A horizontally acquired group II intron in the chloroplast psbA gene of a psychrophilic Chlamydomonas: In vitro self-splicing and genetic evidence for maturase activity

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ABSTRACT
The majority of known group II introns are from chloroplast genomes, yet the first self-splicing group II intron from a chloroplast gene was reported only recently, from the psbA gene of the euglenoid, Euglena myxocylindracea. Herein, we describe a large (2.6-kb) group II intron from the psbA gene (psbA1) of a psychrophilic Chlamydomonas sp. from Antarctica that self-splices accurately in vitro. Remarkably, this intron, which also encodes an ORF with putative reverse transcriptase, maturase, and endonuclease domains, is in the same location, and is related to the E. myxocylindracea intron, as well as to group IIB2 introns from cyanobacteria. In vitro self-splicing of Chs.psba1 occurred via a lariat, and required Mg2+ (>12 mM) and NH4+. Self-splicing was improved by deleting most of the ORF and by using pre-RNAs directly from transcription reactions, suggestive of a role for folding during transcription. Self-splicing of Chs.psba1 pre-RNAs showed temperature optima of ≈44°C, but with a broad shoulder on the low side of the peak; splicing was nearly absent at 50°C, indicative of thermolability. Splicing of wild-type Chs.psba1 also occurred in Escherichia coli, but not when the ORF was disrupted by mutations, providing genetic evidence that it has maturase activity. This work provides the first description of a ribozyme from a psychrophilic organism. It also appears to provide a second instance of interkingdom horizontal transfer of this group IIB2 intron (or a close relative) from cyanobacteria to chloroplasts.

Keywords: Chlamydomonas; chloroplast; group IIB2 intron; horizontal transfer; maturase; psbA gene; psychrophilic; self-splicing

INTRODUCTION
Ribozymes play key roles in gene expression likely in all cells; however, relatively little is known about how they have adapted to the wide range of niches that organisms occupy in the biosphere. Adaptation to temperature is one of the best studied aspects of biochemical adaptation (Hochachka and Somero 2002); however, only a few of these studies have been with ribozymes, and all involved thermophily (Brown et al. 1993; Fang et al. 2001; Nesbo and Doolittle 2003). Protein enzymes from psychrophilic organisms typically have lower temperature optiums, reduced thermostability, and are more efficient catalysts than mesophilic enzymes (Hochachka and Somero 2002). Greater flexibility is one mechanism that psychrophilic protein enzymes use to maintain function at low temperature. It is not known how ribozymes have adapted to function at low temperature, because classical ribozymes have not been characterized from psychrophilic organisms.

Group II introns are one of the major classes of large ribozymes, and are found in eukaryotic organelles, prokaryotes (for review, see Michel and Ferat 1995; Bonen and Vogel 2001), and archaea (Dai and Zimmerly 2003). They have conserved terminal sequences and secondary structure elements, and several subgroups have been defined. Some group II introns contain an ORF that has coevolved with the intron core (Fontaine et al. 1997; Toor et al. 2001), and archaea (Dai and Zimmerly 2003). They have conserved terminal sequences and secondary structure elements, and several subgroups have been defined. Some group II introns contain an ORF that has coevolved with the intron core (Fontaine et al. 1997; Toor et al. 2001), and for the few whose functions have been studied, they promote splicing (maturase) and mobility of the host intron (Lambowitz et al. 1999). Several group II introns from mitochondria and bacteria have been shown to self-splice in vitro in the absence of proteins (Michel and Ferat 1995; Bonen and Vogel 2001). Splicing occurs by a two-step
mechanism, with the first step proceeding mainly via a lariat, but it can also occur by hydrolysis (Daniels et al. 1996; Podar et al. 1998; Vogel and Börner 2002). Although the majority of known group II introns are from chloroplast genes (Cannone et al. 2002), the first report of self-splicing by a chloroplast group II intron appeared just recently (Sheveleva and Hallick 2004). Also, chloroplast group II intron-encoded maturases, in particular matK, have been inferred (du Jardin et al. 1994; Liere and Link 1995; Vogel et al. 1997, 1999; Doetsch et al. 1998), but not directly demonstrated to promote splicing.

Species of *Chlamydomonas* are known from cold environments, where they are often referred to as snow or ice algae (Hoham and Duval 2001; Thomas and Dieckmann 2002). The chloroplast genomes of *Chlamydomonas* spp. (mostly mesophilic) are excellent sources of group I intron ribozymes (Herrin et al. 1998), but for unknown reasons, typical cis-spliced group II introns have not been reported. To identify a ribozyme from a psychrophilic organism, we searched chloroplast *psbA* and 23S rRNA genes of snow/ice *Chlamydomonas* by PCR, expecting to find group I introns. Surprisingly, a cis-spliced group II intron (*Csp.psbA1*) containing a large ORF was present in the *psbA* gene of a *Chlamydomonas* from the depths of Lake Bonney, Antarctica, a permanently ice-covered lake (Neale and Priscu 1995). It has been possible to examine the splicing of this intron in vitro, and in vivo in *Escherichia coli* and in its normal host.

**RESULTS**

**The organism**

*Chlamydomonas* strain CCMP-1619 grew well at temperatures ranging from 4°C (the lowest temperature tried) to 15°C, but did not grow at 20°C, confirming that it is a psychrophilic organism. CCMP-1619 also did not grow when the cultures were shaken, consistent with adaptation to its highly nonturbulent environment (Neale and Priscu 1995). Several studies of a prominent *Chlamydomonas* from Lake Bonney referred to it as *C. subcaudata* (Morgan et al. 1998; Morgan-Kiss et al. 2002), which is an egg-shaped species with a protrusion of the posterior cell wall (Ettl 1976). Light microscopy of CCMP-1619 did not reveal any cells with that morphology; however, instead, the cells were highly spherical and a typical example is shown in Figure 1. The larger cells were ∼15 µm in length, and most appeared to be highly vacuolated.

To determine the relationship between CCMP-1619, the species studied by Huner et al. (Morgan et al. 1998; Morgan-Kiss et al. 2002), and *C. subcaudata*, the chloroplast *petA-petD* region was amplified from CCMP-1619 and sequenced (accession no. AY332410). The *petA* (723 bp) and *petD* (37 bp) coding regions from CCMP-1619 are identical to those deposited by Huner et al. (accession no. AY039799), and the intergenic spacer differs by <1% (eight out of 1065 residues). However, the *petA* coding region from authentic *C. subcaudata* is only ∼90% identical to CCMP-1619, and the intergenic spacer is only ∼40% identical (not shown). Thus, these results indicate that CCMP-1619 is essentially the same species studied previously (Morgan et al. 1998; Morgan-Kiss et al. 2002), but it is not *C. subcaudata*. Analysis of nuclear rRNA ITS sequence, and other data, suggests that it is a strain of *Chlamydomonas raudensis* (T. Pocock, M.A. Lachance, T. Proeschold, S. Kim, and N. Huner, pers. comm.).

**The intron**

Amplification and sequencing of the 3′ half of the *psbA* gene from CCMP-1619 (accession no. AY253205) revealed the presence of one large (2572 bp) group II intron that splits the V185 codon. Computer translation of the flanking exon sequences (284 bp) gave an amino acid sequence identical to *Chlamydomonas reinhardtii psbA*. BLAST searches using the exon DNA sequences produced the highest similarity scores with green algal *psbA* genes. The G-C content (38%) and codon usage of the CCMP-1619 exon sequences are also similar to those of green algal chloroplast genes, which are quite different from nuclear genes in these organisms. These results indicate that the amplified *psbA* gene is from the chloroplast of CCMP-1619, which is where it is located in all other photosynthetic eukaryotes that have been examined.

The predicted secondary structure of this *Chlamydomonas* sp. (CCMP-1619) *psbA* intron, which we named *Chs.psbA1* (Fig. 2), has the common group II domains I–VI and all of the tertiary interactions characteristic of group IIB2 (chloroplast-like class 2) introns (Michel et al. 1989; Toor et al. 2001). The intron is most similar in secondary structure to the recently described intron from the chloroplast *psbA* gene of *Euglena myxoxylindracea* (Sheveleva and Hallick 2004) and to introns from the cyanobacteria *Calothrix* sp. (*Cal.x1*; Ferat and Michel 1993) and *Nostoc* sp. PCC 7120 (bp 45,420–48,000; Kaneko et al. 2001), which are not located in *psbA* genes. An idiosyncrasy of the *Chs.psbA1* intron is an unusual insertion in domain ID (nt 275–310).
that is shared with the *E. myxocylindracea* intron, although
the sequence of the insertion has diverged considerably.

Chs.psbA1 is also inserted at the same location in the psbA gene as the euglenoid intron. Finally, both of these introns are atypical in that they end with CC instead of AY (Bonen and Vogel 2001).

The large ORF (600 amino acids) in domain IV of Chs.psbA1 is preceded by a putative Shine–Dalgarno sequence (AGUAGGU), and contains reverse transcriptase (RT, including subregions 0–7), maturase (X), DNA-binding (D), and endonuclease (HNH) domains (Fig. 3A). Phylogenetic analysis of the ORF (Fig. 3B) places it in subgroup 11B2 or the chloroplast-like class 2 clade of Toor et al. (2001). The highest sequence identity (52%) was with the ORF of the *E. myxocylindracea* intron, and the next highest was with the *Calothrix* and *Nostoc* ORFs (46%); the lowest sequence identity was with the *Lactococcus lactis* intron ORF (25%). The fact that both the ORF and core structure of Chs.psbA1 place it in the same subgroup suggests that the ORF and intron core have coevolved.

**Self-splicing of Chs.psbA1 in vitro**

In the initial test for self-splicing of Chs.psbA1, a pre-RNA containing the full-length intron (similar to wild-type pre-RNA in Fig. 4A but with a longer 3′ exon) was used. The pre-RNA was incubated under conditions previously shown to promote self-splicing of other group II introns and analyzed by denaturing gel electrophoresis. An autoradiograph, similar to that shown in Figure 4B (lanes 1,2), indicated the presence of slowly migrating lariat as well as spliced-exon RNAs. When these reactions were subjected to reverse transcription and PCR (RT-PCR) with flanking exon primers, a product of the size expected for spliced exons was obtained (data not shown, but see below).

Because only a very small fraction of the 2.9-kb pre-RNA spliced, steps were taken to improve the splicing efficiency. One was to shorten the 3′ exon to 92 nt by using primer 177 instead of 100 for PCR (as in the wild-type pre-RNA in Fig. 4B); however, that resulted in only a small increase in splicing efficiency. Hence, we deleted 1575 bp of the ORF (amino acids 43–567), creating the ΔORF.1 mutant (boxed sequence above the wild-type structure).

![FIGURE 2. Proposed secondary structure of the 2572-bp Chs.psbA1 intron. Intron nucleotides are in uppercase and numbered from the 5′ end; exon nucleotides are in lowercase. 5′SS and 3′SS are the splice sites. The major domains are labeled with Roman numerals (I–VI) and subdomains of domain I as A, B, C (C1 and C2), and D (D′, D″, D2a, D2b, D3). Tertiary interactions are labeled with Greek letters (α–α′, etc.). EBS and IBS refer to exon and intron binding sites, respectively. The Shine–Dalgarno (SD) sequence and start codon (AUG) of the ORF in domain IV are boxed, and the branch-site A in domain VI is circled. An alternative structure of the unusual insertion in domain D (nt 275–310) is proposed for the ΔORF.1 mutant (boxed sequence above the wild-type structure).](https://www.rnajournal.org/1099)

[Image of secondary structure of Chs.psbA1 intron]

The ORF template (Fig. 4A). We also cloned the ΔORF PCR product, generating the ΔORF.1 plasmid DNA (Fig. 4A). Sequencing of the ΔORF.1 insert, however, revealed four substitutions compared to the original full-length (wild-type intron) PCR product: one change (C304 → U304) was in the domain-I insertion (see the alternative, boxed region in Fig. 2), and the other three (U756 → C756, C844 → A844, and C874 → U874) were in the remnant of the ORF in ΔORF.1. These substitutions were probably introduced during sequential PCR. As Figure 4B shows, the ΔORF and ΔORF.1 pre-RNAs spliced more efficiently than wild type. It was also easier to visualize the smaller lariat RNAs with these constructs. The multiple bands of the
The optimum concentrations of Mg\(^{2+}\) and NH\(_4^+\) ions, which are typically used in group II self-splicing experiments, were examined with Chs.psbA1. Splicing was detected at 12–13 mM MgCl\(_2\) (500 mM NH\(_4\)Cl), and there was a broad optimum of 25–100 mM. Self-splicing of Chs.psbA1 was undetectable in the absence of NH\(_4^+\), and was optimal at 1–1.5 M NH\(_4\)Cl. Spermidine did not stimulate Chs.psbA1 self-splicing.

The effect of a denaturation–renaturation cycle was also examined. Renatured RNAs spliced more efficiently than RNAs purified from denaturing gels without renaturation, but less efficiently than RNAs taken directly from transcription reactions (not shown). Even extracting the transcription reactions with chloroform reduced the splicing efficiency of the various RNAs, as shown by the ΔORF and ΔORF.1 pre-RNAs in Figure 4B. Splicing of extracted wild-

FIGURE 3. Domain structure and phylogenetic analysis of the Chlamydomonas sp. (CCMP-1619) Chs.psbA1 intron ORF. (A) Diagram of the intron and ORF subdomains. (RT) reverse transcriptase; (X) maturase; (D) DNA-binding; (HNH) endonuclease. (B) Phylogenetic analysis of the ORF. The neighbor-joining tree was generated with clustal X using amino acid sequences of selected group II intron ORFs. The scale bar is 0.1 substitutions per site. The bootstrap values are out of 1000 possible trees, and the L. lactis protein was the out-group. The ORFs contained all of the domains indicated in A. The ORF sequences and their accession numbers are: Calothrix sp. (X71464.1:1034–2788); Chlamydomonas, Chlamydomonas sp. CCMP-1619 chloroplast psbA intron; E. coli, Escherichia coli 0157:H7 (AE074613.1:158,849–60,573); E. myxoyclindracea, Euglena myxoylindracea chloroplast psbA intron (AY290861); L. lactis, LtrA of Lactococcus lactis (AAP06503.1); Nostoc 1, Nostoc sp. PCC 7120 (AP003599.1:45,989–47,791); Nostoc 2, Nostoc sp. PCC 7120 (AP003599.1:30,840–32,459); P. purpurea, Porphyra purpurea mitochondrial rRNA intron 1 (AF114794.1:2909–4543); P. littoralis, Pyliella littoralis mitochondrial rRNA intron 2 (AJ277126.1:4664–6370); S. cerevisiae a1, Saccharomyces cerevisiae mitochondrial COXI intron a1 (NP_009310.1); S. cerevisiae a2, Saccharomyces cerevisiae mitochondrial COXI intron a2 (NP_009309.1); T. elongatus 1, Thermosynechococcus elongatus BP-1 (AP005369.1:28791–30,485); T. elongatus 2, Thermosynechococcus elongatus BP-1 (AP005371.1:25,718–27,406).

FIGURE 4. In vitro self-splicing of 32P-labeled Chs.psbA1 pre-RNAs. (A) Maps of the pre-RNAs. Wild-type and ΔORF pre-RNAs have the same 5′ (67 nt) and 3′ (92 nt) exons, whereas the 5′ and 3′ exons of ΔORF.1 have an additional 5 nt. The wild-type intron is 2572 nt, and 997 nt in the ΔORF and ΔORF.1 pre-RNAs. The nucleotide substitution in the ΔORF.1 core is indicated. See Figure 3 for explanation of ORF domains. (B) Gel analysis of splicing reactions with 32P-labeled pre-RNAs. The reactions were incubated at 0°C or 42°C for 90 min and then analyzed on a denaturing polyacrylamide gel. "Unextracted" RNAs were taken from transcription reactions and used for splicing without treatment; "extracted" RNAs were extracted with chloroform and ethanol precipitated before splicing. The location of excised lariat, precursor, and spliced-exon RNAs are indicated to the left.
type pre-RNA was not included in Figure 4B, because it was so inefficient. These results suggest that some folding of the pre-RNA occurs in the transcription reaction that improves subsequent self-splicing. The folding is presumably incomplete, however, because there was no evidence of splicing during transcription. By incorporating all of the above findings, plus the temperature optimization (see below), ~20% of the ΔORF and ~30% of the ΔORF.1 pre-RNAs self-spliced in vitro after 180 min.

Reverse transcription coupled with PCR (RT-PCR) was used to judge the accuracy of self-splicing and to determine the branch site. As shown in Figure 5A, the RT-PCR product corresponding to spliced-exons RNA (E1–E2) depended on incubation of the splicing reaction at elevated temperature, as expected, and was also reverse transcriptase dependent (not shown). Sequencing of this product from wild-type as well as ΔORF and ΔORF.1 splicing reactions confirmed that the exons had been spliced correctly.

A nested RT-PCR assay was used to map the branch site in the wild-type intron lariat as described by Vogel et al. (1997). This assay depends on the ability of reverse transcriptase to read through a 2’–5’ linkage in the intron lariat at the branch site (diagram in Fig. 5B). In the PCR step, a nested downstream oligonucleotide (primer 316 in Fig. 5B) and an upstream primer (317) are used. A single product was obtained with this assay on wild-type pre-RNA that had been spliced at 42°C, but no product was obtained with pre-RNA incubated for splicing at 0°C (Fig. 5B, lower panel). The sequence of this product (on the sense strand) at the junction of the 3’ and 5’ regions of the intron was 3’-GCGTGTGCTGG-5’, where the underlined nucleotide is the branch site. This corresponds to the bulge A2565 in domain VI, similar to other group II introns. It should be noted that with this method, the reverse transcriptase inserts an A instead of a T when it encounters the branched A (Vogel et al. 1997).

The temperature optimum for self-splicing was tested with all three unextracted pre-RNAs, as well as extracted forms of ΔORF and ΔORF.1. Figure 6 shows the fraction of each RNA that spliced in 90 min, based on lariat accumulation. The lariat was used because it was relatively easy to quantify, and its accumulation correlated well with accumulation of spliced-exons RNA. The optimum splicing temperature for the different pre-RNAs was close to 44°C. Figure 6 also shows that there is a broad shoulder on the low side of the peak, and, moreover, that splicing of the more active pre-RNAs was observed below 20°C. At temperatures above the optimum, self-splicing dropped off quickly; in fact, with the most active pre-RNA (ΔORF.1,
unextracted), there was almost no splicing at 50°C. The steep drop in self-splicing at the higher temperature most likely reflects a loss of structure.

To determine rate constants for Chs.psbA1 self-splicing, time-course reactions of the shorter pre-RNAs (ΔORF and ΔORF.1) were performed at 41°C; a representative analysis for the ΔORF (extracted) RNA is shown in Figure 7A. The reaction was fairly linear at the early time points (10–20 min) and was nearly complete by 100 min (Fig. 7B). The linearity of the semi-log plot of the decay of active pre-RNA over time (Fig. 7C) suggested that the active fraction was one kinetic component. The \( k_{obs} \), for the initial splicing rate of this pre-RNA was 0.017 min\(^{-1}\). The \( k_{obs} \) for self-splicing by the ΔORF and ΔORF.1 pre-RNAs, both unextracted and extracted, was also determined, and found to vary less than twofold, ranging from 0.016 min\(^{-1}\) to 0.020 min\(^{-1}\); these rates are similar to other self-splicing group II introns (Hebbar et al. 1992; Jarrell et al. 1988; Adamidi et al. 2003).

**Splicing of Chs.psbA1 in E. coli**

Because chloroplast genes, including intron-encoded ones (Lee 2003), usually express at some level in *E. coli*, we decided to see if the Chs.psbA1 intron would splice in *E. coli* and, if so, if its splicing would be ORF dependent. The recombinant intron plasmids also contained a T7 promoter upstream of the inserted DNA, which enabled transcription in *E. coli* strains producing the T7 RNA polymerase. RT-PCR with primers flanking the intron was used to assess splicing of Chs.psbA1. As Figure 8 shows, a product of the size expected for spliced-exons RNA (E1–E2) was obtained with the wild-type, but not the ΔORF.1 intron; the figure also shows that the E1–E2 product is reverse transcriptase dependent. An RT-PCR product of the size expected for unspliced RNA (pre-RNA) was obtained for both wild-type and ORF.1 introns, verifying that the pre-RNA was transcribed in both clones. Analysis of three other wild-type clones and one other ΔORF.1 clone produced the same results (data not shown). Sequencing of the E1–E2 product verified that it was from correctly spliced RNA. We also sequenced the RT-PCR products migrating at ~0.6 kb and just faster than the E1–E2 product in Figure 8, which revealed that these were due to spurious priming.

Two other Chs.psbA1 intron mutants were examined for splicing in *E. coli* using the same assay; one has a frame-shift mutation after amino acid 10 of the ORF (FS), and the other has two point (i.e., missense) mutations (PM), one in the mature domain (G456→D456), and the other in the reverse transcriptase domain (L333→P333). Figure 8 shows that there was little or no splicing of Chs.psbA1 with these two mutants. Subjecting the products of the RT-PCR reactions in Figure 8 to another round of PCR with the same or a nested downstream primer still did not produce any spliced-exons product with the ΔORF.1 and missense (PM) mutants, but a small amount was visible with the frame-shift mutant (not shown). The latter result may be due to some translation from an in-frame GUG codon, which is 96 nt from the normal start codon and preceded by a possible Shine–Dalgarno sequence (AAGGA), or from the first internal methionine (M123) of the ORF, which is also preceded by a possible Shine–Dalgarno sequence (GAAG). In summary, these results provide genetic evidence that the Chs.psbA1 ORF encodes a maturase.

**Splicing of Chs.psbA1 in Chlamydomonas CCMP-1619**

The splicing efficiency of Chs.psbA1 in CCMP-1619 cells growing photoautotrophically at 8°C (Morgan et al. 1998) was assessed by Northern blot hybridization (Fig. 9). The exon-specific probe gave a very strong signal with a 1.1-kb RNA that comigrated with *C. reinhardtii* psbA mRNA (lane 1). The intron-specific probe detected RNAs of 2.6, 3.2,
DISCUSSION

We have identified a group II intron in the chloroplast psbA gene of a psychrophilic Chlamydomonas sp. that shows self-splicing activity in vitro. Chs.psbA1 is only the second group II intron from a chloroplast gene shown to be autocatalytic—the first was reported while this article was in preparation (Sheveleva and Hallick 2004), and, remarkably, is closely related to this Chlamydomonas intron, although it is from an unrelated euglenoid. The relative paucity of autocatalytic group II introns from chloroplasts may be because most of the known ones are from vascular plants and euglenoids, which appear to rely mainly on protein-dependent splicing (Jenkins et al. 1997). The Chs.psbA1 intron is also the first cis-spliced group II intron from Chlamydomonas, and it is not closely related to the two trans-spliced group II introns in the psaA1 gene (for review, see Herrin et al. 1998).

The proposed secondary structure for Chs.psbA1 places it in subgroup IIB2; however, it has an idiosyncratic insertion in domain ID that can affect splicing. The latter conclusion is based on the fact that the ΔORF1 pre-RNA, which has a nucleotide substitution in the insertion, splices more efficiently than the ΔORF pre-RNA, which does not. Comparison of the predicted wild-type and mutant structures in this region suggests that there is a little better stability in the mutant, which, in turn suggests that this unusual insertion may inhibit self-splicing in vitro by forming alternate structures with other parts of the intron.

We obtained genetic evidence that the ORF in Chs.psbA1 has maturation activity. A putative group II intron maturation in the trnK (lysyl tRNA) gene (matK) of angiosperm chloroplasts has been implicated in splicing of its own and other introns.
group II introns (du Jardin et al. 1994; Liere and Link 1995; Vogel et al. 1997, 1999); however, because only binding of the protein to its intron was reported (Liere and Link 1995), the implication is that it does not promote splicing of the trnK intron by itself. The fact that we observed ORF-dependent splicing of Chs.psbA1 in E. coli bodes well for possibly overproducing this protein for biochemical studies. SDS-PAGE analysis of total soluble protein from the wild-type Chs.psbA1 transformants did not reveal an abundant polypeptide of the size expected for the ORF, suggesting that production of the protein using its own translation signals was rather low (O. Odom and D. Herrin, unpubl. results). This may explain why splicing of wild-type Chs.psbA1 in E. coli was somewhat inefficient.

The only group II intron maturase that has been studied biochemically is from the group IIA intron, Ll.LtrB, of the bacterium L. lactis. The intron-ORF protein, LtrA, binds with high affinity to subdomain IVa, which includes the beginning of the LtrA coding region (Wank et al. 1999). Secondary contacts are then made to conserved regions of the intron core, with the effect of promoting tertiary folding of the intron (Matsuura et al. 2001). A predicted fold of domain IV in Chs.psbA1 (Fig. 2) reveals a possible IVa subdomain that also contains the putative Shine–Dalgarno sequence, but it is considerably reduced compared to LLtrB. Moreover, we could not find an analogous structure upstream of the ORFs in the related Calothrix and Nostoc introns, suggesting that group IIB maturases may bind their introns differently than LtrA. In this respect, we also note that the IVa subdomain of the Saccharomyces cerevisiae mitochondrial COXI a2 group IIA intron seems to play a reduced role in maturase-dependent splicing of this intron in vivo, although it is critical for a2 homing (Huang et al. 2003).

To our knowledge, the Chs.psbA1 intron is the first ribozyme described from a psychrophile. The ion requirements for Chs.psbA1 self-splicing are similar to those of other group II introns in requiring fairly high Mg$^{2+}$ and high NH$_4^+$, although the minimal Mg$^{2+}$ requirement (13 mM in the absence of a polycation) appears to be lower than for some group II introns (Jarrell et al. 1988; Hebbar et al. 1992; Adamidi et al. 2003). Evidence was also presented for productive, cotranscriptional folding of the pre-RNA in vitro. The proposed folding is likely only partial, however, because splicing did not occur in the transcription reactions. Cotranscriptional folding is presumably what happens in vivo, and could help prevent the formation of some nonproductive isomers that form during denaturation–renaturation cycles in vitro. Cotranscriptional folding has been shown to improve group I intron self-splicing (Heilmann-Miller and Woodson 2003).

Self-splicing of Chs.psbA1 occurred via a lariat; however, there could also have been a fraction of the pre-RNA that spliced via a hydrolysis mechanism. There is a band in the gels of $^{32}$P-labeled RNA splicing reactions that might represent excised linear intron (e.g., Fig. 4B). We did not try to quantify this band, but it is clearly weaker than the intron lariat band. The A residue utilized in the lariat pathway is in domain VI, its usual location, although it is somewhat atypical in that it is in an asymmetric internal loop (A$^{2563}$ in Fig. 2). The rates of self-splicing of the ΔORF and ΔORF.1 pre-RNAs via the lariat mechanism are in the range of other self-splicing group II introns (Daniels et al. 1996; Adamidi et al. 2003). With the most active preparation (ΔORF.1, unextracted), 25–30% of the pre-RNA self-spliced. Thus, the extent of Chs.psbA1 self-splicing seems to be somewhat less than the more robust group II introns from mitochondria and prokaryotes (Daniels et al. 1996; Costa et al. 1997; Adamidi et al. 2003). With the full-length Chs.psbA1 intron, only 4%–5% of the pre-RNA spliced, suggesting that Chs.psbA1 is an inefficient folder. It is possible that inefficient folding to the active state reflects an adaptation to low temperature. If increased flexibility enhances ribozyme function at low temperature as it does for protein enzymes (Hochachka and Somero 2002), then such flexibility could manifest itself as a reduction in stability of the catalytically active state of the ribozyme. Of course that raises the question of how splicing is made efficient in vivo.

The temperature optimum for Chs.psbA1 self-splicing was determined to be ~44°C for the different forms of the pre-RNA. This seems to be similar to, or somewhat lower in some cases, than other group II introns (Jarrell et al. 1988; Winkler and Kück 1991; Hebbar et al. 1992; Costa et al. 1997; Adamidi et al. 2003). On the other hand, the poor self-splicing of Chs.psbA1 at 50°C suggests that Chs.psbA1 is more thermolabile than other group II introns. It may also be significant that there is a shoulder on the cold side of the temperature peak (Fig. 6), and that self-splicing of Chs.psbA1 was clearly detected at low (<20°C) temperatures. In contrast, Hebbar et al. (1992) reported that the yeast a1 intron was essentially devoid of self-splicing activity at 30°C. It is also interesting to note that self-splicing of the Pylaella littoralis mitochondrial rRNA intron, PlLSU/2, was readily detected at 25°C (Costa et al. 1997). Although not shown to be a true psychrophile, to our knowledge, P. littoralis is common in northern temperate waters (Voipio 1987; Milchakova 1999). We conclude that this ribozyme shows some evidence of adaptation to low temperature, but the adaptation is not as dramatic as with many protein enzymes (e.g., Loppes et al. 1996; Irwin et al. 2001), which often show great differences in temperature optima between mesophilic and psychrophilic homologs. However, additional ribozymes from psychrophilic organisms need to be examined before general conclusions are drawn.

The analysis of psbA transcripts in CCMP-1619 indicated that Chs.psbA1 splicing is efficient in these cells at 8°C. Thus, it seems likely that, in vivo, proteins function to promote splicing of this intron ribozyme at low temperature. As indicated here, the intron-encoded ORF appears to be one such factor, but there could be others (Lambowitz et al.
solution (0.5 mL/L) and ampicillin (100 µg/mL) were added after MES (0.2 g/L; pH 6.8). Filter-sterilized f/2 (Guillard 1975) vitamin fold, using Hutner’s trace metals (1 mL/L), and buffering with chloroplast genes of Chlamydomonas spp. are evolutionarily derived from cyanobacterial-like organisms. However, chloroplasts are evolutionarily derived from cyanobacterial-like organisms. Thus, CCMP-1619 should not require as tight a control over psbA expression as does C. reinhardtii.

Another interesting evolutionary question concerns the origin of the Chs.psbA1 intron. The similarity of this intron to several introns from cyanobacteria is intriguing, because chloroplasts are evolutionarily derived from cyanobacterial-like organisms. However, psbA genes in most photosynthetic organisms, including Calothrix and Nostoc, lack group II introns. It was suggested previously that the Chs.psbA1-related intron from the E. myxocylindracea chloroplast psbA gene was acquired horizontally from a cyanobacterium (Sheveleva and Hallick 2004). The chloroplast genome of Euglena spp. is believed to be of green algal origin, having been acquired by secondary endosymbiosis. Because Sheveleva and Hallick (2004) found that several other Euglena species tested lack this intron in their psbA gene and we have also found that the Chs.psbA1 intron is lacking from several other species of Chlamydomonas that were examined (O. Odom, D. Shenkenberg, and D. Herrin, unpubl. results), it seems unlikely that these introns were acquired vertically from a common chloroplast ancestor. We suggest, therefore, that the Chs.psbA1 intron was acquired by an independent horizontal transfer from a cyanobacterium. This suggestion is also supported by the fact that the GC content of the Chs.psbA1 intron (45%) is much closer to the genes of Nostoc and Calothrix (43%) than to chloroplast genes of Chlamydomonas spp. (34%). It is likewise intriguing that Nostoc spp. are found at Lake Bonney, Antarctica (Paerl et al. 2000). In summary, this group II intron (or a close relative) has apparently jumped across the prokaryotic–eukaryotic divide at least twice. It will be interesting to see if it is a mobile intron.

MATERIALS AND METHODS

Cell culture

Chlamydomonas sp. CCMP-1619 was obtained from the Provasoli-Guillard Center and C. subeaudata (SAG 12.87) from the Götingen Collection. The growth medium was soil–water extract (Pringsheim 1946) or Sager–Granick minimal medium (Harris 1989) modified by eliminating FeCl₃, reducing MgSO₄•H₂O three-fold, using Hutner’s trace metals (1 mL/L), and buffering with MES (0.2 g/L; pH 6.8). Filter-sterilized E2 (Guillard 1975) vitamin solution (0.5 mL/L) and ampicillin (100 µg/mL) were added after autoclaving. For RNA analysis, the cells were grown at 8°C under continuous illumination (60 µmoles/m²/sec) and slow bubbling with 5% CO₂.

Isolation of algal DNA, PCR, and DNA constructs

Total nucleic acids (TNA) were isolated from liquid cultures as described (Durrenberger et al. 1996), except RNase A was omitted. PCR amplification of psbA used primers 176 (TTTACCTCCTGA CCGTATGCTTATTAGG) and 100 (TAGGATCCACGACTTAA GAAGAGTGTAATG), 30 ng TNA, Taq DNA polymerase, and an annealing temperature of 60°C. The 2.9-kb product was sequenced by primer walking (accession no. AJ325305). Amplification of the petA-petD region was performed with primers 330 (GTGTTTACCAGATACGTTTTTG) and 332 (CCATACCTTAGGATATTAGCTT) and an annealing temperature of 45°C. The 1.9-kb product was sequenced (the CCMP-1619 sequence was deposited in GenBank, accession no. AY332410). The transcription template containing the full-length intron and partial flanking exons (wild-type) was synthesized by PCR of Chlamydomonas sp. CCMP-1619 TNA with primers 310 (TAAAGCTTCCTACATTATA GGGATCCGGGTGTTCATTCTCTGACGTTAAGCTTATTAGG) and either 177 (GTGATGATGAAAGAATGAAAGC) or 100 (GCTGTTTTACCAGATACGTTTTTG) and either 177 (GTGATGATGAAAGAATGAAAGC) or 100 (GCTGTTTTACCAGATACGTTTTTG) and either 177 (GTGATGATGAAAGAATGAAAGC) or 100 (GCTGTTTTACCAGATACGTTTTTG) and either 177 (GTGATGATGAAAGAATGAAAGC) or 100 (GCTGTTTTACCAGATACGTTTTTG) and either 177 (GTGATGATGAAAGAATGAAAGC) or 100 (GCTGTTTTACCAGATACGTTTTTG). The ORF-deleted (ΔORF) DNA was generated as follows. The 310/177 PCR product was digested with BstBI and StuI, giving fragments of 872, 1441, and 435 bp. The 435-bp fragment was digested with HpaII, which yielded fragments of 134 and 301 bp, and the 301-bp fragment was ligated to the 872-bp fragment via the complementary 5’-CG overhangs generated by HpaII and BstBI. An aliquot of the ligation mixture was used for PCR with primers 310 and 177, and the expected 1.2-kb product was sequenced. The ΔORF.1 plasmid was generated by cloning the ΔORF DNA into pluBluescript SK+ after an additional round of PCR with primers 322 (TAGGATACCTCATTCTCTGACGTTA AGCTTATTAGG) and 323 (TAAAGCTTCGATGAAATGAAAGC) and digestion with KpnI and HindIII. A ΔORF.1 clone was identified and the insert sequenced. The constructs for analyzing Chs.psbA1 splicing in E. coli were obtained by PCR of Chlamydomonas sp. CCMP-1619 TNA with exon primers 322 and 323 and cloning into Bluescript SK+. Sequencing of several clones did not yield one without mutations, so one was created by exchanging Nhel-KpnI fragments.

In vitro RNA synthesis and self-splicing

RNA was synthesized from PCR products or HindIII-digested plasmid DNA (200 ng) with T7 RNA polymerase in 20-µL volume. After 1 h at 37°C, the reactions were either frozen at −70°C or extracted with chloroform/EDTA and precipitated with ethanol. Standard self-splicing reactions (5 µL) contained ~80 ng RNA, 40 mM Tris–HCl (pH 7.5), 30 mM MgCl₂, 1.3 M NH₄Cl, and were incubated at 42°C for 90–180 min; control reactions were incubated at 0°C. Variations in these conditions are indicated in the text. The reactions were stopped with EDTA and cooled on ice. For gel analysis, RNAs that had not been extracted previously were extracted with chloroform, and all samples were precipitated with ethanol. The 32P-labeled RNAs were separated on denaturing 4% polyacrylamide gels and exposed to X-ray film (BioMax MS) or to a phosphorimager screen that was quantified with ImageQuant (Molecular Dynamics).
The kinetics of self-splicing of ΔORF and ΔORF.1 pre-RNAs were determined at 41°C by quantifying gels with a phosphorimager after verifying the integrity of the samples. Semi-log plots of precursor decay were linear for over 100 min, indicative of a single kinetic component. The observed rate constants (k_{obs}) were determined from these plots (after correcting for the fraction of active ribozyme, which was determined from a 4-h incubation) or from early time points when the rate of splicing was nearly linear.

**Splicing in E. coli**

Plasmids were transformed into the *E. coli* Rosetta(DE3)pLysS strain (Novagen), and the appropriate clones verified by restriction analysis. Selected transformants were grown in Luria Broth at 37°C to an A_{600} of 0.7, and then induced at 30°C with IPTG (1 mM). After 2 h, RNA was isolated as described previously (Herrin and Schmidt 1988), except for resuspending the cells in lysozyme (0.5 mg/mL) plus 200 U/mL Superase·In (Ambion), and including two acid–phenol extractions. Most of the residual DNA was removed by degradation with TurboDNase (Ambion).

**RT-PCR**

Reverse transcription was performed with either 2–3 µg of *E. coli* RNA or ~2 ng of self-spliced RNA in 20 µL, described as prepared by the manufacturer; an aliquot (1 µL) was used for PCR (30 cycles). To examine in vitro splicing, primer 177 was used, and for splicing in *E. coli*, a T3 promoter primer (GCCGCAGATTAACCCCTAAGGGA) was used for reverse transcription, and then the same primer plus 176 was used for PCR (annealing at 60°C). The products (either 148 or 256 bp, respectively) were sequenced using the same primers. To map the intron branch point (Vogel et al. 1997), reverse transcription was performed with primer 313 (GTTCCTTACGTGAGGGGTGAGTACT TAG), then PCR was carried out with the nested primers 316 (CCACGAGTTGGTATGGGTAGATTCAG) and 317 (GAGCGTAGATGAGCAAAGTGACAG). The product (109 bp) was sequenced using the PCR primers. RT–PCR products were analyzed and purified either by ethidium bromide–agarose gel electrophoresis or by polyacrylamide gel electrophoresis and SYBR Green I staining (Molecular Probes).

**Algal RNA isolation and Northern blot hybridization**

RNA was isolated from *Chlamydomonas* and Northern blotted as described (Herrin and Schmidt 1988). The exon-specific probe (343 bp) was obtained by RT-PCR of self-spliced pre-RNA (wild-type) with primers 100 and 176. The *Chs.psbA1* intron—internal probe (731 bp) was obtained by PCR of TNA with primers 312 (CTAGGCTATACCTATCAGG) and 313. The probes were labeled by single-sided PCR with α^{32}P-dCTP (3000 Ci/mmol), and the blots were hybridized and washed at 65°C (Church and Gilbert 1984).

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Chloroplast group II intron splicing


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A horizontally acquired group II intron in the chloroplast psbA gene of a psychrophilic *Chlamydomonas*: In vitro self-splicing and genetic evidence for maturase activity

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