate a histogram quantifying the number of sequences obtained for each size class. (C) Sequence composition of the 5’ ends of the small RNAs. (D) Genome-wide density analysis of the small RNAs on an artificial *Chlamydomonas* genome assembled by linked scaffolds. The number of small RNAs with perfect matches in either a direct or complementary strand within each 50-kb sliding window was plotted. A small RNA production hot spot in Scaffold_58 is shown. The black bar represents the 310-nt hot spot region. Short thin lines above the black bar represent small RNAs derived from the sense strand, and lines below the bar represent small RNAs from the antisense strand. Red lines represent small RNAs that are mapped uniquely to this hot spot, while green lines represent small RNAs that are mapped to multiple genomic sites including the hot spot.
Chlamydomonas reinhardtii miRNAs have a set of small RNAs matching different genomic loci, prompting us to examine whether intergenic and intronic regions should have a set of small RNAs (Lu et al. 2005). The two small RNAs with exceptionally high cloning frequency were found in the NCBI database. The two small RNAs start sites and differed only by one nucleotide at their 3' ends. No identifiable homologs of these two small RNAs were derived from the same genomic loci with identical cloning frequency. One with a size of 22 nt (UUGUGCCGGCCGACACUGCGG) was cloned 54,353 times, the other with a size of 21 nt (UUGUGC) was cloned 5,786 times. These two small RNAs matched four genomic loci with one located in an exon of a protein-coding gene of unknown function and the other three within intergenic regions. Two of the intergenic loci could also produce a 23-nt-long small RNA, whose sequence is reverse-complementary to the two small RNAs but has additional nucleotides at both ends. The cloning frequency of this small RNA ranked third among all small RNAs (cloned 1694 times) (data not shown). The exceptionally high transcription rate may suggest the unique functional significance of these small RNAs.

### miRNAs in Chlamydomonas

The identification of a large number of small RNAs in *Chlamydomonas* prompted us to examine whether some of them were miRNAs. We selected small RNAs originating from intergenic or intronic regions to examine whether their surrounding sequences could form hairpin secondary structures, a distinguishing feature of known plant and animal miRNAs.

Consistent with the SYBR-gold staining result, the sizes of the small RNAs range from 17 nt to 26 nt, with 91.5% of small RNAs in the redundant data set and 72.2% in the unique data set being 20–22 nt in length [Fig. 1B; Supplementary Table 1]. In both the unique and redundant small RNA data sets, a preference for a U was observed at the 5' end of small RNA sequences [Fig. 1C]. A similar 5' nucleotide bias has been noted in small RNAs from other organisms (Aravin et al. 2003; Bartel 2004; Girard et al. 2006). Among the 4182 unique small RNAs mapped to the genome, 2247 sequences match only one site in the genome, and 781 sequences match twice. The remaining sequences match the genome from several to a few hundred times, with one exception that has 1152 genomic matches. These small RNA-generating regions include intergenic regions, annotated protein-coding genes, repetitive sequences, and regions producing noncoding RNAs (tRNAs or rRNAs) [Table 1]. The number of unique small RNAs matching the intergenic regions exceeds the number of those matching genes and repeats, similar to what has been observed for *Arabidopsis* small RNAs (Lu et al. 2005).

We summed the number of unique small RNAs in a 50-kb sliding window and plotted it against the artificial genome assembled from 1266 scaffolds. A few hot spots for small RNA production were observed [Fig. 1D]. One such hot spot included 427 small RNAs within a 310-nt intergenic region [Fig. 1D]. The entire region was covered by small RNAs on both strands. In many cases, multiple small RNAs with identical start sites but different lengths were cloned from the same position.

We found two small RNAs with exceptionally high cloning frequency. One with a size of 21 nt [UUGUGC CGGCCGCACAGCGCCG] was cloned 54,353 times, the other with a size of 22 nt [UUGUGCCCGCCGACACU GCGG] was cloned 5,786 times. These two small RNAs were derived from the same genomic loci with identical start sites and differed only by one nucleotide at their 3' ends. No identifiable homologs of these two small RNAs were found in the NCBI database. The two small RNAs matched four genomic loci with one located in an exon of a protein-coding gene of unknown function and the other three within intergenic regions. Two of the intergenic loci could also produce a 23-nt-long small RNA, whose sequence is reverse-complementary to the two small RNAs but has additional nucleotides at both ends. The cloning frequency of this small RNA ranked third among all small RNAs (cloned 1694 times) (data not shown). The exceptionally high transcription rate may suggest the unique functional significance of these small RNAs.

### Table 1. Chlamydomonas small RNAs matching different categories of sequences

| Categories                          | Redundant | Unique |
|-------------------------------------|-----------|--------|
| Nuclear genome                      | 82,350    | 4182   |
| Protein-coding genes only           | 1985      | 742    |
| Exons                               | 1702      | 629    |
| Introns                             | 278       | 108    |
| Exons and introns                   | 5         | 5      |
| Antisense to protein-coding genes only| 3563      | 535    |
| Intergenic regions only             | 8066      | 1712   |
| Protein-coding genes and intronic regions | 66,751    | 1193   |
| Repeats                             | 278       | 167    |
| Noncoding RNAs (tRNAs or rRNAs)    | 37        | 27     |
| Antisense to noncoding RNAs         | 1         | 1      |
| Chloroplast genome                  | 116       | 65     |
| Mitochondia genome                  | 20        | 14     |

The plant and animal miRNAs contain a phosphate at the 5' end. The small RNA cloning procedure used has a step of ligation of an RNA adapter to the 5' end of small RNAs by T4 RNA ligase, a process that requires a 5' phosphate [Lau et al. 2001]. Thus, the miRNAs we have cloned from *Chlamydomonas* should have a 5' phosphate. We used enzymatic methods to further confirm that the *Chlamydomonas* miRNAs have a 5' phosphate. The small RNAs isolated from *Chlamydomonas* cells were subjected to calf intestinal phosphatase (CIP) treatment followed by treatment with polynucleotide kinase in the presence of ATP. The CIP treatment caused miR912 to have a lower mobility in polyacrylamide gel electrophoresis.
| miRNA | Sequence (5’/3’) | Len | Start | End | Strand | Intergenic or Intronic | Predicted target |
|-------|-----------------|-----|-------|-----|--------|------------------------|-----------------|
| miR905 | UCCGAGGCGAUAUUUUCUU | 21 | 4 | 10125326 | + | Intergenic | jgi|Chlre3.178355/Chlre2.37.C_scaffold.21001903 |
| miR906-3p | UCCGAUAAAGCUUCCCCCUGC | 21 | 22 | 186500 | + | Intronic | jgi|Chlre3.148620/Chlre2.37.C_scaffold.21000500 |
| miR906-5p | CGGUUGGUGGGCGUGAUCAGC | 21 | 22 | 186416 | + | Intronic | jgi|Chlre3.148620/Chlre2.37.C_scaffold.21000500 |
| miR907 | UCUUCUGCGAGCGGUGCGAGC | 21 | 7 | 78141466 | + | Intergenic | jgi|Chlre3.173855/Chlre2.37.C_scaffold.21001903 |
| miR908 | ACGCGUUUGAUAGCAGGAUC | 20 | 4 | 112304014 | + | Intronic | jgi|Chlre3.173855/Chlre2.37.C_scaffold.21001903 |
| miR909-3p | UUCAGGGUCAAGUUUGCAUGC | 21 | 1 | 84678492 | + | Intergenic | jgi|Chlre3.173855/Chlre2.37.C_scaffold.21001903 |
| miR909-5p | AUGCAAACAUGACCCUGAAUG | 21 | 1 | 84678206 | + | Intergenic | jgi|Chlre3.173855/Chlre2.37.C_scaffold.21001903 |
| miR910 | AGCAGCGUCGGGCUCGACCGC | 21 | 17 | 46319955 | + | Intronic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR911 | ACAAUGGUAGUACGGCCAUUU | 21 | 8 | 23356104 | − | Intergenic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR912 | UGGGAUUGAUCCCAGCCAGGCG | 21 | 12 | 75416949 | − | Intronic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR913 | ACGACAGCGACGGAGGGCGAGC | 21 | 4 | 23356104 | − | Intergenic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR914 | UGGGAUUGAUCCCAGCCAGGCG | 21 | 12 | 75416949 | − | Intronic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR915 | GGCGCGCGCCGGAAUCCGUGG | 21 | 3 | 22482189 | − | Intronic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR916 | UGGCAAUAAGGCAAUCGUUGC | 21 | 7 | 58303884 | − | Intergenic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR917 | CCAAGGUCAUCGGUUCGACUCC | 21 | 4 | 104739429 | − | Intronic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR918 | UUGGUUGAUCCCAGCCAGGCG | 21 | 12 | 75416949 | − | Intronic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR919.3p.1 | UACCCUCCAGCAGCUUGCUAC | 21 | 13 | 20656080 | − | Intergenic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR919-3p.2 | UAGUUGAUCCCAGCCAGGCG | 21 | 58 | 20656080 | − | Intergenic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR919-5p.1 | UCUCAGGAGGACAUCGCCACU | 21 | 9 | 20656144 | − | Intronic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR919-5p.2 | AGCAAGCUACUUCAGGGUAUC | 21 | 14 | 20656120 | − | Intronic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |
| miR919-5p.3 | CAAGAUGUCCGCUUCCAGGUAC | 21 | 24 | 20656079 | − | Intronic | jgi|Chlre3.181462/Chlre2.37.C_scaffold.21002223 |

Length of mature miRNA sequence.

Starting position of pre-miRNA in the assembled artificial Chlamydomonas genome.

Predicted target sites. The mispairing score is given in parentheses. A complete list of predicted targets, with associated target site alignments and scores, is given in Supplementary Database 3.
Supplementary Fig. 2), suggesting that the miRNA had one or more phosphates at the 5’ end. Subsequent treatment with polynucleotide kinase and ATP restored the original mobility (Supplementary Fig. 2). This indicates that the miRNA contains one phosphate group at the 5’ end, like plant and animal miRNAs.

Unlike plant miRNA genes that are mostly located in intergenic regions [Jones-Rhoades et al. 2006], a considerable proportion of the Chlamydomonas miRNA/Candidate genes is intronic, while the remaining miRNA genes reside in intergenic regions [Table 2, Supplementary Table 2]. Unlike instances of polycistronic microRNAs in other systems, where multiple cotranscribed stem–loops give rise to independent microRNAs, we found that multiple Chlamydomonas miRNAs/Candidates can be derived from a single stem–loop (Fig. 2A; Supplementary Database 1). For example, miR906-5p and miR906-3p as well as miR919-3p.1, miR919-3p.2, miR919-5p.1, miR919-5p.2, and miR919-5p.3 are produced from the same precursors [Fig. 2A]. We also found that some miRNA genes are clustered in the genome, likely representing more conventional microRNAs polycistrons. One of the clusters [Cluster 4] only contains miRNAs in the same family [Supplementary Table 3]. These two genes [Candidate160c and Candidate160d] are nearly identical, suggesting that they were recently evolved through duplication [Supplementary Database 1].

The majority of the miRNAs/Candidates have a single precursor, whereas others have more than one precursor [Fig. 2, Supplementary Table 2]. For most miRNAs with multiple precursors, the mature miRNAs are located on the same arm of the stem–loops, suggesting that these genes might share a common ancestry [Fig. 2B, Supplementary Database 1]. We also identified some miRNAs that are located on either the 5’ or the 3’ arm of different

Figure 2. Representative miRNAs in Chlamydomonas. (A–C) Predicted fold-back structures of selected miRNA precursors. Sequences corresponding to the mature miRNAs are shown in red, and sequences corresponding to the miRNA*s are shown in blue. Protruding stem–loops in the 3’ arm of the hairpins are indicated by slashes (\). (D) Confirmation of miRNA expression by Northern blot using end-labeled oligonucleotide probes as indicated. A synthetic 32P-labeled 21-nt RNA oligo was used as a size marker.

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precursors [Fig. 2C; Supplementary Database 1], suggesting that genes encoding a particular miRNA could have evolved from more than one ancestor.

To confirm expression of some of our miRNAs/Candidates, we performed Northern analysis on RNAs prepared from Chlamydomonas vegetative cells. Many of the miRNAs/Candidates were readily detectable [Fig. 2D], whereas some are likely expressed at lower levels and were below the detection limit of our Northern blot protocol [data not shown].

Are miRNAs conserved in green algae?

Land plants are thought to be descendants of green algae-like ancestors (Lewis and McCourt 2004; Palmer et al. 2004). It was recently shown that some miRNAs are conserved in land plants and have ancient origins (Axtell and Bartel 2005). The existence of miRNAs in the green alga Chlamydomonas raised the possibility that plants and Chlamydomonas might share some common miRNAs. We compared Chlamydomonas miRNAs with all known plant [and also animal] miRNAs, and found no homologs. As a matter of fact, no Chlamydomonas small RNAs within the entire data set are homologs of known miRNAs, or of reported Arabidopsis small RNAs [data not shown]. The lack of universally conserved miRNA genes among plants, animals, and green algae suggests that miRNA genes may have evolved independently in the lineages leading to animals, plants, and green algae.

Next we asked whether Chlamydomonas miRNAs are conserved in other green algae. We searched for miRNA homologs in the three other green algae—Ostreococcus tauri, Ostreococcus lucimarinus, and Volvox carteri—whose draft genome sequences are available. V. carteri is a colonial multicellular green alga in the class Chlorophyceae, while O. tauri and O. lucimarinus belong to the class Prasinophyceae. We found that most of the Chlamydomonas miRNAs/Candidates did not have identifiable orthologs in the other three green algae. When up to three nucleotide substitutions were allowed, 11, 22, and 25 miRNAs/Candidates had homologous sequences in V. carteri, O. tauri, and O. lucimarinus, respectively. However, many of these potential orthologs are likely false positives, since they are located in the nonhomologous arms of the predicted stem–loops (Supplementary Database 2).

Prediction of miRNA targets in Chlamydomonas

As an initial step toward understanding the biological function of miRNAs in Chlamydomonas, we searched for miRNA targets among annotated protein-coding transcripts and ESTs of Chlamydomonas, adapting criteria for plant miRNA target prediction (Jones-Rhoades and Bartel 2004; Allen et al. 2005).

Applying a cutoff mispairing score of \( \leq 3.0 \), we predicted a total of 697 genes as miRNA targets [Table 2; Supplementary Table 2; Supplementary Database 3], with a signal/noise ratio of 5.6:1 as estimated by summing the number of targets predicted for miRNAs/Candidates and comparing with the average predicted targets for 10 shuffled cohorts. Most of the target sites are located in the coding sequences, with 40 being in the untranslated regions [UTRs]. The putative target genes appear to be involved in various biological processes. However, since the Chlamydomonas genome is not fully annotated and a great proportion of Chlamydomonas protein-coding genes have no known functions, it is difficult to draw a conclusion as to whether these miRNA targets have any functional bias. Among the predicted targets with annotated functions, many genes are involved in cellular metabolism and physiological processes, whereas transcription factors appear to be underrepresented. This is in contrast to the situation in plants where miRNAs have a remarkable propensity to target transcription factor gene families (Jones-Rhoades et al. 2006). It is noteworthy that many genes encoding Flagellar-Associated Proteins [FAPs] were predicted as miRNAs/Candidate targets. We also predicted histone 2A family proteins as the targets of Candidate136. In Arabidopsis, two histone 2A.F/Z genes are the predicted targets of miR841 (Rajagopalan et al. 2006).

Chlamydomonas miRNAs can direct target mRNA cleavage

In plants and animals, miRNAs directly interact with AGOs to form nucleoprotein complexes that direct target mRNA cleavage or translational repression. Chlamydomonas encodes two AGO proteins and one protein containing a Piwi domain but not a PAZ domain [Supplementary Fig. 1]. The expression of the two AGO proteins is supported by EST sequences and RT–PCR [data not shown], whereas that of the Piwi domain-only protein has yet to be verified. We reasoned that Chlamydomonas miRNAs should reside in AGO protein complexes in order to carry out their functions.

As a first step, we examined whether such a complex existed using size chromatography. We prepared total extracts from Chlamydomonas cultures, and then fractionated the extracts by gel filtration. RNAs were prepared from each fraction and used for small RNA Northern blotting. As shown in Figure 3A, several miRNAs including Candidate82 could be detected in the fractions corresponding to \( \sim 30–440 \) kDa with a peak around 150 kDa as determined by standard size markers. The average size of a 21-nt RNA is \( \sim 7 \) kDa; thus, the miRNAs must exist in complexes with other components, presumably including an AGO protein.

Next we sought to test through which mode [mRNA cleavage or translational repression] the Chlamydomonas complex acts on mRNA targets. We incubated an in vitro transcribed \(^{32}\)P-labeled RNA [jgiChlre3]196819|OVA_estExt_GenewiseH_1.C_540109; a putative di- or tricarboxylate translocator, the predicted target for Candidate82] with each fraction collected from size exclusion chromatography. We found that the transcripts were specifically cleaved into prod-
ucts of the expected size [Fig. 3B]. The peak of the cleavage activity corresponded to the peak of the miRNA signal as detected by Northern blot, suggesting that Candidate82 can direct the cleavage of its target. To examine whether Candidate82 can also cleave its target in vivo, we used a 5′/3′ RACE procedure to detect the cleavage products and to map precise cleavage sites (Llave et al. 2002). We could detect a 5′ RACE PCR product of predicted size from C_540109 on an agarose gel, suggesting that the target was cleaved in vivo [Fig. 3C]. Sequencing of the PCR product revealed that the majority of the cleavage sites were located between the target nucleotides that pair with nucleotides 10 and 11 of the miRNA (Fig. 3D), which is the characteristic feature of RISC-mediated cleavage (Elbashir et al. 2001; Llave et al. 2002; Rivas et al. 2005).

We tried to confirm the predicted targets of other miRNAs/Candidates using 5′/3′ RACE. For the FAP274 gene (Chlre3_159889), the predicted target of Candidate9, a few of the cleavage sites were mapped to the target regions, albeit more sites were found in the downstream region of the target regions [Fig. 3E]. For many other predicted targets, cleavage products were not detectable or cleavage sites were mapped outside the predicted target regions [very often in the downstream region of the predicted target regions]. This might attribute to quicker turnover of cleavage products in Chlamydomonas, compared with that in Arabidopsis, or to false-positive prediction of some of the miRNA targets.

Considered together, our results indicate that Chlamydomonas miRNAs can direct the cleavage of their targets.

Expression patterns of miRNAs/Candidates during Chlamydomonas gametogenesis

In multicellular organisms, miRNAs have been shown to play regulatory roles in cellular differentiation, developmental patterning, and timing (Bartel 2004). The discovery of a large number of miRNAs in unicellular Chlamydomonas raises pressing question about their biological functions. The life cycle of Chlamydomonas is rather simple compared with that of plants and animals. Cells of this species are haploid and divide asexually under normal growth conditions. Under stress conditions (e.g., lack of nitrogen), haploid cells of opposite
mating types become gametes and conjugate to form diploid zygotes. The zygote loses its flagella and produces a thick cell wall, forming a resting spore. When conditions become normal, the diploid zygote undergoes meiosis and produces four haploid cells that resume asexual reproduction (Harris 1989). As a first step to address the question whether miRNAs are involved in the progression of the Chlamydomonas life cycle, we sought to examine their expression profiles in vegetative cells and cells undergoing gametogenesis.

We probed the expression of eight randomly selected miRNAs/Candidates in vegetative cells and gametes using small RNA Northern blotting. Different expression patterns were observed for these miRNAs (Fig. 4). The expression of miR919-3p-2, Candidate12, and Candidate112 remained similar in both cell types. The expression of miR911 and miR912 was elevated in gametes, whereas that of miR906-3p, Candidate57, and Candidate82 was decreased at different degrees. The changed expression level of these miRNAs in gametes suggests that they might play roles in gametic differentiation.

Endogenous siRNAs in Chlamydomonas

Besides miRNAs/Candidates, we found that many small RNAs were derived from various genomic locations including protein-coding genes and intergenic regions. Interestingly, we identified 11 loci producing small RNAs (Fig. 5; Supplementary Table 4), the majority of which are phased relative to each other, reminiscent of plant ta-siRNAs. It remains to be examined whether these phased siRNAs are produced by a similar mechanism to that used by plant ta-siRNAs, the generation of which requires the actions of miRNAs and RDRs. However, no RDR homologs can be predicted from the current version of the Chlamydomonas genome sequence.

Besides the ta-siRNA-like small RNAs, we also found a large proportion of siRNAs derived from sense or antisense strands of other protein-coding genes or intergenic regions (Table 1). The biogenesis mechanism of these siRNAs remains to be elucidated. It also remains to be examined whether these siRNAs can regulate gene expression in cis and/or in trans.

It has been shown in several organisms that small RNAs can play an important role in silencing of repetitive sequences and formation of heterochromatin (Lippman and Martienssen 2004). We asked whether small RNAs could also act to silence repeats in Chlamydomonas. As a first step, we examined whether there were any siRNAs originating from repeats. By the RepeatMasker program (Kohany et al. 2006), 6.11% of the Chlamydomonas genome could be annotated as transposable elements, low complexity, or simple repeats [Supplementary Table 5]. Intriguingly, we only identified 26 unique siRNAs derived from transposable elements, 114 from simple repeat regions, and 46 from low complexity regions. These three groups of unique small RNAs represent 41, 206, and 65 sequences from the redundant data set, respectively [Supplementary Table 5]. Therefore, it seems that small RNAs in Chlamydomonas may not be important for silencing of repetitive elements.

Discussion

The diversification of small RNA pathways has been well demonstrated in multicellular systems but remains largely unknown in unicellular organisms. Here we have shown that the unicellular green alga Chlamydomonas contains a variety of small RNAs. These include miRNAs and phased siRNAs, as well as other siRNAs originating from intergenic regions, sense and antisense strands of protein-coding genes, and repeat regions. The presence of these small RNAs indicates an unexpected complexity of RNAi-related processes in this unicellular organism. Our discovery of miRNAs in a unicellular alga also supports the concept that the miRNA pathway is a conserved mechanism of gene regulation in eukaryotes that arose prior to the developmental patterning needed for multicellular body plans [Bartel 2004].

We did not find that the Chlamydomonas miRNAs we discovered have any identifiable homologs in higher plants. However, the lack of sequence homology between plant and Chlamydomonas miRNAs does not necessarily reflect the fundamental differences in miRNA silencing pathways. In fact, Chlamydomonas miRNAs share several common features with plant miRNAs. These include being ~21 nt in length and having a preference for uracil at the 5' end, as well as being generated from a stem–loop by Dicer and being competent in directing target RNA cleavage. This suggests a conserved machinery for miRNA production and action.
in Chlamydomonas and in higher plants. It has been proposed that plant miRNAs may have arisen from inverted duplication of their target genes [Allen et al. 2004]. Supporting this proposal, several presumably newly evolved Arabidopsis miRNA loci have been shown to share significant sequence similarity to their respective target genes [Allen et al. 2004; Rajagopalan et al. 2006]. Moreover, a few small RNA-producing loci that can form relatively long hairpins were identified as transitional forms leading to miRNAs [Allen et al. 2004]. Intriguingly, a substantial number of the Chlamydomonas miRNAs/Candidates are located in long hairpins, many of which have nearly perfect base-pairing between the arms and have the potential to produce multiple small RNAs. These long hairpins might represent the evolutionary intermediates of miRNA genes.

It has been well demonstrated in plants and animals that miRNAs play key roles in multicellular development [Bartel 2004]. In this study, we identified many miRNAs in the unicellular Chlamydomonas. The biological functions of these miRNAs remain to be understood. It is of great interest to examine whether they play regulatory roles in sexual and/or asexual reproduction, responses to a variety of environmental changes, or metabolic pathways. We found that the expression of some miRNAs/Candidates was changed during gametogenesis [Fig. 4], suggesting the involvement of miRNAs in Chlamydomonas sexual reproduction. We predicted many genes encoding FAPs as miRNAs/Candidates targets, and validated FAP274 as the target of Candidate9, suggesting the involvement of miRNAs in flagellar development. We also have predicted many other targets for Chlamydomonas miRNAs. However, since the Chlamydomonas genome awaits better annotation and functions of many of the annotated genes have yet to be assigned, validation of miRNA targets and elucidation of their functions in Chlamydomonas biology is a major but exciting challenge. Analyzing transgenic lines over-expressing miRNAs and cleavage-resistant targets may reveal which biological process/es each miRNA is controlling. Additionally, using microarrays to profile the expression of miRNAs and their targets at different steps of the life cycle or under different physiological conditions will help us understand the biological roles of Chlamydomonas miRNAs.

Most animal miRNAs regulate expression of their targets by repressing productive translation with the one exception that mouse miR196 guides the cleavage of its target HoxB8 mRNA [Bartel 2004; Yekta et al. 2004]. In contrast, mRNA cleavage is the predominant mechanism used by plant miRNAs to regulate their targets. It remains unclear whether some plant miRNAs can also function as translation repressors [Jones-Rhoades et al. 2006]. Through the detection of miRNA cleavage products in vivo by the 5’ RACE method and in vitro by cleavage assay, we showed that Chlamydomonas miRNAs can direct the cleavage of their target mRNAs. However, we do not exclude the possibility that Chlamydomonas miRNAs can also function at the level of translation, in addition to mRNA cleavage. We also point out that the criteria we used for target prediction were adapted from plants; thus, it is possible that we have missed some target genes that are regulated by Chlamydomonas miRNAs through modes similar to those used in animals.

It has been well established that RNAi plays an important role in the silencing of selfish genetic elements and in the formation of heterochromatin in plants, ani-
It has been proposed that siRNAs guide heterochromatic silencing complexes to homologous target loci through RNA/RNA recognition [Motamedi et al. 2004; Sugiyama et al. 2005; Qi et al. 2006]. Screening for *Chlamydomonas* mutants that are defective in silencing of a transgene at either the transcriptional or post-transcriptional level led to the identification of three genes, *Mut11* encoding a WD40-repeat protein, *Mut9* encoding a Set/Thir kinase, and *Mut6* encoding a DEAH-box RNA helicase [Wu-Scharf et al. 2000; Jeong Br et al. 2002; Zhang et al. 2002]. In these mutants, several transposons were reactivated. However, evidence for direct involvement of these proteins in RNAi is still lacking, and the role of RNAi in transposon silencing in *Chlamydomonas* remains to be evaluated. Among the 4182 unique small RNAs we obtained from *Chlamydomonas*, <4% of small RNAs were derived from repeats [Supplementary Table 5]. This fraction is much less than what would be expected based on a random distribution, considering that the repeats account for 6.11% of the *Chlamydomonas* genome as defined by RepeatMasker [Kohany et al. 2006]. Thus, RNAi might not be a general mechanism used to silence transposons and repetitive elements. Alternatively, RNAi-controlled silencing of repeats may not be dependent on small RNA accumulation in *Chlamydomonas* under vegetative growth conditions. In Tetrahymena, scnRNAs are only induced during conjugation to direct developmentally programmed DNA elimination [Mochizuki and Gorovsky 2004a; Lee and Collins 2006]. It remains possible that RNAi may play a role in silencing repeats when *Chlamydomonas* undergoes sexual reproduction. Profiling of small RNAs of *Chlamydomonas* during this process will be necessary to address this possibility.

In plant and animal species studies so far, there are multiple Dicer and Argonaute family members that are genetically and functionally diversified [Carmell et al. 2002; Carmell and Hannon 2004]. For instance, in *Arabidopsis*, there are four Dicer-like and 10 Argonaute proteins. It has been demonstrated that DCL1 has a unique role in miRNA biogenesis, DCL3 produces ~24-nt hc-siRNAs, and DCL2 and DCL4 are involved in antiviral defense and the production of nat-siRNAs or ta-siRNAs [Vacheret 2006]. AGO1 binds miRNAs to catalyze the cleavage of miRNA target mRNAs [Baumberger and Baulcombe 2005; Qi et al. 2005], while AGO4 recruits hc-siRNAs to direct DNA/histone methylation, sometimes resulting in transcriptional gene silencing [Zilberman et al. 2003; Qi et al. 2006]. The *Chlamydomonas* genome encodes three Dicer-like and at least two Argonaute proteins [Schroda 2006]. It will be of great interest to examine whether the different types of small RNAs that we have uncovered are processed by the same or different Dicers, and whether they enter distinct complexes containing different Argonaute proteins. Genetic analysis of RNAi lines or mutants for each Dicer-like and Argonaute gene, in combination with biochemical purification of their complexes, will shed light on the mechanisms of RNAi pathways in *Chlamydomonas*.

Materials and methods

*Chlamydomonas* strains and culture conditions

*Chlamydomonas* strain CC-503cw92 mt+ was used as the source of small RNA cloning. CC-621 [mt−] was used to produce gametes. *Chlamydomonas* cells were grown in Tris-acetate-phosphate (TAP) liquid medium [Harris 1989] on a rotary shaker at 23°C, under constant light. Gametes were generated essentially as described [Wegener et al. 1989]. Gametic differentiation was induced by transferring vegetative cells in mid-log phase to N-free TAP medium for 18–20 h under the same culture conditions.

Small RNA preparation and cloning

Total RNA was extracted from *Chlamydomonas* cultures at vegetative growth in TAP medium using Trizol (Invitrogen), and small RNAs were enriched by PEG precipitation method. The isolated small RNAs were separated by 15% denaturing PAGE and visualized by SYBR-gold staining, and small RNAs of 18–28 nt were gel-purified. Small RNAs were ligated to a 3′ adaptor and a 5′ acceptor sequentially, and then RT–PCR-amplified as described [Lau et al. 2001]. PCR products were reamplified using a pair of 454 cloning primers and provided to 454 Life Sciences for sequencing as previously described [Girard et al. 2006].

The genome and transcript sequences

The *Chlamydomonas* genome (version 3.0) was downloaded from the JGI *Chlamydomonas* Web site [http://genome.jgi-psf.org/Chlam3/Chlam3.home.html]. Since the current *Chlamydomonas* genome sequences were provided as 1266 scaffolds, to facilitate data analysis, we linked all these scaffolds together following the order of their numbers to build an arbitrary genome. All analyses described in the text were carried out using information extracted from this arbitrary genome. Repeat regions and transposable elements of the *Chlamydomonas* genome were identified by RepeatMasker (version 3.1.3) according to RepBase [version 10.09] [Jurka et al. 2005].

The sequences of 15,143 annotated *Chlamydomonas* protein-coding genes [including pseudogenes] were also obtained from the *Chlamydomonas* Web site [http://genome.jgi-psf.org/Chlre3/Chlre3.home.html]. The genomic location and intron-exon structure of each gene were determined by mapping gene sequences to the arbitrary *Chlamydomonas* genome using the BLAT program [Kent 2002].

The complete genome sequences of *O. tauri* and *O. lucimarus* were downloaded from the *O. tauri* Web site [http://genome.jgi-psf.org/Ostab4/Ostab4.home.html] and the *O. lucimarus* Web site [http://genome.jgi-psf.org/Ost9901_3/Ost9901_3.home.html].

Small RNA analysis

After removing adaptor/acceptor sequences from the raw reads obtained from 454 Life Sciences, the remaining small RNA sequences were mapped to the arbitrary *Chlamydomonas* genome using Perl scripts. Relationships of small RNAs to annotated genes were determined by comparing the genomic loci of small RNAs with the BLAT results of gene sequences. Small RNAs derived from known noncoding RNAs were identified by comparing small RNAs with the sequences of noncoding RNAs collected in Rfam [http://www.sanger.ac.uk/Software/Rfam] [Griffiths-Jones et al. 2005].
Predictions of miRNAs

The prediction of *Chlamydomonas* miRNAs was carried out using criteria similar to those applied for *Arabidopsis* miRNA prediction previously (Wang et al. 2004). Basically, the upstream and downstream genomic sequences of each miRNA locus within intergenic and intronic regions were extracted and subjected to RNA secondary structure check using MFold (Zuker 2003). Genomic sequences with lengths ranging from 70 to 350 nt with 20-nt increments were extracted from each miRNA locus. Since mature miRNAs could be located on either the 5‘ arm or the 3‘ arm of their hairpin-structured precursors, two genomic sequences with the same length were extracted from each miRNA locus, with the cloned small RNA sequences mapped near the 5‘ end or the 3‘ end, respectively. If the lowest energy form of any genomic sequence extracted as described above contained a relatively long double-stranded region that included the cloned small RNA sequence, the small RNA was selected as a miRNA candidate and the genomic sequence was considered as a miRNA precursor. All the sequences and structures of the precursors were manually checked. In some cases, the precursor sequences were manually extended up to ~1 kb to get structures of the lowest free energy. In cases in which two small RNAs were mapped to the same stem–loop and they formed a duplex with a 2-nt overhang, the one with higher cloning frequency was chosen as a miRNA, while the other was considered as miRNA*.* When the two small RNAs had the same cloning frequency, then both were selected as miRNAs. In cases in which two small RNAs were mapped to the same stem–loop but they could not form a duplex with a 2-nt overhang, both small RNAs were selected as miRNA candidates.

Prediction of miRNA targets

We modified criteria that were developed for plant miRNA target prediction to predict targets of *Chlamydomonas* miRNAs (Jones-Rhoades and Bartel 2004; Allen et al. 2005). The putative target sites of all miRNA candidates were identified by aligning miRNA sequences to the annotated gene sequences and ESTs of *Chlamydomonas* using Perl script. The method enables identification of multiple target sites of a single miRNA on the same target sequence. An initial pool of predicted targets was created, with at most four unpaired nucleotides (including G:U pairs, mismatches, and single-nucleotide bulges) and two single-nucleotide bulges allowed between a miRNA and its targets. A mispair scoring system was then applied to these initial targets. Mismatches and single-nucleotide bulges were each scored as 1, and G:U pairs were each scored as 0.5. The scores were doubled if mismatches, G:U pairs, and bulges were located at positions at 2–13 as counted from the 5‘ end of a miRNA. Genes with a mispair score ≥3 were selected as putative miRNA targets. In order to assess the specificity of the prediction method, we generated 10 shuffled cohorts for all the miRNAs/Candidates. The signal/noise ratio was calculated by comparing the total number of targets [the “signal”] predicted for miRNAs/Candidates and that for shuffled cohorts.

Detection of phased siRNAs

Small RNAs with a phased position relationship were identified by checking the distance among all small RNAs within every 1000-nt window, anchored at one end of each small RNA. Small RNAs with interval distances that were a multiple of 21 nt (allowing ±2 nt divergence) were selected as phased small RNAs. Every small RNA within the 1000-nt window was examined to identify most small RNAs with the same phased relationship.
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