An inorganic carbon transport system responsible for acclimation specific to air levels of CO₂ in *Chlamydomonas reinhardtii*

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Many photosynthetic microorganisms acclimate to CO₂ limited environments by induction and operation of CO₂-concentrating mechanisms (CCMs). Despite their central role in CCM function, inorganic carbon (Ci) transport systems never have been identified in eukaryotic photosynthetic organisms. In the green alga *Chlamydomonas reinhardtii*, a mutant, *pmp1*, was described in 1983 with deficiencies in Ci transport, and a Pmp1 protein-associated Ci uptake system has been proposed to be responsible for Ci uptake in low CO₂ (air level)-acclimated cells. However, even though *pmp1* represents the only clear genetic link to Ci transport in microalgae and is one of only a very few mutants directly affecting the CCM itself, the identity of Pmp1 has remained unknown. Physiological analyses indicate that *C. reinhardtii* possesses multiple Ci transport systems responsible for acclimation to different levels of limiting CO₂ and that the Pmp1-associated transport system is required specifically for low (air level) CO₂ acclimation. In the current study, we identified and characterized a *pmp1* allelic mutant, *air dier 1* (ad1) that, like *pmp1*, cannot grow in low CO₂ (350 ppm) but can grow either in high CO₂ (5% CO₂) or in very low CO₂ (<200 ppm). Molecular analyses revealed that the Ad1/Pmp1 protein is encoded by *LciB*, a gene previously identified as a CO₂-responsive gene. LciB and three related genes in *C. reinhardtii* compose a unique gene family that encode four closely related, apparently soluble plastid proteins with no clearly identifiable conserved motifs.

bicarbonate | CO₂-concentrating mechanism | microalgae | photosynthesis

Although present in small quantities in the air, carbon dioxide (CO₂) has profound influences on the living environments by serving as the major substrate for photosynthesis. In nature, the ambient CO₂ concentrations for photosynthetic organisms can vary across orders of magnitude and often become the limiting factor for carbon acquisition. Many aquatic photosynthetic microorganisms use a CO₂-concentrating mechanism (CCM) to maximize photosynthesis under limiting CO₂ conditions. Photosynthesis of these microorganisms grown in limited CO₂ environments displays a characteristic similar to that in C₄ photosynthesis, with much higher apparent affinity for CO₂ (1, 2). However, unlike the CO₂ enrichment in C₄ plants, CCMs in aquatic photosynthetic microorganisms operate by accumulating a large amount of dissolved inorganic carbon (Ci; CO₂ and/or bicarbonate) intracellularly, the uptake of which is driven by energy-coupled Ci transport systems.

As vital components of CCMs, Ci transport systems have been extensively studied in the prokaryotic model organisms, cyanobacteria. With the aid of mutant studies and the recent availability of several genomes, at least four transport modes involved in Ci uptake have been identified and characterized in cyanobacteria, including two bicarbonate transporters and two CO₂ uptake systems associated with the operation of specialized NDH-1 complexes (3). However, little information other than its physiological demonstration is available regarding Ci transport in eukaryotic photosynthetic microorganisms. The unicellular green alga *Chlamydomonas reinhardtii* has served as a key model system to study CCMs for many years, and several genes required for acclimation to limiting CO₂ have been cloned and characterized in this organism (4–10), but no transport system for Ci uptake has been definitively identified and characterized. Even though specific defects in several mutants requiring elevated CO₂ for survival have been identified, only three characterized mutants can be argued as having defects in genes required unambiguously for operation of the CCM, and one of these mutants, *cia5* (and various alleles, including *ccm1*), appears to be defective in a master regulator (Cia5 or Ccm1) for induction of the CCM and other proteins required for limiting CO₂ acclimation (4, 6, 11), rather than a functional component of the CCM. Another key mutant, *cal* (and various alleles, including *cia3*), corresponds to a thylakoid lumen carbonic anhydrase (Cah3) apparently required for the rapid dehydration of intracellular bicarbonate accumulated by active Ci transport (5, 7, 12). The third of these key mutants, *mpm1*, was characterized more than two decades ago as being impaired in Ci transport (13) and, thus, far represents the only mutant identified with a specific defect in Ci transport in a eukaryotic photosynthetic organism. These mutants form the foundation for our understanding of the *C. reinhardtii* CCM, demonstrating the requirement for active Ci transport (*mpm1*) to accumulate intracellular Ci and a thylakoid lumen CA (*cal*) for dehydration of the intracellular Ci accumulated as a bicarbonate.

Among these classic mutants, the defective gene in *mpm1* and the identity of Pmp1 protein, thus far, have resisted identification. Although initially identified as a probable Ci transport mutant, a recent study reported that the expression profiles of several CO₂ responsive genes in *mpm1* differ from those in wild type and suggested that the *Pmp1* gene product might regulate the expression of Ci transporter genes (14). Another recent observation regarding *mpm1* is its unusual, air-dying phenotype (15, 16): it grows well in either high (5%) or very low (<200 ppm) CO₂ but dies in low (air-level) CO₂ (350–450 ppm). This conspicuous phenotype distinguishes *mpm1* from most other high CO₂-requiring mutants and indicates the existence of multiple Ci transport systems in *C. reinhardtii* corresponding to multiple CO₂ level-dependent acclimation states. Indeed, at least three distinct CO₂ acclimation states have been demonstrated in *C. reinhardtii*, corresponding to: high CO₂, ≥0.5% CO₂; low CO₂, 0.4–0.03% CO₂; and very low CO₂, ≤0.01% CO₂ (17). Therefore, the Pmp1 protein must play either a functional or a regulatory role in a Ci transport system specific for the low (air-level) CO₂ acclimation state.

To understand the mechanism of limiting CO₂ acclimation in eukaryotic photosynthetic organisms, we have taken an insertional mutagenesis approach to identify functional components
involved in limiting CO2 acclimation in C. reinhardtii. Here we describe the identification and characterization of a mutant that displays an air-dying phenotype, \textit{air dier} (\textit{ad1}). Our results demonstrate that the defective gene in \textit{ad1} is allelic to \textit{pmp1} and that it belongs to a small family of genes encoding an apparently unique group of proteins in \textit{C. reinhardtii}.

**Results**

**Identification of the \textit{air dier} 1 (\textit{ad1}) Mutant.** To isolate and identify \textit{C. reinhardtii} mutants unable to acclimate to air levels of CO2, we performed insertional mutagenesis by using a \textit{BleR}\textsuperscript{R} containing plasmid (pSP24s) (18) to transform a wallless, wild-type strain, \textit{cw10}. The \textit{air diering} phenotype was evaluated in spot tests, based on the ability to grow in high, low (air-level), or very low CO2 concentrations. From \(\approx 2,500\) transformants, two mutants displayed the “\textit{air dier}” (\textit{ad}) phenotype, and one, \textit{ad1}, was selected for further investigation. As with \textit{pmp1}, \textit{ad1} and wild type grow in either high CO2 or very low CO2 but dies in low (air-level) CO2 (Fig. 1). In contrast, another classic mutant, \textit{cia5}, previously identified as defective in acclimation responses to limiting CO2, grows similar to wild type in high CO2, somewhat more slowly in low (air-level) CO2, but dies in very low CO2.

Photosynthetic O\textsubscript{2} evolution in response to Ci concentrations for low CO2 acclimated and very low CO2 acclimated wild-type and \textit{ad1} cells was compared (Fig. 2 A and B). The \textit{ad1} mutant cells acclimated in low CO\textsubscript{2} showed dramatically decreased photosynthetic Ci affinity compared with wild-type cells grown under the same conditions. In contrast, when acclimated to very low CO\textsubscript{2}, the photosynthetic O\textsubscript{2} evolution of \textit{ad1} and wild type exhibited similar responses to external Ci concentrations. Furthermore, \textit{ad1} acclimated to low CO\textsubscript{2} exhibited dramatically reduced Ci accumulation compared with wild type (Fig. 2C), similar to that observed with \textit{pmp1} (13). These results demonstrate that the \textit{air dier} phenotype in \textit{ad1} is due to an impaired photosynthetic affinity in low (air-level) CO\textsubscript{2}, which is caused by a deficiency in Ci transport in this CO\textsubscript{2} concentration.

**Identification of the \textit{Ad1} Gene.** The \textit{ad1} strain was crossed with wild-type strain CC620 to determine whether the \textit{air dier} phenotype in the \textit{ad1} mutant cosegregated with the inserted \textit{BleR}\textsuperscript{R} gene. More than 100 random progeny were tested for their growth in different levels of CO\textsubscript{2} and their resistance to zeocin, which indicates the presence of the \textit{BleR}\textsuperscript{R} insert. Although 50 random progeny with the \textit{air dier} phenotype all exhibited zeocin resistance, all zeocin-sensitive progeny showed wild-type growth in low CO\textsubscript{2}, indicating cosegregation of the \textit{air dier} phenotype with the \textit{BleR}\textsuperscript{R} insert. DNA gel blot analysis with probes specific for the \textit{BleR}\textsuperscript{R} gene and pBluescript sequences indicated a single insert present in \textit{ad1} (data not shown).

DNA flanking the \textit{BleR}\textsuperscript{R} gene in \textit{ad1} was cloned from \textit{ad1} genomic DNA by inverse PCR. This sequence was used in a BLAST search against the \textit{C. reinhardtii} genome, and the insertion site was shown to be located on scaffold 4 of the genome draft (version 3.0 of the \textit{C. reinhardtii} genome, http://genome.jgi-psf.org/Chlre3/Chlre3.home.html). Further PCR and DNA gel blot analyses revealed a large deletion of a large segment of the genomic DNA sequence in \textit{ad1} at the site of insertion, presumably caused by the integration of the \textit{BleR}\textsuperscript{R} insert. DNA gel blot analysis with probes specific for the \textit{BleR}\textsuperscript{R} gene and pBluescript sequences indicated a single insert present in \textit{ad1} (data not shown).

A sequence flanking the \textit{BleR}\textsuperscript{R} insert was used as probe to identify \textit{C. reinhardtii} bacterial artificial chromosome (BAC) clones containing wild-type genomic DNA overlapping the site of the insertion, and DNA from two identified clones was demonstrated to complement the \textit{air dier} phenotype of \textit{ad1}. Using Southern and PCR analyses to determine whether any of the predicted genes located within the deleted region were present in complemented lines, only one gene, \textit{LciB} (14), was found to be present in all BAC complemented \textit{ad1} lines, and other predicted genes either were not present at all or were present only in some complemented lines (data not shown).

To confirm whether the \textit{LciB} gene is the \textit{Ad1} gene, we PCR-amplified a genomic DNA fragment containing \textit{LciB} from a wild-type BAC and found the DNA fragment could complement \textit{ad1}. Complemented \textit{ad1} lines and wild type grew in both low and very low CO\textsubscript{2} (Fig. 1), and Southern analysis indicated that all putative complemented lines carried the genomic DNA.
of the LciB gene, which is absent from the ad1 mutant (Fig. 3B). In addition, RNA gel blot analysis showed that all complemented lines recovered the expression of LciB (Fig. 3C). Complementation of ad1 also was achieved by expressing an LciB cDNA under control of the constitutive PsaD promoter and terminator (data not shown).

ad1 Is a pmp1 Allele. The ad1 mutant appears very similar to pmp1 in its air-die phenotype and impaired photosynthesis and Ci transport. Miura et al. (14) reported that induction of three CO2-responsive genes by limiting CO2 was abolished in pmp1. These genes include two putative Ci transporter genes, LciA and Mrp1, and LciB, which also was suggested by Miura et al. (14) to be a putative Ci transporter gene. When we compared the low CO2-induced expression of LciA and Mrp1 in ad1 and pmp1 with that in wild type (Fig. 4), we found that expression of both LciA and Mrp1 was induced in all three strains upon exposure of high CO2-grown cells to either low (air-level) CO2 or very low CO2. Although we did observe a slightly reduced expression of LciA and Mrp1 in ad1 and pmp1 to a variable extent, it seems unlikely that this is a direct result of the lesion in LciB. The failure of Miura et al. (14) to observe the induction of LciA and Mrp1 gene expression in pmp1 might be explained by their short induction time (2 h) and different growth conditions. Nevertheless, our results demonstrate ad1 is very similar to pmp1 in its expression of these putative Ci transporter genes and that any reduced induction of LciA and Mrp1 in pmp1 or ad1 is relatively minor and likely to be pleiotropic.

Crossovers between ad1 and pmp1 failed to produce recombinants or diploids with a wild-type growth phenotype, suggesting that ad1 is likely to be a pmp1 allele. This conclusion was confirmed by the complementation of pmp1 with LciB. Both the genomic and cDNA forms of LciB that complemented ad1 also successfully complemented pmp1 (Fig. 1). Comparison of the DNA sequence of LciB from pmp1 and that from wild type revealed a point mutation (C > A) at nucleotide position 105 in pmp1. This mutation would result in a stop codon in place of tyrosine at amino acid 35 of the wild-type gene product and, therefore, result in an extremely truncated LciB gene product in pmp1 (Fig. 5).

LciB Gene Family. BLAST searches and domain searches of several databases with LciB revealed no significant recognizable domains nor significant homologies, except for three additional genes in the C. reinhardtii genome (http://genome.jgi-psf.org/Chlre3/Chlre3.home.html; Fig. 5): a similar CO2 responsive gene, LciC (14), on scaffold 12, and two previously unreported genes, LciD and LciE, both on scaffold 15. As noted by Miura et al. (14), the LciB and LciC gene products are predicted to be soluble proteins, probably targeted to the plastid. This situation also is the case for LciD and LciE. However, considering the low sensitivity of currently available prediction tools for discriminating subcellular targeting for C. reinhardtii proteins, especially for differentiating between plastid and mitochondrion proteins, one cannot exclude the possibility that the products of LciB gene family are mitochondrion localized.

The LciB and LciC gene products are quite similar in their predicted amino acid sequence (57% identity; 73% similarity), as are LciD and LciE (71% identity; 78% similarity), with these two protein pairs also sharing substantial similarity with each other (40–44% identity; 62–65% similarity), thus constituting an LciB gene products are predicted to be soluble proteins, probably targeted to the plastid. The inverted regions flank another set of CO2 responsive genes, Cahl and Cuh2 (20), forming a cluster of six CO2 sensitive genes within a 75-kb region on scaffold 15. The significance of this arrangement, if any, is not clear.

Genes of the LciB Family as CO2 Responsive Genes. Miura et al. (14) demonstrated that LciB and LciC were up-regulated by limiting CO2. Because the two genes share high similarity in their coding sequence, we used gene-specific 3′ UTR probes for LciB and LciC in Northern blot analyses and showed that the LciB and LciC genes had very similar patterns of limiting CO2-induced expression (Fig. 6). In wild type, both genes showed constitutive expression with low mRNA abundance under high CO2 conditions, whereas both mRNA levels increased dramatically when cells were transferred into either low CO2 (400 ppm) or very low CO2 (100 ppm; Fig. 6), or into a level of CO2 (1,500 ppm) intermediate between low and high CO2 (data not shown), indicating that up-regulation of LciB and LciC expression was not confined to a specific level of limiting CO2.

There were no ESTs available for LciD and LciE, suggesting that they either are genes with low expression or are genes expressed under conditions different from those used for EST identification and from those inducing LciB/LciC expression. We amplified predicted 3′ UTR sequences of LciD and LciE from a cDNA library based on their predicted genomic se-
quences and used these specific, amplified probes to analyze the expression of these genes. On RNA gel blots, the \( \text{LciD} \) gene showed two bands hybridizing to the \( \text{LciD} \) probe (Fig. 6) and a limiting CO2-inducible expression pattern similar to those of \( \text{LciB} \) and \( \text{LciC} \) but with relatively lower mRNA abundance. \( \text{LciE} \) expression was not detectable on RNA gel blots by using a predicted \( \text{LciE} \)-specific probe, although we did successfully verify this 3’/H11032 UTR and the expression of \( \text{LciE} \) by amplifying a partial \( \text{LciE} \) cDNA from a cDNA library and subsequently sequencing the PCR product. Attempts to identify full-length \( \text{LciD} \) and \( \text{LciE} \) cDNAs by screening a cDNA library with predicted \( \text{LciD} \) and \( \text{LciE} \) coding region probes yielded five cDNA clones, all of which were determined by sequencing to be from \( \text{LciD} \). Comparison of 3’/H11032 UTR sequences of the \( \text{LciD} \) cDNA clones revealed two 3’ UTR sequences with different lengths, indicating that alternative termination occurred, possibly explaining the two \( \text{LciD} \)-hybridizing bands on gel blots. Because \( \text{LciE} \) expression was undetectable in Northern analyses and attempts to identify a full-length \( \text{LciE} \) cDNA from the cDNA library were not successful, it appears that the expression of \( \text{LciE} \) may be relatively low compared with other genes in this family, at least under the conditions explored in this work and under conditions used for construction of the cDNA library.

Discussion

\( \text{Ad1/Pmp1}: \) Transporter or Regulator? Despite being major components in the CCM, Ci transport systems in eukaryotic photosynthetic organisms remain largely unknown. Since its identification more than two decades ago, the \( \text{pmp1} \) mutant has been touted as demonstrating a Ci transport requirement in the CCM (1, 13). In this work, we generated a new mutant allele of \( \text{pmp1} \), \( \text{ad1} \), by insertional mutagenesis, and identified the \( \text{Ad1}/\text{H20862 Pmp1} \) gene. We demonstrated that a lesion in \( \text{LciB} \) (\( \text{Ad1}/\text{H20862 Pmp1} \)) caused the air deficient phenotype and greatly decreased Ci transport and photosynthetic activity in \( \text{ad1} \) and \( \text{pmp1} \), presumably only in low CO2.

Physiological and biochemical characterization of \( \text{pmp1} \) suggests that Pmp1 is a functional component involved in Ci transport. Although both \( \text{pmp1} \) and \( \text{ad1} \) also have reduced CO2 assimilation in low CO2, the dramatically decreased internal Ci accumulation in low CO2 argues strongly for a defect in Ci transport rather than internal Ci utilization. However, being predicted to be a soluble protein with no obvious transmembrane regions, \( \text{LciB} \) (\( \text{Pmp1/Ad1} \)) seems unlikely to perform as an intact active Ci transporter by itself, because hydrophobic transmembrane domains are signatures for almost all identified transporters. Alternatively, Pmp1 as a general regulator for multiple Ci uptake systems also has been suggested. Miura et al. (14)
reported the lack of induction or up-regulation of several putative transporter genes in pmpl1, including LciB itself. These authors therefore proposed that Pmp1 is involved in regulation of multiple Ci transporters. In the current study, we have shown that transcripts of these putative transporters still were present in pmpl1 and ad1 (except LciB in ad1), although we did observe that their mRNA abundance in low CO2 often was reduced in both the ad1 and pmpl1 mutants, but to a variable extent. Because LciB is predicted to localize in plastids (or possibly mitochondria), this protein is not expected to be a transcription factor and directly involved in transcription regulation, like Cia5. If LciB affects the synthesis of new transcripts or the stability of these putative transporter transcripts, it is more likely that it affects these processes in an indirect way.

However, given the probable plastid localization of LciB and the nonreproducibility of the decreased expression of putative Ci transporters in pmpl1/ad1, the direct involvement of LciB in Ci uptake seems more plausible than the regulation by LciB of the expression of other Ci transporter genes. In fact, the physiological evidence from pmpl1/ad1 for nearly a complete lack of Ci transport, even though the expression of Mrp1 and LciA still is present, argues for a direct involvement of LciB with Ci transport.

It is not clear why a defect in only one gene (LciB itself) from the LciB gene family causes the air dier phenotype in ad1 and pmpl1 despite the high sequence similarity and similar limiting CO2-inducible expression patterns among the genes in the LciB family, especially the strong similarities between LciB and LciC. It is possible that interaction of LciC and LciB is required for a functional transporter complex, or a regulator complex for a Ci transporter(s), and this possibility should be investigated.

Acclimation to Multiple Levels of CO2. The photosynthesis of ad1 was impaired only in low (air-level) CO2-acclimated cells, which apparently is caused by defective Ci transport. In very low CO2-acclimated cells, photosynthesis of ad1 recovered to a level similar to that in wild type. These results confirm the existence of distinct states for very low CO2 acclimation and low (air-level) CO2 acclimation in C. reinhardtii. Therefore, limiting CO2 acclimation in C. reinhardtii must require at least two (probably non-overlapping) suites of proteins that are differentially expressed or activated in different levels of limiting CO2. LciB obviously is associated with and required for the low (air-level) CO2 acclimation. Photosynthetic measurements also showed that very low CO2-acclimated cells have a relatively higher affinity for Ci but lower photosynthesis at near-saturated Ci concentrations, relative to cells acclimated to low (air-level) CO2. This observation is consistent with a recent report on different physiological states for limiting CO2 acclimation in C. reinhardtii, in which very low CO2-acclimated cells exhibited lower $K_c$ (CO2) and $V_{max}$ compared with low CO2-acclimated cells (17). The difference in $K_c$ (CO2) and $V_{max}$ between low CO2-grown cells and very low CO2-grown cells implies that the Ci transport system specific for low (air-level) CO2 has a relatively lower affinity for Ci but higher transport capacity, whereas the system specific for very low CO2 has a higher affinity for Ci but a lower capacity. This acclimation may represent an excellent survival strategy in C. reinhardtii for acclimation to different levels of limiting CO2: In very low CO2, a Ci uptake system with a high affinity and low capacity would allow C. reinhardtii cells to grow at a reasonable rate without depleting all available Ci, whereas in low (air-level) CO2, a high capacity for Ci uptake could maintain optimal growth, and a transporter with relatively low affinity would be sufficient to accommodate the Ci uptake in low CO2.

Identification of the defect responsible for the Ci transport deficiency in pmpl1 and ad1 represents a critically important step toward understanding Ci transport, its role in the CCM, and its regulation in eukaryotic microalgae. It clearly will be important to fully understand the role of LciB and the other members of the LciB gene family in limiting CO2 acclimation, including any role they may play in distinguishing the low CO2 and very low CO2 acclimation states. The majority of past research on limiting or low CO2 acclimation in C. reinhardtii and other microalgae has focused mainly on air level CO2 acclimation, whereas targeted research on very low CO2 acclimation has been limited. Future investigation of this distinct state should help fill the gap in our understanding of the multiple levels of CO2 acclimation in C. reinhardtii or other eukaryotic photosynthetic cells.

Materials and Methods

C. reinhardtii Strains, Culture, and Gas Conditions. C. reinhardtii strains CC849, CC620, and CC125 were obtained from the Chlamydomonas Genetics Center, Duke University, Durham, NC. The pmpl1 and cia5 mutants have been described in refs. 11 and 13. Wild-type cells and high CO2-requiring mutants were maintained on agar plates with CO2 minimal medium (21) and kept in Plexiglas chambers at room temperature. Liquid cultures were grown in Erlenmeyer flasks on an orbital shaker at 125 rpm. In both plate and liquid cultures, continuous gas flow was maintained through either the growth chambers or the culture flasks. Three gas conditions used in this study were: high CO2 (5% CO2 in air vol/vol), obtained by mixing compressed CO2 with normal air; low CO2 (normal air, 350–400 ppm); and very low CO2 (50–150 ppm), obtained by mixing normal air with either compressed CO2-free air or CO2-depleted air (air passed through a saturated sodium hydroxide solution).

Isolation of air dier Mutants, Growth Spot Tests, and Genetic Analysis. C. reinhardtii wall-less strain CC849 (cw10, mt−) was transformed with linearized pSP124s plasmid (ref. 18; a gift from Saul Burton, University of London, London) by the glass bead method (22). Transformed cells were kept in high CO2 and selected on minimal medium plates supplemented with 10 µg/ml zeocin. Zeocin-resistant transformants were transferred to duplicate plates for screening by growth spot tests in high CO2, low CO2, and very low CO2. Mutants exhibiting an air dier phenotype were maintained in the high CO2 chamber. Spot growth tests were performed by suspending actively growing cells in minimal medium to similar, low-cell densities (<105 cells/ml), then spotting 3 µl of each cell suspension onto agar plates and kept in high CO2, low CO2, or very low CO2 chambers for 8 days.

Genetic crosses and tetrad analyses were performed as described by Harris (23).

Photosynthetic O2 Evolution and Ci Uptake. Photosynthetic O2 evolution was measured at 25°C with a Clark-type O2 oxygen electrode (Rank Brothers, Cambridge, U.K.). Cells from liquid cultures were collected by centrifugation and suspended in N2-saturated Mops-Tris (25 mM, pH 7.3) to a final chlorophyll concentration of 20 µg/ml. Internal and external Ci first were depleted under illumination (500 µmol photons·m−2·s−1) as judged by cessation of O2 evolution before measurements were initiated by addition of various concentrations of NaHCO3.

Ci uptake by C. reinhardtii cells was measured by the silicone oil filtration technique (24, 25) by using one of the walled ad1 progeny from the cross with CC620, because Ci uptake experiments were found to be unreliable with wall-less strains.

DNA and RNA Blot Analysis. Genomic DNA was isolated and purified in the presence of CTAB as described by Ausubel et al. (26). Total RNA was purified by the acid guanidinium thiocyanate-phenol-chloroform method described by Chomczynski and Sacchi (27).

Southern and Northern analyses were performed by standard
procedures (28), and membranes were scanned by using a phosphorimager (Storm).

Isolation of Sequences Flanking the Ble\textsuperscript{R} Insert from ad1 by Inverse PCR. Based on information from Southern blot analysis, BamHI was used to digest the genomic DNA isolated from ad1 to produce a fragment with a size \( \sim 1.5\)kb, including part of the inserted pSP124s vector and its flanking genomic DNA. The BamHI-digested ad1 genomic DNA (0.2 \( \mu\)g) was circularized with 1 unit of T4 DNA ligase (Invitrogen), precipitated, and the circularized product was used as template for inverse PCR by using standard PCR procedures. Three pairs of primers were designed, with each pair complementing the pSP124s sequence in opposite orientations. All three primer pairs produced PCR products with the correct predicted sizes, and amplified DNA from one primer pair (5\'-CTGGAGGCGGCTGTAGTGAACA-3\' and 5\'-GGAGGTCGTGTCACAGACT-3\') was sequenced to determine the sequence flanking the insert.

Identification of BAC Clones Containing the Wild-Type Ad1 Gene and Complementation of ad1 and pmp1. DNA flanking the site of insertion in ad1 was PCR-amplified based on the sequence of the DNA from inverse PCR and the C. reinhardtii genome (http://genome.jgi-psf.org/Chlre3/Chlre3.home.html). Using this amplified DNA as a probe, six BAC clones containing wild-type DNA from the inserted region were identified from a BAC library (Clemson University; www.genome.clemson.edu/groups/bac).

All complementation was performed by the glass bead transformation procedure (22). After transformation, cells were kept in low (air-level) \( \text{CO}_2 \) to observe wild-type growth of complemented mutants. Cells transformed with the empty vector or mock DNA were used as controls. For BAC complementation of ad1, DNA isolated from BAC clones 26B2 and 14L14 were used to transform ad1.

In complementing ad1 and pmp1 with LciB genomic DNA, a 3.6-kb fragment of genomic DNA containing the LciB coding region and putative promoter region was PCR-amplified from a BAC clone (26B2) by using a pair of primers: upper primer, 5\'-GAGTAGGCGTCGTCGTAA-3\', lower primer, 5\'-CGACACTGACGGCGCAATT-3\', and primer 5\'-CGACACTGACGGCGCAATT-3\'. They were used to transform ad1 and pmp1.

In complementing ad1 and pmp1 with LciB cDNA, LciB cDNA was PCR-amplified from a cDNA library (an expression cDNA library described below) with specific primers that introduced an NdeI site overhanging the start codon ATG at the 5\' end, and an EcoRI site after the stop codon at the 3\' end: upper primer, 5\'-AGCGAGCATATGGTGCTCTGCTTCTT-3\'; lower primer, 5\'-TTGAATTCTAGACCCGCAAGAG-3\'. The amplified cDNA was digested by EcoRI and NdeI and ligated into EcoRI/NdeI-digested pGenD plasmid (29), which placed the LciB cDNA between the PsdD promoter and terminator. This plasmid was linearized and used to transform ad1 and pmp1.

Construction and Screening of the cDNA Expression Library. Pooled mRNA (a gift from John Davies, Exelixis Plant Sciences, Portland, OR) isolated from cells grown to mid-log phase in tris-acetate-phosphate (TAP) (acetate-containing) medium in the light, TAP medium in the dark, high salt (HS) (minimal) medium in ambient levels of \( \text{CO}_2 \), and HS medium bubbled with 5\% \( \text{CO}_2 \) and identical to that used for constructing the core libraries described by Shragar et al. (30), was used to construct the C. reinhardtii cDNA expression library by using the HybriZAP 2.1 two-hybrid system (Stratagene, La Jolla, CA) according to the manufacturer’s protocol. To reduce secondary structure in the mRNA template, the reverse transcription reaction was performed by using SuperScript II reverse transcriptase (Invitrogen, Carlsbad, CA) at 50°C as suggested by Shragar et al. (30).

For identification of the LciD cDNA, a pair of primers was designed based on the sequence flanking the 3\' end of the predicted LciD coding region (http://genome.jgi-psf.org/Chlre3/Chlre3.home.html); upper primer, 5\'-AAGAAAGGCCTCGCCTTAACG-3\', and lower primer, 5\'-GGTACTGGGTGAAGCTAAAT-3\', and was used to amplify the putative 3\' UTR of LciD from the HybriZAP2.1 library by PCR. The amplified PCR product was used as a probe to screen the HybriZAP2.1 library. Five cDNA clones were identified and sequenced.