

Spontaneous Dominant Mutations in *Chlamydomonas* Highlight Ongoing Evolution by Gene Diversification

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We characterized two spontaneous and dominant nuclear mutations in the unicellular alga *Chlamydomonas reinhardtii*, *ncc1* and *ncc2* (for nuclear control of chloroplast gene expression), which affect two octatricopeptide repeat (OPR) proteins encoded in a cluster of paralogous genes on chromosome 15. Both mutations cause a single amino acid substitution in one OPR repeat. As a result, the mutated NCC1 and NCC2 proteins now recognize new targets that we identified in the coding sequences of the chloroplast *atpA* and *petA* genes, respectively. Interaction of the mutated proteins with these targets leads to transcript degradation; however, in contrast to the *ncc1* mutation, the *ncc2* mutation requires on-going translation to promote the decay of the *petA* mRNA. Thus, these mutants reveal a mechanism by which nuclear factors act on chloroplast mRNAs in *Chlamydomonas*. They illustrate how diversifying selection can allow cells to adapt the nuclear control of organelle gene expression to environmental changes. We discuss these data in the wider context of the evolution of regulation by helical repeat proteins.

INTRODUCTION

During and after transcription, sequence-specific RNA binding proteins control the processing, transport, localization, translation, and stability of coding and noncoding RNAs. Modular proteins, made up of tandem repeats of simple structural motifs (20 to 50 amino acids in length), most often comprising antiparallel α -helices and thus also termed helical repeat proteins, are particularly well suited to develop interactions with macromolecules, including RNA. Repeated motifs fold independently and stack on each other to form elongated or concave surfaces. While tetratricopeptide repeat (34 amino acids), Huntington, Elongation factor 3, protein phosphatase 2A, and yeast kinase TOR1 (HEAT; 39 amino acids), Armadillo (38 amino acids), Ankirin (33 amino acids), and leucine-rich repeat (23 to 24 amino acids) repeats are involved in protein-protein interactions, Pumilio and fem-3 binding factor (PUF; 36 amino acids), Transcription Activator Like Effector (TALE; 34 amino acids), pentatricopeptide repeat (PPR; 35 amino acids), Half A Tetratricopeptide (HAT; 34 amino acids), and mitochondrial termination factor (mTERF; ~30 amino acids) motifs mediate protein-nucleic acid interactions (reviewed in Rubinson and Eichman, 2012). Crystallographic structures of PUF (Wang et al., 2001, 2002; Miller et al., 2008), TALE (Deng et al., 2012; Mak et al., 2012), and PPR (Ke et al., 2013; Yin et al., 2013; Gully et al., 2015) proteins in complex with

their RNA/DNA targets confirmed that nucleic acids bind in an extended conformation to the inner concave surface of the solenoid, with each nucleotide contacting one, or at most two, consecutive repeats. Thus, repeats act in a modular fashion, with each repeat interacting with one nucleotide. Within a repeat, the side chain of a few residues at specific positions determines the recognized nucleotide, mainly by establishing hydrogen bonds with the nucleotide base. Nucleotide recognition by specific amino acid combinations was recently successfully predicted for PUF (Wang et al., 2002; Cheong and Hall, 2006; Filipovska et al., 2011), TALE (Boch et al., 2009; Moscou and Bogdanove, 2009), and PPR (Barkan et al., 2012; Takenaka et al., 2013; Yagi et al., 2013) proteins. Based on this “recognition code,” recombinant PUF and TALE proteins can be engineered to bind virtually any RNA or DNA target of interest (Christian et al., 2010; Cooke et al., 2011; reviewed in Bogdanove and Voytas, 2011; Filipovska and Rackham, 2012; Yagi et al., 2014). Furthermore, this modular architecture endows helical repeat proteins with a great versatility in vivo, as module reorganization through genetic recombination or substitutions of nucleotide-specifying amino acids will allow recognition of new targets.

Nuclear control of organelle gene expression is a key feature of eukaryotic cells that emerged after endosymbiosis (Choquet and Wollman, 2002; Woodson and Chory, 2008; Barkan, 2011). Indeed, every posttranscriptional step of organelle gene expression, RNA editing, splicing, processing, trimming, or translation activation is controlled in a gene- and, thus, sequence-specific manner by nucleus-encoded RNA binding proteins (denoted ROGEs for regulators of organelle gene expression; reviewed in Barkan and Goldschmidt-Clermont, 2000; Choquet and Wollman, 2002; Schmitz-Linneweber and Small, 2008; Woodson and Chory, 2008; Germain et al., 2013). Predictably, most ROGEs belong to helical repeat protein families, such as PPR, HAT, and mTERF (reviewed in Barkan and Small, 2014; Hammani et al., 2014). The great expansion of ROGEs in photosynthetic organisms contrasts with the

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poor conservation of ROGEs between different lineages. Despite their common structural organization, the various families of modular proteins do not generally share a common origin as their respective consensus sequences are not related. This suggests a high flexibility of nucleo-organelle interactions, well suited for rapid adaptation to new environmental constraints or ecological niches. For instance, PPR proteins, predominantly targeted to mitochondria or chloroplasts (Lurin et al., 2004) are particularly numerous in land plants with more than 450 members identified in *Arabidopsis thaliana* or rice (*Oryza sativa*). By contrast, the unicellular green alga *Chlamydomonas reinhardtii* possesses only 14 PPR proteins (Tourasse et al., 2013) but more than 120 members of another family of helical repeat proteins, poorly represented in land plants, the OPR (octatricopeptide repeat) proteins, defined by a degenerate motif of 38 residues (Auchincloss et al., 2002; Merendino et al., 2006; Loiselay, 2007; Eberhard et al., 2011; Rahire et al., 2012; Lefebvre-Legendre et al., 2015). As PPR repeats, OPR repeats are predicted to fold into a pair of antiparallel α -helices. Most OPR proteins are predicted to be targeted to organelles (Loiselay, 2007) where several have been shown to control the posttranscriptional steps of gene expression.

Most mutants affected in ROGEs described to date were screened for photosynthetic or respiratory defects after mutagenesis. They display a recessive phenotype, being defective in a gene product that, in the wild type, binds specifically to a given target transcript, usually in its 5' untranslated region (5' UTR). In contrast, the two nuclear mutations that we describe below in *Chlamydomonas* appeared spontaneously and are dominant. They correspond to single amino acid substitutions in two OPR proteins that gain a new function by recognizing a new target in the coding region of two chloroplast transcripts, thus providing insights into the evolution of the nuclear control of organelle gene expression.

RESULTS

Isolation of the *ncc2* Mutation, Which Alters the Stability of the *petA* Transcript

To better understand the regulation of chloroplast gene expression and identify new nuclear mutants, we used a chimeric construct to express a photosystem II (PSII) core protein under the control of the 5' UTR of an unrelated chloroplast gene. This chimera was expressed to reduced levels, insufficient for phototrophic growth, allowing us to isolate strains that had recovered phototrophic growth. The 5' *petA-psbB* chimera is made of the *psbB* coding sequence (CDS), encoding the PSII core antenna PsbB, translated under the control of the 5' UTR of the chloroplast *petA* gene, encoding the cytochrome *f* subunit of the cytochrome *b₆f* complex (Figure 1A). When inserted by chloroplast transformation at the *psbB* locus (see Table 1 for transformants generated in this work), this chimera only allows a low level of PsbB expression, insufficient to sustain the phototrophic growth of {5' *petA-psbB*} transformed cells, which display a PSII defective phenotype (Figures 1B to 1D). While plating the T9 {5' *petA-psbB*} transformant on minimal medium, we found one spontaneous phenotypic revertant, *Su0*, which had recovered phototrophic growth capability and increased accumulation of PsbB (Figures

1D and 1E). In crosses to a wild-type strain (see Table 2 for crosses performed in this work), all tetrad progeny inherited the 5' *petA-psbB* chimera, uniparentally transmitted by the *mt⁺* {5' *petA-psbB*} parent, but the photoautotrophic capability segregated 2:2. This was indicative of a single nuclear suppressor mutation that we called *ncc2* for nuclear control of chloroplast gene expression. We then crossed one phototrophic progeny (*mt⁻*) to the wild type (*mt⁺*) to eliminate the 5' *petA-psbB* chimera; two tetrad progeny had a wild-type phenotype, while the other two, presumably carrying the *ncc2* mutation, although phototrophic, presented fluorescence induction kinetics typical of cytochrome *b₆f* leaky mutants (Figure 2A). Indeed, their Φ_{PSII} (0.2), much lower than that of the wild type (0.59), indicates a decreased electron flow downstream of PSII (Maxwell and Johnson, 2000). This correlates well with the 6-fold reduced abundance in cytochrome *f*, when compared with the wild type (Figure 2B). In 5-min pulse labeling experiments, cytochrome *f* synthesis was much reduced and hardly detectable in the *ncc2* mutant (Figure 2C). When probing chloroplast transcripts for the major photosynthetic protein subunits by mRNA hybridization, we observed a selective drop in accumulation of the mature *petA* mRNA (below 5% of the wild-type level; Figure 2D), whereas *petA* processing intermediates were little affected (Figure 2D, asterisks, top panel). This suggests a preserved transcription but an accelerated degradation of the mature *petA* transcript in the mutant.

This phenotype readily explains the suppressor effect of the *ncc2* mutation on the expression of the 5' *petA-psbB* chimera. Cytochrome *f* is a CES (for controlled by epistasy of synthesis) protein, whose rate of synthesis is controlled by its assembly within cytochrome *b₆f* (Choquet et al., 1998, 2003; reviewed in Choquet and Wollman, 2008). In the wild type, a small fraction of cytochrome *f* remains unassembled and downregulates the translation of the *petA* mRNA or of any chimera driven by the *petA* 5' UTR, such as the 5' *petA-psbB* chimera here, by 3-fold. The *ncc2* mutant synthesizes reduced amounts of cytochrome *f* (Figure 2C), which precludes significant accumulation of unassembled cytochrome *f*, thereby releasing the translational downregulation of 5' *petA*-driven genes. The 5' *petA-psbB* chimera, now expressed 3-fold more in the *ncc2* mutant than in the wild type, allows PsbB synthesis at rates sufficient to sustain photoautotrophic growth, even though PsbB levels remain lower than those produced by the endogenous *psbB*.

We previously identified two nuclear genes, *MCA1* and *TCA1* (for maturation/stability and translation of the cytochrome *b₆f* complex subunit PetA, respectively), which contribute to the protection of the *petA* transcript against exonucleolytic degradation (Wostrikoff et al., 2001; Loiselay et al., 2008). However, crosses of *ncc2* to either *mca1-2* or *tca1-2* mutants (Supplemental Table 1) clearly showed genetic independence, indicating that the *ncc2* phenotype was not due to altered expression of *MCA1* or *TCA1*.

ncc2 Is a Dominant Mutation

To determine whether the *ncc2* mutation was recessive or dominant, we constructed vegetative diploid strains either homozygous or heterozygous for the *ncc2* locus, as described in Table 2. In contrast to homozygous *NCC2/NCC2* diploids, the heterozygous *ncc2/NCC2* and homozygous *ncc2/ncc2* diploids displayed the same 20-fold reduced accumulation of the *petA* mRNA as the

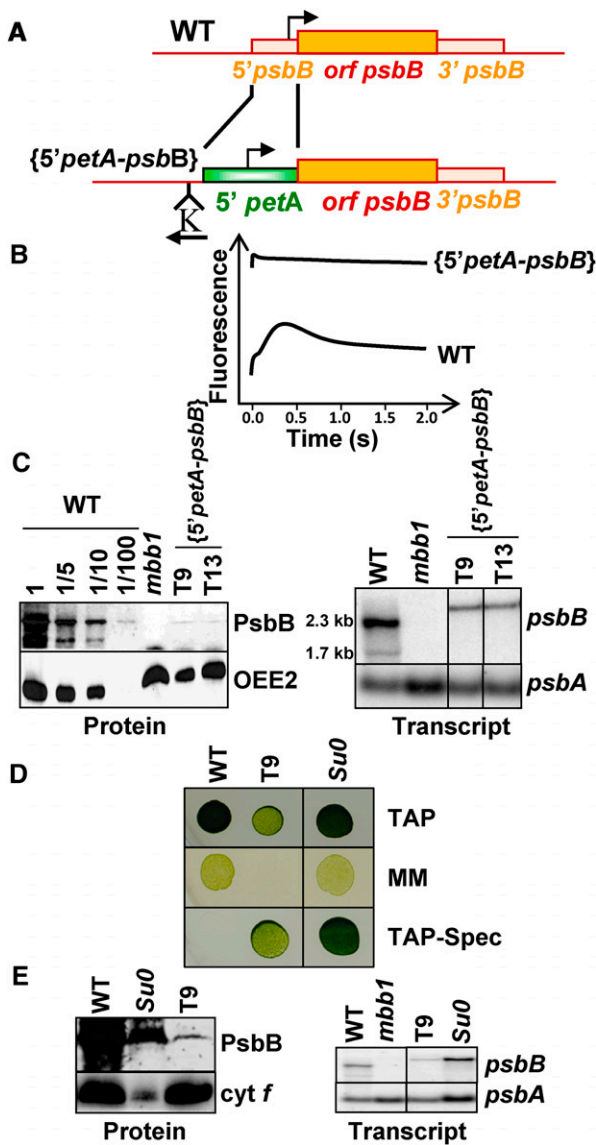


Figure 1. Isolation of the *ncc2* Mutant.

(A) Schematic description of the *psbB* gene in wild-type and {5'*petA-psbB*} transformants. Bent arrows indicate promoters, and the position and orientation of the selection cassette (K) are indicated.

(B) Fluorescence induction kinetics of dark-adapted wild-type and {5'*petA-psbB*} cells. The nearly constant fluorescence intensity over time in strain {5'*petA-psbB*}, as well as its high initial level, almost similar to the stationary level, is typical of leaky PSII mutants.

(C) Phenotypic characterization of {5'*petA-psbB*} transformed strains. Left: PsbB accumulation probed with a specific antibody in two independent transformants, in a dilution series of the wild type and in the *mbb1*-222E strain, defective for the accumulation of the *psbB* mRNA (Monod et al., 1992), as a specificity control. OEE2, whose accumulation is independent of PSII assembly (de Vitry et al., 1989), provides a loading control. Right: *psbB* mRNA accumulation in the same strains. Because of the larger size of the *petA* versus *psbB* 5'UTRs, the chimeric mRNA migrates more slowly than the endogenous *psbB* transcript.

(D) Growth properties of the wild-type, {5'*petA-psbB*}, and *Su0* strains. Drops of liquid culture (2×10^6 cells \cdot mL $^{-1}$) were spotted on TAP medium

haploid *ncc2* parental strain (Figure 2E). However, some gene dosage effect partially damped the effect of the *ncc2* mutation at the protein level, since cytochrome *f* abundance only decreased 2-fold in *ncc2/NCC2* heterozygotes (Figure 2E). In contrast to almost all mutations in ROGE genes described to date in *Chlamydomonas*, the *ncc2* mutation is not recessive but dominant. It thus strikingly resembles the other dominant nuclear mutation acting on chloroplast transcript isolated in *Chlamydomonas*, *ncc1*, which specifically destabilizes the monocistronic transcript of the chloroplast *atpA* gene encoding the α subunit of ATP synthase (Drapier et al., 2002). Similar to the *ncc1* mutation, the *ncc2* mutation is unlikely to be a loss-of-function allele, but rather likely modifies some gene product, so that it now acts on the *petA* mRNA.

The *ncc2* Mutation Causes NNC2 to Act on the *petA* CDS, Similar to the Action of Mutated NCC1 on *atpA*

In *Chlamydomonas*, the known ROGEs that control the stability of a chloroplast transcript target its 5'UTR, with the exception of the protein affected by the dominant *ncc1* mutation, which targets the *atpA* CDS (Drapier et al., 2002). We thus tested which part of the *petA* mRNA was targeted for degradation in the *ncc2* mutant, using two distinct *petA* chimeras.

At the *petA* locus, we first substituted the sequence coding for mature cytochrome *f* by that encoding the ATP synthase subunit α , truncated to maintain a similar mRNA length, fused in frame with the sequence coding for the lumen targeting peptide of cytochrome *f* ($\Delta f::\alpha_{Tr}$ chimera; Figure 3A). All chimeras used in this study were associated with an *aadA* selection cassette (Table 1), and transformants were selected for resistance to spectinomycin. One of the resulting transformants, { $\Delta f::\alpha_{Tr}$ }, was crossed to the *ncc2* mutant. RNA gel blot analysis showed no hybridization with an intragenic *petA* probe in the parental strain { $\Delta f::\alpha_{Tr}$ } because this sequence had been replaced by that of *atpA*. An *atpA* probe detected, in addition to the endogenous *atpA* transcripts, the $\Delta f::\alpha_{Tr}$ chimeric transcript in the parental transformant and in all progeny (Figure 3B). This was also true for a larger and minor transcript (indicated by an asterisk), resulting from cotranscription of the chimera with the downstream *aadA* cassette. Strikingly, these two bands accumulated to the same level in all progeny, irrespective of their *ncc2* or *NCC2* genotypes. Thus, chimeric transcripts lacking the sequence encoding mature cytochrome *f* are no longer destabilized by the *ncc2* mutation.

Symmetrically, the *petA* coding region, fused to the *atpA* 5'UTR and *rbcl* 3'UTR in the *dAfr* chimera (Figure 3A), was introduced by chloroplast transformation in a wild-type strain. After crossing with *ncc2*, all tetrad progeny expressed the chimeric *petA* transcript, larger than the genuine *petA* transcript. However, two

and grown under dim light ($10 \mu\text{E} \cdot \text{m}^{-2} \cdot \text{s}^{-1}$, top), on minimal medium (MM) under high light ($150 \mu\text{E} \cdot \text{m}^{-2} \cdot \text{s}^{-1}$, intermediate), or on TAP supplemented with spectinomycin ($500 \mu\text{g} \cdot \text{mL}^{-1}$; lower panel). Pictures were taken after 10 d of growth.

(E) Phenotypic characterization of the *Su0* strain. Accumulation of PsbB and cytochrome *f* (left) and of the *psbB* mRNA (right) in the same strains. *psbA* was used as a loading control.

Table 1. Strains Generated by Transformation in This Study

	Recipient Strain	Transforming Plasmid	Transformed Strain
Chloroplast transformation ^a	Wild type	pK5' <i>petA-psbB</i>	{5' <i>petA-psbB</i> }
	Wild type	pKΔf::α _{Tr}	{Δf::α _{Tr} }
	Wild type	p5' <i>dAfrR</i>	{ <i>dAfrR</i> }
	Wild type	<i>pf</i> ₄₂ St	{ <i>f</i> ₄₂ St}
	Wild type	<i>pf</i> ₁₄₅ St	{ <i>f</i> ₁₄₅ St}
	<i>ncc1</i>	pKatpA _{St} ^b	<i>ncc1</i> { <i>atpA</i> _{St} }
	Wild type	pKatpA ^M	{ <i>atpA</i> ^M }
	<i>ncc1</i>	pKatpA ^M	<i>ncc1</i> { <i>atpA</i> ^M }
	Wild type	pKpetA ^M	{ <i>petA</i> ^M }
	<i>ncc2</i>	pKpetA ^M	<i>ncc2</i> { <i>petA</i> ^M }
	Wild type	pK5' <i>petD</i> ::T2	{5' <i>petD</i> ::T2}
	<i>ncc1</i>	pK5' <i>petD</i> ::T2	<i>ncc1</i> {5' <i>petD</i> ::T2}
	<i>ncc2</i>	pK5' <i>petD</i> ::T2	<i>ncc2</i> {5' <i>petD</i> ::T2}
	Wild type	pKpetD _{Cod} ::T2	{ <i>petD</i> _{CDS} ::T2}
	<i>ncc2</i>	pKpetD _{Cod} ::T2	<i>ncc2</i> { <i>petD</i> _{CDS} ::T2}
Nuclear transformation ^c	Wild type	pNCC1 ^M -HA	NCC1 ^M -HA
	Wild type	pNCC2 ^M -HA	NCC2 ^M -HA

^aAll recipient strains were mt⁺. Thanks to the uniparental inheritance of the mt⁺ chloroplast genome, this allowed the transmission of the chimeras introduced by transformation to the whole progeny of crosses. Recipient strains were also spectinomycin sensitive, with all chimera being associated with a spectinomycin resistance cassette for selection of transformants. Transformants were selected for resistance to spectinomycin (100 mg · mL⁻¹) under low light (5 μE · m⁻² · s⁻¹) and subcloned in darkness until they reached homoplasmy.

^bReference: Eberhard et al. (2011).

^cTransformed strains were selected for resistance to paromomycin (10 mg · mL⁻¹) under low light (5 μE · m⁻² · s⁻¹).

members of each tetrad, likely carrying the *ncc2* allele, showed a markedly reduced accumulation of the chimeric transcript, similar to the endogenous *petA* mRNA in the *ncc2* parent (Figure 3C). The *petA* CDS is thus not only required but also sufficient for transcript degradation in the *ncc2* background.

Active Translation of the Target Transcript Is Required for Its Destabilization in *ncc2*, but Not *ncc1*, Backgrounds

The *ncc1* and *ncc2* mutations target the CDSs of *atpA* and *petA*, respectively; we wondered whether translation of these transcripts was required for their degradation in the *ncc1* and *ncc2* mutants. To test this, we used {*atpA*_{St}} and {*petA*_{St}} strains, which express untranslatable *atpA* and *petA* mRNAs that have their initiation codons replaced by a stop codon (Boulouis et al., 2011; Eberhard et al., 2011). We crossed these strains with the *ncc1* and *ncc2* mutants. In the {*atpA*_{St}} cells, the *atpA* transcript pattern is similar to that of the wild-type *atpA* gene, which comprises four mRNAs transcribed from the tetracistronic *atpA* gene cluster (Drapier et al., 1998). The *ncc1* mutation markedly decreases the amount of monocistronic *atpA* transcript relative to polycistronic forms (Drapier et al., 2002). We observed a strong reduction in the abundance of monocistronic *atpA* transcripts in two progeny (1 and 2) from the {*atpA*_{St}} × *ncc1* cross (Figure 4A). Thus, the *ncc1* mutation destabilizes the *atpA* transcript, even when not translated. In contrast, destabilization of the *petA* transcript by the *ncc2* mutation likely depends on active translation, as the accumulation of the *petA*_{St} transcript remained high in all tetrad progeny (Figure 4B).

Alternatively, the *petA* AUG initiation codon could be part of the site recognized by the mutated NCC2. We thus studied transcript accumulation over time in cultures treated with

lincomycin, an inhibitor of chloroplast translation. The drug did not affect the accumulation of *atpA* transcripts in *ncc1* cells (Figure 4C). By contrast, the accumulation of *petA* transcripts in the *ncc2* mutant increased spectacularly from barely detectable before lincomycin addition, up to wild-type levels after a 4-h treatment (Figure 4D). Thus, active translation of *petA* transcripts is required for their *ncc2*-dependent destabilization, whereas the

Table 2. Strains Generated by Crosses in This Study

mt ⁺ Parent	mt ⁻ Parent	Progeny
<i>ncc2</i> {5' <i>petA-psbB</i> }	Wild type	<i>ncc2</i> mt ⁻ {5' <i>petA-psbB</i> }
(Su0)		
Wild type	<i>ncc2</i> {5' <i>petA-psbB</i> }	<i>ncc2</i>
	(Su0)	
<i>arg2</i> [1]	<i>ncc2</i>	<i>ncc2 arg2</i>
<i>arg7</i> [1]	<i>ncc2</i>	<i>ncc2 arg7</i>
<i>ncc2 arg7</i>	<i>ncc2 arg2</i>	Diploid <i>ncc2/ncc2</i>
<i>arg7</i>	<i>ncc2 arg2</i>	Diploid <i>ncc2/NCC2</i>
<i>arg7</i>	<i>arg2</i>	Diploid <i>NCC2/NCC2</i>
{Δf::α _{Tr} } [2]	<i>ncc2</i>	<i>ncc2</i> {Δf::α _{Tr} }
{ <i>dAfrR</i> } [2]	<i>ncc2</i>	<i>ncc2</i> { <i>dAfrR</i> }
{ <i>petA</i> _{St} } [3]	<i>ncc2</i>	<i>ncc2</i> { <i>petA</i> _{St} }
{ <i>f</i> ₄₂ St} [2]	<i>ncc2</i>	<i>ncc2</i> { <i>f</i> ₄₂ St}
{ <i>f</i> ₁₄₅ St} [2]	<i>ncc2</i>	<i>ncc2</i> { <i>f</i> ₁₄₅ St}
{FAFA} <i>ncc1</i> [4]	<i>ncc2</i>	<i>ncc1 ncc2</i> {FAFA}
{ <i>atpA</i> _{St} } [5]	<i>ncc1</i>	<i>ncc1</i> { <i>atpA</i> _{St} }

References: [1] (Ebersold, 1967; Harris, 1989); [2] this work; [3] (Boulouis et al., 2011); [4] (Drapier et al., 2002); [5] (Eberhard et al., 2011). By convention, chloroplast genotypes, when relevant, follow the nuclear genotype and are written between braces.

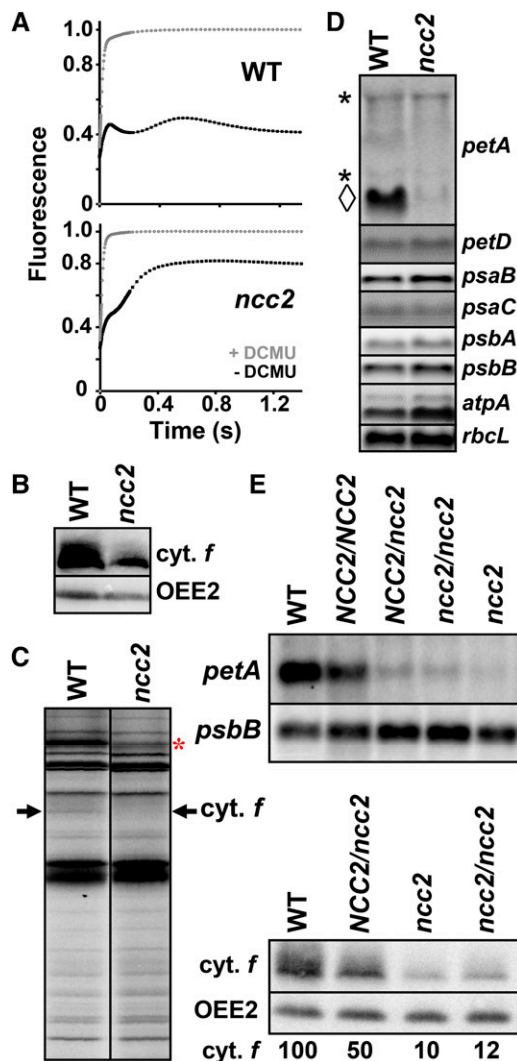


Figure 2. The Dominant *ncc2* Mutation Confers a b_6f Leaky Phenotype Due to Reduced Accumulation of the *petA* mRNA.

(A) Fluorescence induction kinetics of dark-adapted *ncc2* and wild-type cells. Black and gray curves show the kinetics recorded in the absence and in the presence, respectively, of DCMU (5 μ M), which blocks electron transfer downstream of PSII. Maximal fluorescence levels in the presence of DCMU were normalized to 1.

(B) Cytochrome *f* accumulation in wild-type and *ncc2* strains. OEE2 provides a loading control.

(C) Translation of chloroplast genes determined by 5 min 14 C-acetate (5 μ Ci·mL $^{-1}$) pulse-labeling experiments performed in the presence of cycloheximide (10 μ g·mL $^{-1}$) to block cytosolic translation. The arrow indicates the position of cytochrome *f*. The origin of the reduced rate of synthesis of Rubisco LS (red asterisk) in the *ncc2* mutant, despite the unchanged accumulation of the *rbcL* transcript (**D**), is unknown.

(D) Accumulation of representative transcripts for chloroplast photosynthesis genes in wild-type and *ncc2* strains, assessed by RNA gel blots using the probes indicated on the right of the panel. For the *petA* gene, the diamond indicates the mature transcript, while asterisks point to precursor RNA species.

(E) Accumulation of *petA* mRNA and cytochrome *f*, determined as above, in diploid strains either homozygous or heterozygous for the *ncc2* mutation.

ncc1 mutation destabilizes the *atpA* transcript independent of translation.

Toward a More Accurate Localization of the Targets of the *ncc2* and *ncc1* Mutations

To better understand how translation triggers *petA* mRNA degradation in strain *ncc2*, we introduced frame shifts within the *petA* CDS after nucleotides 93 or 390, causing premature abortion of translation after codons 42 and 145, respectively (Figure 5A). The f_{42} St mutation preserves the translation of the lumen targeting peptide, which is not sufficient to confer *ncc2* sensitivity to a translated sequence (Figure 3C). Tetrad analysis after crossing transformants $\{f_{42}$ St $\}$ and $\{f_{145}$ St $\}$ to a *ncc2* strain showed that the *ncc2* mutation still decreased the abundance of the f_{145} St transcript 20-fold (Figure 5B). In contrast, transcripts from the f_{42} St construct were much less sensitive to the *ncc2* mutation as their abundance only decreased 2-fold in *ncc2* progeny (Figure 5C). Thus, the *ncc2*-mediated degradation of *petA* transcripts starts early after the lumen targeting peptide has been synthesized and is completed before translation reaches the 145th codon of cytochrome *f*.

The target of the NCC1 mutated factor was previously localized in the last 1360 bases of the *atpA* mRNA (Drapier et al., 2002). We narrowed down this region by transforming the chloroplast of the *ncc1* strain with the $\Delta f::\alpha_{Tr}$ construct, which contains the *atpA* CDS deprived of its last 579 nucleotides (Figure 3A). In contrast to the endogenous *atpA* transcript, the chimeric $\Delta f::\alpha_{Tr}$ mRNA was not destabilized in the *ncc1* background (Figure 5D), suggesting that the *ncc1* mutation targets the last 579 nucleotides of the *atpA* transcript.

Identification of the *ncc2* and *ncc1* Mutations

Because the *ncc1* and *ncc2* mutations are dominant and allow photoautotrophic growth, we could not clone the genes by complementation. Thus, we mapped the mutations onto the *Chlamydomonas* nuclear genome by crossing the two mutants with the S1D2 strain, which shows a profusion of polymorphisms compared with laboratory strains (Gross et al., 1988; Kathir et al., 2003; Rymarquis et al., 2005). We harvested \sim 50 tetrads from each cross and picked one mutant per tetrad. *ncc2* progeny were identified based on their fluorescence phenotype. To identify *ncc1* progeny, we used for the cross the nonphotosynthetic strain *ncc1* {FAFA}, bearing the CDS of *atpA* fused to the 5' and 3' UTRs of the *petA* gene. This chimera is expressed at a low level in a wild-type background and, when combined with the *ncc1* mutation, prevents phototrophic growth (Drapier et al., 2002). By PCR-based mapping with diagnostic markers on all chromosome arms (Kathir et al., 2003), we found linkage for both mutations to the ZYS3 marker on the long arm of chromosome 15 (96 and 71% for the *ncc2* and *ncc1* mutations, respectively; Figure 6A). Using the new markers listed in Supplemental Table 2, the *ncc2* mutation was localized between nucleotides 659,176 and 1,063,367 (*Chlamydomonas*

Wild-type and *ncc2* strains are shown for comparison. *psbB* mRNA and OEE2 provide loading controls in RNA and immunoblots, respectively.

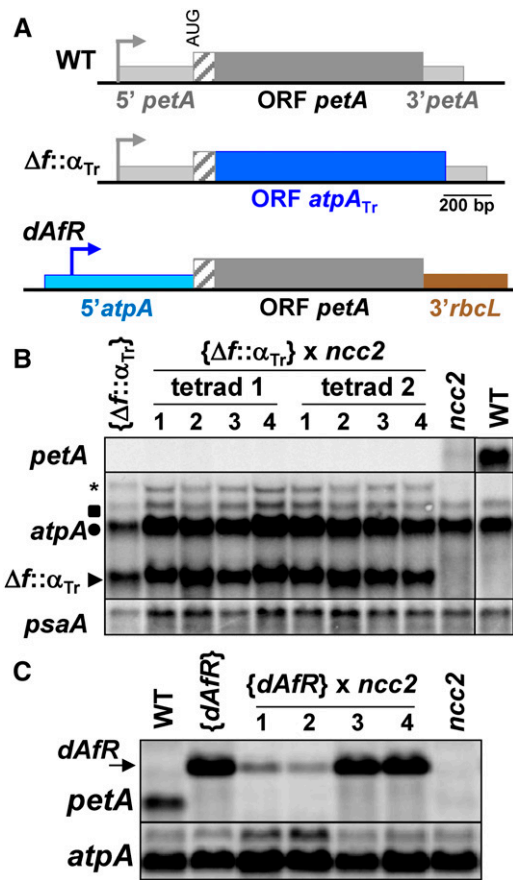


Figure 3. The *ncc2* Mutation Targets the *petA* CDS.

(A) Scheme of the chimeras. Sequence encoding the mature cytochrome *f* is shown as a dark-gray rectangle and that encoding the lumen targeting peptide as a hatched rectangle. *petA* 5' and 3'UTRs are represented by thinner light-gray rectangles. *petA* promoter is indicated by a bent arrow. In $\Delta f::\alpha_{Tr}$ chimera, the region encoding mature cytochrome *f* was replaced by the first 944 nucleotides of the *atpA* CDS, depicted as a blue rectangle. In *dAfR* chimera, the *petA* promoter and 5'UTR regions were replaced by the corresponding *atpA* regions (pale-blue rectangle), while *petA* 3'UTR was replaced by that of the *rbcL* gene (brown rectangle).

(B) Transcript accumulation in tetrad progeny from the cross $\{\Delta f::\alpha_{Tr}\} \times ncc2$ and in wild-type and parental strains. RNA gel blots were hybridized with probes derived from *petA*, *atpA*, and *psaA* (loading control) CDSs, as indicated on the left. Positions of the endogenous mono- and dicistronic *atpA* transcripts are indicated by a circle and a square, respectively. Arrow points to the position of the major $\Delta f::\alpha_{Tr}$ chimeric transcript, while the asterisk indicates a minor cotranscript that includes the downstream *aadA* resistance cassette.

(C) *petA* and *atpA* (loading control) transcript accumulation in tetrad progeny from the cross $\{dAfR\} \times ncc2$ and in wild-type and parental strains. Arrow points to the position the chimeric *dAfR* transcript.

genome v5.5, available on Phytozome at <http://phytozome.jgi.doe.gov/>; Merchant et al., 2007; Blaby et al., 2014).

We then crossed strains *ncc2* and *ncc1* {FAFA} and recovered a few double mutants among 80 tetrads, identified by their *ncc2*-like fluorescence induction kinetics and their poor phototrophic growth, due to the *ncc1* {FAFA} combination. They showed

decreased levels of both *atpA* and *petA* transcripts, which were comparable to those in the respective parental *ncc1* {FAFA} and *ncc2* strains. Accordingly, the accumulation of subunit α and cytochrome *f* was decreased, as shown in Supplemental Figure 1 for one typical double mutant. Its genome was sequenced using the Illumina platform (2×100 bp). After eliminating synonymous and intergenic polymorphisms and those also present in a collection of quasi-isogenic photosynthetic mutants carrying neither *ncc1* nor *ncc2* mutation, we were left with two point mutations in the interval determined by molecular mapping: an A \rightarrow C substitution at position 693,478 and a double substitution AT \rightarrow GG at position 1,001,095/6. These mutations cause a single amino acid substitution in two distinct genes encoding

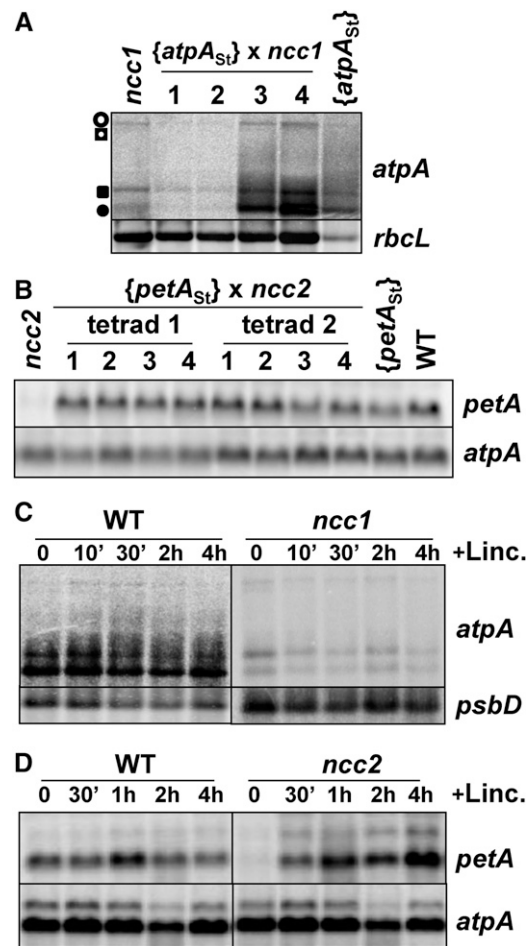


Figure 4. The *ncc2*, but Not *ncc1*, Phenotype Is Observed Only When the Target RNA Is Translated.

(A) *atpA* transcript accumulation in tetrad progeny from the cross $\{atpA_{St}\} \times ncc1$ and in parental strains. Position of the four transcripts from the *atpA* tetracistronic transcription unit is indicated. Loading control: *rbcL* mRNA.

(B) *petA* transcript accumulation in tetrad progeny from the cross $\{petA_{St}\} \times ncc2$ and in wild-type and parental strains (loading control: *atpA*).

(C) and (D) Accumulation of *atpA* (C) and *petA* (D) transcripts in *ncc1* (C), *ncc2* (D), and wild-type strains incubated in the presence of lincomycin for the indicated times. Loading controls: *psbD* (C) and *atpA* (D).

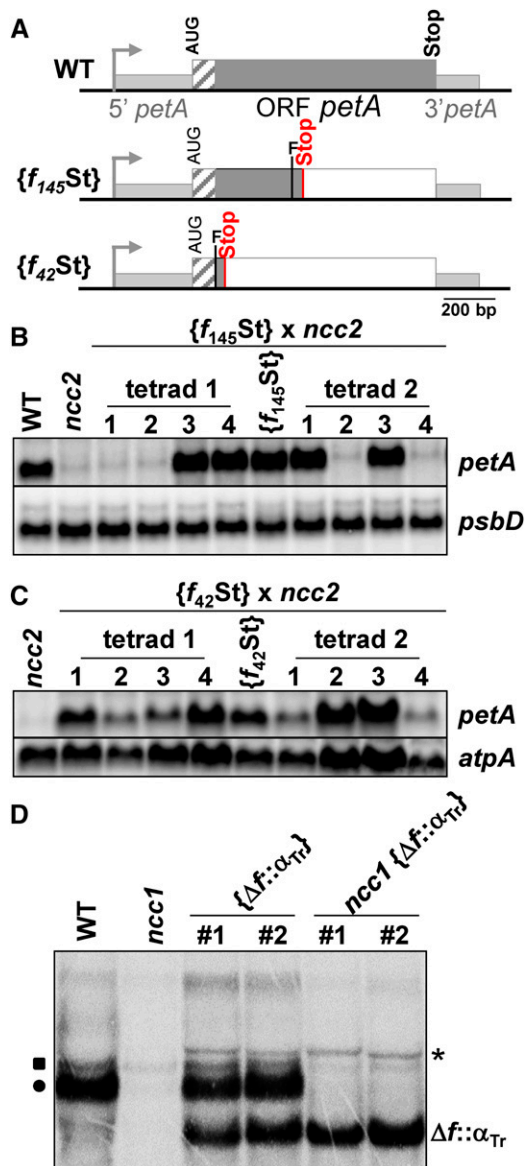


Figure 5. Narrowing Down the Target Sequences of the *ncc1* and *ncc2* Mutations.

(A) Schematic representation of the mutant *petA* genes, presented as in Figure 3A. The “F” indicates the position of the introduced frame shifts, while gray and white rectangles show translated and untranslated *petA* sequences, respectively.

(B) and **(C)** *petA* transcript accumulation in tetrad progeny from crosses *{f₁₄₅St}* × *ncc2* **(B)** and *{f₄₂St}* × *ncc2* **(C)**. Loading controls: *psbD* **(B)** and *atpA* **(C)**.

(D) Accumulation of *atpA*-hybridizing transcripts in wild-type and *ncc1* strains transformed with the Δf:α_{Tr} chimera (Figure 3A). The probe hybridizes with the chimeric transcript, either alone (Δf:α_{Tr}) or cotranscribed with the *aadA* cassette (asterisk) and with endogenous monocistronic (circle) and dicistronic (square) *atpA* mRNA.

OPR proteins: Cre15.g638950.t1 and Cre15.g640400.t1, whose gene models are fully supported by EST evidence displayed at <http://genomes.mcdb.ucla.edu/Cre454/>.

Direct sequencing of PCR products amplified from *ncc1* and *ncc2* strains (for primers, see Supplemental Table 3) localized the A → C substitution in the genome of the *ncc2* mutant but not in that of the *ncc1* mutant. It leads to a S₄₃₁ → R substitution in Cre15.g640400.t1, therefore named NCC2. Conversely, strain *ncc1*, but not strain *ncc2*, carried the AT → GG mutation, leading to the D₅₆₈ → A substitution in Cre15.g638950.t1, hereafter called NCC1. The NCC1 and NCC2 proteins both contain nine OPR repeats, the mutations changing the 6th residue of the 6th OPR repeat of NCC1 and the 8th residue of the 6th OPR repeat of NCC2 (Figures 6B and 6C). In addition, both proteins also contain a RAP domain (RNA Binding Abundant in Apicomplexa; Lee and Hong, 2004) at their C terminus.

Transgenic Expression of the Mutated NCC1 or NCC2 Protein Confers the *ncc1* or *ncc2* Phenotype to Transformed Strains

Since the *ncc1* and *ncc2* mutations are dominant, mutated copies of the NCC1/NCC2 genes, introduced in the wild type, should confer the *ncc1/ncc2* phenotypes to transformants. The genomic sequences coding for the mutated NCC1 and NCC2 proteins (hereafter referred to as NCC1^M and NCC2^M) were fused to a triple HA tag to allow their immunodetection and introduced into *Chlamydomonas* by transformation, using a vector carrying the *aphVIII* gene (Sizova et al., 1996). Clones, selected for paromomycin resistance, were screened with an HA-specific antibody for expression of NCC1^M or NCC2^M. As predicted from their respective molecular mass, NCC2^M migrates slightly faster than NCC1^M (Figure 7A). When assessing the accumulation of the *atpA/petA* transcripts by RNA gel blots, transformants expressing NCC1^M showed a marked decrease in the accumulation of the *atpA* monocistronic transcript. In independent transformants, the higher the accumulation of NCC1^M, the less *atpA* monocistronic transcripts (Figure 7B), supporting an NCC1^M-mediated degradation of the *atpA* mRNA. Similarly, the accumulation of the *petA* mRNA was strongly reduced in transformants expressing NCC2^M (Figure 7A), even though still higher than in the *ncc2* strain, probably because of insufficient expression level of the transgenic NCC2^M. In two transformants expressing NCC2^M, the extent of *petA* mRNA destabilization correlated with the abundance of the protein (Figure 7B).

Identification of the NCC1^M and NCC2^M Targets

Based on a preliminary version of the code for nucleotide recognition by OPR repeats (Y. Choquet, unpublished results), we predicted that NCC1^M and NCC2^M should target the NAGN-GATTA and GTGAGGNTA sequences, respectively, found at positions 1066 to 1075 and 130 to 139 of the *atpA* and *petA* CDS (Figures 8A and 9A). The NCC2^M target is located 5 bp downstream of the premature stop codon in the chimera *f₄₂St*, a region critical for the degradation of the *petA* transcript in the *ncc2* mutant (Figures 5C and 9A).

To confirm these target sequences, we substituted the nucleotide presumably recognized by the mutated OPR repeat in

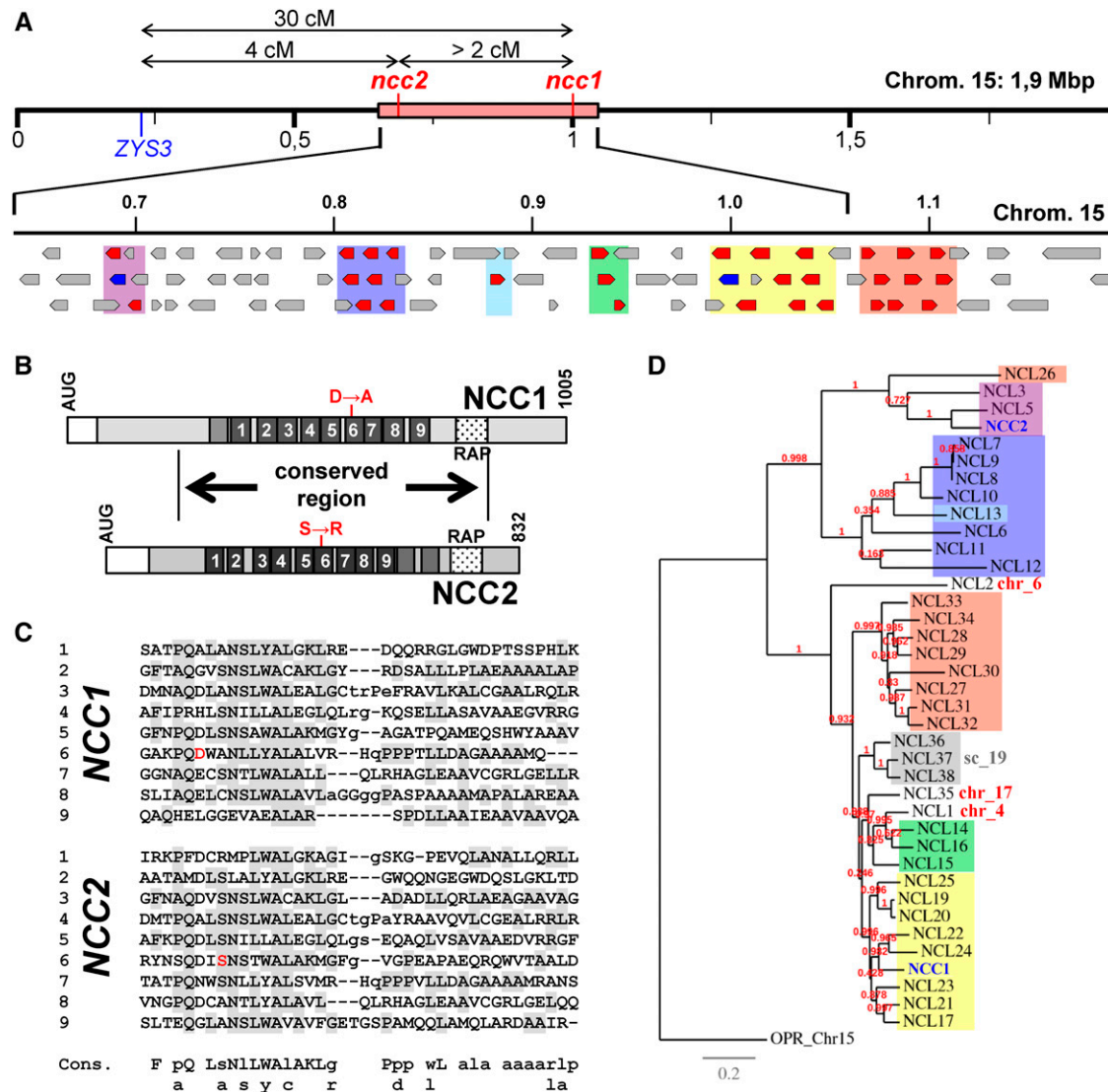


Figure 6. The *NCC1* and *NCC2* Genes.

(A) Top: Genetic and molecular map of the *ncc1* and *ncc2* loci. Locations on chromosome 15 of the *ZYS3* marker and *ncc1* and *ncc2* mutations are shown, along with genetic distances. The origin of the discrepancy between genetic distances determined in the three point test has not been investigated but likely originates from the poor fluorescence identification of some double mutants. The pink rectangle represents the molecular region containing the *ncc2* mutation, as determined by map-based cloning. Bottom: Physical map of the *NCL* gene cluster on chromosome 15. *NCL* genes are drawn in red, *NCC1* and *NCC2* in blue, and non-OPR genes in gray.

(B) Schematic representation of the *NCC1* and *NCC2* proteins. White rectangles depict the chloroplast transit peptide predicted by the ChloroP program. Dark-gray boxes represent the OPR repeats. Punctuated rectangles show the position of the RAP domains. The positions of the two substitutions in the *ncc1* and *ncc2* strains are shown in red. A highly conserved region (57% identity and 69% similarity) between the two proteins is indicated.

(C) Alignment of OPR repeats in the *NCC1* (top) and *NCC2* (bottom) proteins, with residues corresponding to the consensus (bottom) shaded in gray. Mutated amino acids in *ncc1* and *ncc2* strains are shown in red.

(D) Phylogeny of *NCL* proteins. Maximum likelihood tree of the *NCL* proteins using *Chlre_OPR68* as an outgroup. Branch length represents the estimated rate of amino acid substitution. Colored boxes indicate the genomic location of the corresponding *NCL* genes, as indicated in **(A)**. *NCC1* and *NCC2* are written in blue.

the predicted target of *NCC1^M*, leading to the conservative mutation I₃₉₁→V (Figure 8A). After transformation of wild-type and *ncc1* strains, the mutated *atpA* gene, *atpA^M*, replaced the endogenous *atpA* gene. In the wild-type background, this mutation slightly decreased (~20%) the accumulation of the monocistronic

atpA transcript and slightly increased that of the dicistronic *atpA* transcript (Figure 8B). In contrast, in the *ncc1* background, the mutation increased the level of *atpA* transcripts, with a 4-fold increase in monocistronic transcript and a 2-fold increase in dicistronic transcript. Thus, the ability of *NCC1^M* to destabilize the

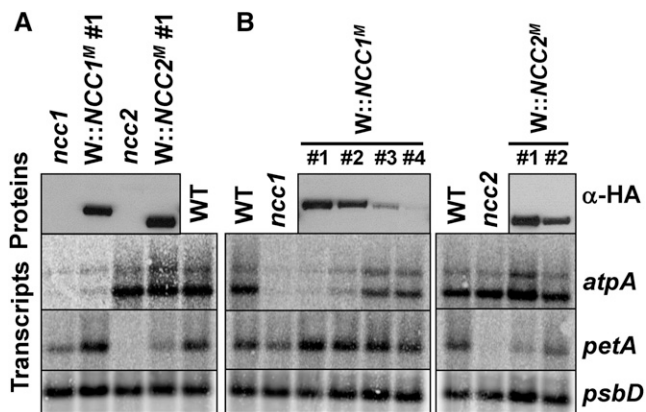


Figure 7. Expression of *NCC1^M* (*NCC2^M*) Confers the *ncc1* (*ncc2*) Phenotype to Transformed Strains.

(A) Accumulation of the *petA* and *atpA* mRNAs in wild-type, *ncc1*, and *ncc2* strains and in two transformed strains expressing *NCC1^M* and *NCC2^M*, as shown with an antibody against the HA tag (upper panel).

(B) Left: Decreasing accumulation of *atpA* mRNA in a series of transformants accumulating increasing amounts of *NCC1^M*. Right: Two transformants illustrating the negative correlation between accumulations of *petA* mRNA and *NCC2^M*.

atpA transcripts was reduced, but not abolished, by a point mutation in the predicted target.

The partial suppression of the *NCC1^M* effect, when only one nucleotide was substituted in its target, prompted us to introduce several silent mutations when assessing the target sequence of *NCC2^M*, converted from GTGAGGCTA to GAGAAGCAA (*petA^M*; Figure 9A). Following transformation, the mutation *petA^M* had no effect on the *petA* mRNA steady state level in the wild type. In stark contrast, it completely abolished the *ncc2* phenotype as it led to a spectacular restoration of the accumulation of *petA* mRNA in the *ncc2* strain, up to wild-type level (Figure 9B), demonstrating that we indeed identified the target of *NCC2^M*.

Wondering whether the target of *NCC2^M* would still be recognized when placed in a different nucleotide context, we inserted it either in the *petD* 5'UTR (*5'petD::T2*) or within the *petD* CDS (*petD_{CDS}::T2*) (Figure 9C). We used these two insertion sites to assess whether active translation would still be required for the destabilization of the *petD* transcript. When transformed with the *5'petD::T2* construct, both wild-type and *ncc2* strains remained phototrophic and the mutated *petD* transcript accumulated to the same level as does the original *petD* transcript in the wild type (Figure 9D). In contrast, chloroplast transformants expressing the *petD_{CDS}::T2* chimera were incapable of phototrophic growth, in wild-type or in *ncc2* backgrounds, as expected from the probable impairment of transmembrane helix II insertion within the membrane. In the *ncc2* background, the accumulation of the chimeric *petD* mRNA was largely prevented, whereas it remained unaffected in the wild type, when compared with that of the wild-type *petD* transcript (Figure 9D). Moreover, a 4-h lincomycin treatment of the *ncc2* {*petD_{CDS}::T2*} strain restored the accumulation of the chimeric *petD* mRNA to a wild-type level, confirming the dependence of the *ncc2* phenotype on the translation of its target (Figure 9E).

NCC1 and *NCC2* Belong to the *NCL* Subfamily of Paralogs Encoding OPR-RAP Proteins

In BLAST searches against the *Chlamydomonas* proteome, *NCC1* and *NCC2* hit each other with low E-values ($< 10^{-100}$), along with a set of 36 closely related proteins (Supplemental Table 4). The similarity between these proteins was much higher than with any other protein in the nonredundant Protein Database (<http://www.ncbi.nlm.nih.gov/protein/>), including those of the closely related alga *Volvox carteri* ($E > 10^{-30}$). Thus, *NCC1* and *NCC2* are part of a *Chlamydomonas*-specific group of 38 highly similar paralogs that will hereafter be called *NCL*, for *NCC-Like* (described in Supplemental Data Set 1). As shown in the alignment (Supplemental Figure 2A), *NCL* proteins comprise a highly conserved central region, containing 7 to 12 OPR repeats and an ~60-residue C-terminal RAP domain. By contrast, N- and C-terminal extensions, upstream and downstream of this conserved block, are more divergent.

Strikingly, 32 of these 38 *NCL* genes are clustered on the long arm of chromosome 15 between positions 686,690 and 1,113,927 (Figure 6A), while three are found on the as yet unassembled scaffold 19 and 3 isolated genes lie on chromosomes 4, 6, and 17.

At variance with the bulk of OPR-encoding genes, which have an average number of 12 introns regularly scattered along the CDS, most *NCL* CDSs contain a single intron, at a conserved position with respect to the protein sequence (Supplemental

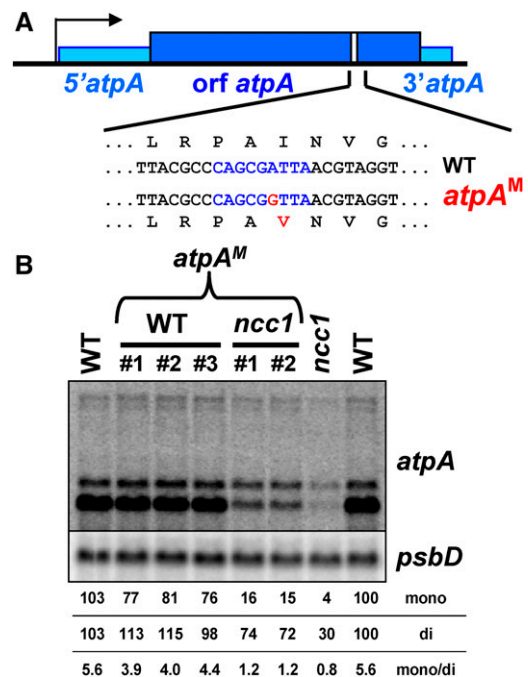


Figure 8. Identification of the Target of *NCC1^M*.

(A) Location of the target of *NCC1^M*, written in blue along the *atpA* gene. The mutation introduced in the *atpA^M* construct is shown in red.

(B) Accumulation of the *atpA* transcript in wild-type and *ncc1* strains transformed with the *atpA^M* construct. Independent transformants are presented for each background. Untransformed wild-type and *ncc1* strains are shown for comparison. Loading control: *psbD*.

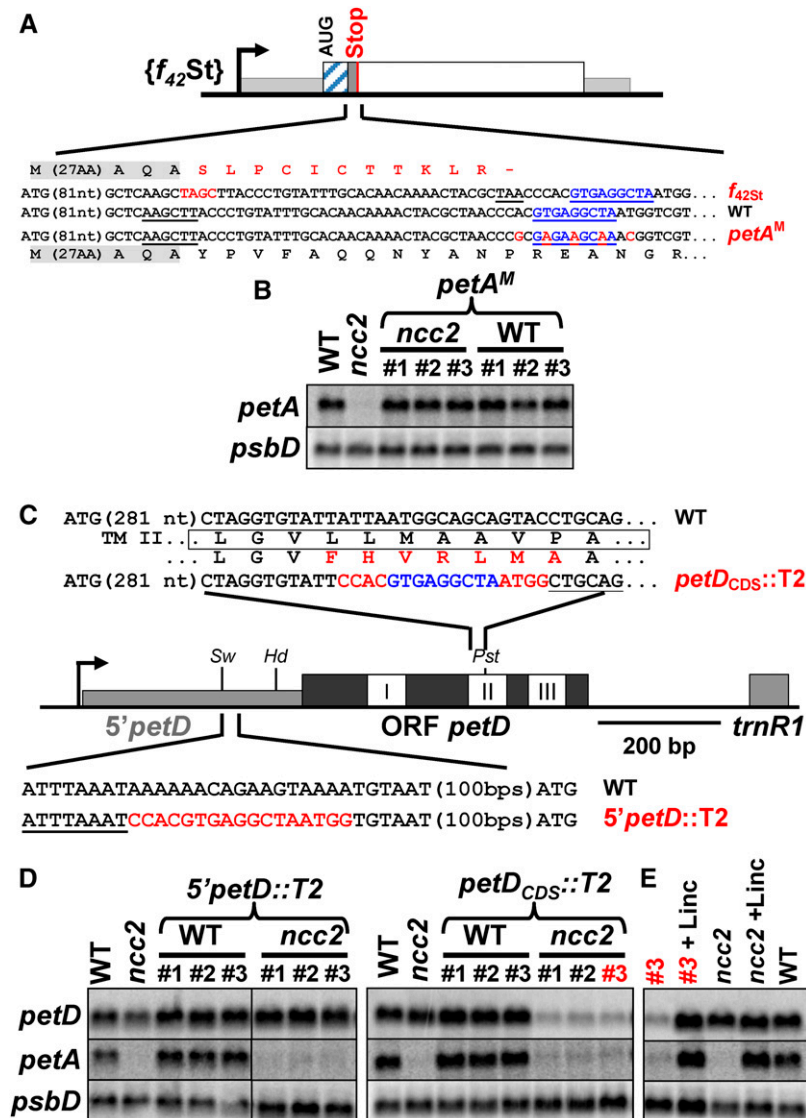


Figure 9. Identification of the Target of NCC2^M.

(A) Location of the target of NCC2^M, written in blue along the *petA* gene. Silent mutations introduced in the *petA*^M construct are written in red. Residual translation of *petA* in the *f*₄₂St mutant, downstream of the frame shift, is shown. The *Hind*III site used to introduce the frame shift indicated in red is underlined. (B) Accumulation of the *petA* transcript in wild-type and *ncc2* strains transformed with the *petA*^M construct. Three independent transformants (#1 to #3) are presented for each background. Untransformed wild-type and *ncc2* strains are shown for comparison. Loading control: *psbD*. (C) Insertion sites of the NCC2^M target within the *petD* gene. Schematic representation of the *petD* gene with the 5'UTR and CDS drawn as thin light-gray and thick dark-gray rectangles, respectively, while the three white boxes represent transmembrane helices. Relevant restriction sites (*Swa*I, *Hind*III, and *Pst*I) are indicated. Nucleotide regions surrounding the NCC2^M target are shown with restriction sites underlined. (D) Accumulation of *petD* transcript in wild-type and *ncc2* strains transformed with the 5'*petD*::T2 (left) and *petD*_{CDS::T2} (right) constructs. Three independent transformants (#1 to #3) are presented for each background. Untransformed wild-type and *ncc2* strains are shown for comparison. A *petA* probe reveals *ncc2* background. Loading control: *psbD*. (E) *petD* and *petA* mRNAs accumulation in strains *ncc2* and *ncc2* {*petD*_{CDS::T2}} #3, before and after a 4-h incubation with lincomycin. The wild type is shown for comparison, and *psbD* was the loading control.

Data Set 1 and Supplemental Figure 2). Most probably this intron was already present in the ancestor gene that gave rise to the NCL family by duplications. Phylogeny of NCL proteins was studied using maximum likelihood inference (Figure 6D) and was well correlated with the position and orientation of NCL genes

along the cluster (Figures 6A and 6D), suggesting that local tandem gene duplications played a major role in the expansion of the NCL family. This evolution is probably still going on: the adjacent NCL7, NCL8, and NCL9 genes probably originated from very recent duplication events, as NCL7 and NCL8 only

differ from *NCL9* by 6 and 15 bases, respectively, along the 2277-bp CDS. This leads to three amino acid substitutions in *NCL8* and to very limited changes in the very C-terminal end of *NCL7*. Conversely, the *NCL* family contains at least two inactivated genes, *NCL2* and *NCL21*, whose sequences, although similar to that of other *NCL* genes over 598 and 1171 codons, respectively, are interrupted by premature stop codons at positions 200 and 305. According to expression data available on Phytozome, *NCL2* is probably not transcribed. Some other *NCL* genes, such as *NCL12* and *NCL16*, encode truncated proteins comprising only the N-terminal extension and the first four OPR repeats, which explains their higher BLAST E-value (Supplemental Table 4).

***NCL* Genes Evolved under Diversifying Selection Pressure**

Within the conserved block, successive OPR repeats, although all obeying the OPR consensus, significantly differ one from another (Supplemental Figure 2B). By contrast, the sequence of a given repeat is remarkably conserved between the different *NCL* proteins (Supplemental Figure 2A), except at some variable positions (e.g., 3 and 6). This, together with the spontaneous appearance of the *ncc1* and *ncc2* mutations, suggests that the *NCL* proteins may be under diversifying selection, i.e., that nonsynonymous substitutions are selected at some sites, probably because they provide enhanced fitness. We tested this hypothesis by computing the nonsynonymous versus synonymous substitution ratio ($\omega = dN/dS$) for *NCL* proteins, using the program suite PAML, which reconstitutes the evolution of codons based on an alignment. dN/dS ratios were compared with predictions of evolution models allowing the presence of sites with $\omega > 1$ (diversifying selection) or not (nearly neutral selection) (Yang and Swanson, 2002; Yang, 2007). Likelihood ratio tests shown in Table 3 clearly show that models allowing classes with $\omega > 1$ much better fit the observed values, as judged by the P values of χ^2 tests. In the alignment, 26 sites were considered to be under diversifying selection at 99% confidence (49 at 95%), most of them within OPR repeats.

In conclusion, *NCL* genes are still under dynamic evolution and undergo a “birth, diversification, and death” process that is

driven by diversifying selection pressure and likely generates new RNA targets via mutations in OPR repeats.

DISCUSSION

***ncc1* and *ncc2*, Dominant and Atypical Mutations in Two OPR Proteins, Lead to RNA Degradation Rather Than RNA Protection**

All mutations affecting ROGEs characterized so far in *Chlamydomonas* are recessive, as they inactivate a protein that binds specifically to a given target transcript, usually in its 5'UTR. Thus, ROGEs likely coevolved with the 5'UTR of their target mRNA. By contrast, the *ncc1* and *ncc2* mutations described here represent a different category of ROGE mutations in *Chlamydomonas*. Both appeared spontaneously in our laboratory strains, are dominant, and act on the CDS of their target mRNA. Their unusual dominant nature results from their molecular basis: a single amino acid substitution in one OPR repeat of two different OPR proteins. By introducing a mutant copy of either *NCC1* or *NCC2* in the wild type, we show that these single substitutions are sufficient to destabilize the *atpA* or *petA* transcripts, respectively. They would change the recognized nucleotide and create new targets for the mutated proteins that fortuitously lie within CDSs. As observed for PPR proteins and in agreement with our preliminary version of the OPR code, well-conserved positions (e.g., 4, 5, and 7 to 15; Supplemental Figure 2) within the first antiparallel α -helix of the OPR repeats should mostly contribute to protein scaffolding, whereas more variable positions (e.g., 3 and 6) would be involved in nucleotide recognition. The *ncc1* mutation, that changes the variable 6th residue of the repeat, was thus expected to alter nucleotide recognition. Surprisingly, the *ncc2* mutation changing the quite conserved 8th residue nevertheless leads to the recognition of a new target.

ROGEs that bind the 5'UTR of their target transcript either activate its translation or protect it from exonucleolytic degradation. In contrast, the interaction of *NCC1^M* and *NCC2^M* with the *atpA* and *petA* CDSs leads to the degradation of the *atpA* and *petA* mRNAs by a mechanism that remains to be studied:

Table 3. Comparison of Codon Evolution Models in *NCL* Genes

	Likelihood ^a		$2\Delta l^b$	P Value ^c	p^d	ω^e
	Nearly Neutral	Positive Selection				
A	M1 −43839.38	M2 −43673.49	(M2 vs. M1) −331.78	1.33 E ^{−72}	0.093	3.05
B	M7 −43806.28	M8 −43600.42	(M7 vs. M8) −411.72	5.66 E ^{−90}	0.147	2.35

Part A is a comparison of model M1 versus M2. M1 allows two classes of codons ($\omega = 0$, negative selection; $\omega = 1$, neutral selection) and M2 an additional class under positive selection ($\omega > 1$). Part B is a comparison of model M7 versus M8. M7 allows a continuous β -distribution of $0 < \omega < 1$, with an additional class ($\omega > 1$) in M8.

^aLog likelihood values for nearly neutral (M1 and M7) or positive selection (M2 and M8) models.

^bLikelihood ratio between the two models.

^cEvaluated from χ^2 distribution ($df = 2$).

^dProportion of sites under positive selection.

^eMean ω value for sites under positive selection.

NCC1^M and NCC2^M may recruit an endonuclease or may themselves carry an endonucleolytic activity, which may be carried by the RAP domain found at the C terminus of both proteins. Indeed, structural modeling of the RAP domains of NCC1 and NCC2 by the I-TASSER software (Zhang, 2008) used four endonucleases as major templates (Supplemental Figures 3 and 4).

A Link with Translation

Strikingly, NCC2^M only degrades its target transcript upon translation. This was observed in two different sequence contexts, within the *petA* and *petD* transcripts, in which the target is involved in widely different secondary structures. It is therefore unlikely that NCC2^M binding to its target site requires the ribosome-mediated unfolding of RNA secondary structures. Rather, the degradation of the transcript would depend on a tight contact between the ribosome and NCC2^M, as supported by the limited decrease in *petA* mRNA observed upon early translation termination, a few nucleotides upstream of the NCC2^M target. This interaction would change the conformation of NCC2^M itself or of its interacting endonuclease, thereby activating nucleolytic activity, as shown for ribosome associated YoeB, RelE, and RegB RNases (Odaert et al., 2007; Neubauer et al., 2009; Feng et al., 2013).

By contrast, the degradation of the *atpA* transcript by NCC1^M does not require ongoing *atpA* translation. However, the monocistronic *atpA* transcript is by far more destabilized than polycistronic *atpA* transcripts. Possibly, the target of NCC1^M, localized 460 bp before the end of the *atpA* CDS, is fully accessible in the monocistronic *atpA* transcript, but trapped within secondary structures with downstream sequences in polycistronic transcripts.

NCC1 and NCC2 Belong to the NCL Family of Paralogous OPR-RAP Proteins

NCC1 and *NCC2* belong to the *NCL* family of highly similar OPR paralogs that differ in many respects from the bulk of genes encoding OPR proteins in *Chlamydomonas* (hereafter “*OPR* genes”). Unlike most *OPR* genes, for which orthologs can be easily found in *V. carteri*, probably because of a conserved role in organelle biogenesis, *NCL* genes appeared after the separation between the *V. carteri* and *Chlamydomonas* lineages. They share a single intron at a fixed position and show high sequence similarity, indicating recent appearance by gene duplication. Their genomic organization is also striking. Whereas *OPR* genes are randomly dispersed throughout the *Chlamydomonas* or *V. carteri* genomes, most *NCL* genes are clustered on chromosome 15. The rapid evolution of this family likely rests on tandem duplications, giving rise to seven subclusters of closely related genes (Figures 6A and 6D). This tandem organization would favor unequal crossovers, leading to gene duplication/loss and to repeat duplications. Interallelic recombination and gene conversion could also participate in repeat shuffling and, thus, to the diversification of binding sites. Indeed, *NCL* genes evolve under diversifying selection pressure, with specific positions showing a high dN/dS ratio. The spontaneous appearance of *ncc1* and *ncc2* mutations in laboratory conditions suggests that this diversification is still active. *NCL* genes also can decay, as evidenced by the presence of inactivated and truncated genes.

Thanks to this vigorous “birth-diversification-death” process, *NCL* genes represent a constant source of RNA binding proteins with new target specificities.

The *NCL* family strikingly resembles two other rapidly evolving gene families in plants. Restorer of Fertility-Like (RFL)-PPR proteins are distributed throughout higher plants and include Restorers of fertility (Rfs) proteins characterized in radish (*Raphanus sativus*), petunia (*Petunia hybrida*), or rice (Fujii et al., 2011; reviewed in Dahan and Mireau, 2013). Rfs repress the expression of mitochondrion-encoded chimeric open reading frames that are generated by recombination between different copies of the mitochondrial genome and cause cytoplasmic male sterility (CMS) in various crop species. RFL proteins are considered as a reservoir for the evolution of Rfs counteracting the expression of new CMS genes. As *NCL* genes, most *RFL* genes are clustered (in two regions of chromosome 1 in Arabidopsis or on chromosome 10 in rice), favoring unequal crossovers and local duplications, which likely contribute to the expansion of the RFL family (Hernandez Mora et al., 2010). *RFL* genes are under diversifying selection, especially those residues of the PPR motifs that are involved in nucleotide recognition (Geddy and Brown, 2007; Fujii et al., 2011), thus favoring the appearance of new RNA targets. They evolve rapidly, with large divergence between species or even between different accessions (Jonietz et al., 2010).

Similarly, pathogen resistance (*R*) genes, which activate plant defense reactions upon recognition of specific pathogen effectors, evolve rapidly in an “arms race” against the constantly evolving plant pathogens. *R* genes predominantly belong to the large and highly dynamic family of nucleotide binding site leucine-rich repeat (NLR) proteins (reviewed in DeYoung and Innes, 2006; Ye and Ting, 2008; Qi and Innes, 2013). They are often clustered on the genome, a situation that, again, favors the expansion and evolution of the gene family (Michelmore and Meyers, 1998), and are under diversifying selection that targets mainly the solvent-exposed residues involved in protein-protein interaction (Wulff et al., 2009; Seeholzer et al., 2010). It is of note that the new alleles created by these diversifying selection processes in *RFL* and *NLR* families confer a dominant phenotype of fertility or pathogen resistance to their host organism (Elkonin, 2005; Song et al., 2006; Moffett, 2009).

The Elusive Physiological Function of *NCL* Proteins

In the above examples, the evolutionary drives for the expansion of gene families are genome warfare and biotic stress. However, the selective pressure that led to the expansion of *NCL* genes in *Chlamydomonas* is not known. To address this question, some knowledge of the physiological function of the wild-type NCC1 and NCC2 proteins would be of interest.

We fail to find additional phenotypes conferred by the *ncc1* or *ncc2* mutations. Either the gain of function provided by the substitutions within NCC1^M and NCC2^M preserves the original function of the wild-type proteins or these functions are subtle and escaped our phenotypic analysis. In Arabidopsis, inactivation of the *RFL* genes *RPF1* or *RPF2*, while modifying the processing of a few mitochondrial transcripts (*nda4* or *cox3* and *nad9*, respectively), do not alter the accumulation of their gene products nor lead to any obvious phenotype (Jonietz et al., 2010; Hölzle

et al., 2011). Furthermore the *RPF2* gene seems inactivated in some accessions (Former et al., 2008; Jonietz et al., 2010). Similarly, would the gene duplication be too recent to have elicited the recruitment of NCC1 and NCC2 for some physiological process, these proteins could have no function, in contrast to their mutant allele. However, their transcription seems regulated: cursory examination of their expression profiles on the Phytozome browser suggests a transient repression of *NCC2* during nitrogen starvation (GSE34585) and a repression of *NCC1* by H₂O₂ treatment (GSE34826).

The spontaneous appearance of the *ncc1* and *ncc2* mutations likely illustrates a “trial and error” process operating on genes whose physiological function is not essential or redundant. It may ultimately lead to the recognition of new RNA targets, which could provide a selective advantage in specific environmental conditions. This is illustrated by the experimental conditions in which we recovered the *ncc2* strain. It appeared when plating a nonphototrophic chimeric strain in phototrophic conditions (Figure 1) but shows a leaky cytochrome *b₆f*-defective phenotype and would have been counterselected in cells with a wild-type chloroplast genome kept in phototrophic conditions.

METHODS

Strains and Growth Conditions

Wild-type, mutant, and transformed *Chlamydomonas reinhardtii* strains, all derived from 137c, were grown at 25°C in Tris-acetate-phosphate (TAP) medium, pH 7.2 (Harris, 1989), under continuous light (5 to 10 μE m⁻² s⁻¹). Strains *ncc1* and *ncc1* {FAFA} were previously described as *mda1-ncc1* and *mda1-ncc1*{FAFA} by Drapier et al. (2002). Crosses were performed according to Harris (1989). Vegetative diploids were selected on arginine-free plates from crosses between strains carrying the *arg2* and *arg7* mutations (Ebersold, 1967; Harris, 1989). After 12 d in low light, dark-green colonies comprised of large cells were checked by PCR for the presence of both *mt* loci (Werner and Mergenhagen, 1998).

Constructs and Nucleic Acid Manipulations

Standard nucleic acid manipulations were performed according to Sambrook et al. (1989). Primers used in this study are listed in Supplemental Table 3.

5' *petA-psbB* Chimera

The *petA* 5' UTR and promoter regions were excised with *EcoRV* and *NcoI* from plasmid p5F (Choquet et al., 1998) and cloned into the p38A.NcoI vector (Vaistij et al., 2000) digested with *StuI* and *NcoI* to yield plasmid p5' *petA-psbB*. The spectinomycin resistance cassette, excised from plasmid pUC-ATPX-AAD (Goldschmidt-Clermont, 1991) by *SmaI* and *EcoRV*, was inserted in the Klenow-treated *AvrII* site, in reverse orientation with respect to the *psbB* gene, yielding plasmid pK5' *petA-psbB*.

pΔf::α_{Tr}

A truncated version of the *atpA* CDS, fused the *petA* lumen targeting peptide, replaced the sequence encoding mature cytochrome *f*. Truncated *atpA* CDS was amplified in two steps from plasmid *patpA2* (Ketchner et al., 1995). A 254-bp fragment amplified with primers *atpA_{FUS}*-FW/*atpA_{FUS}*-RV1 was digested with *HindIII* and *PstI* and cloned into *HindIII*-*PstI* digested *pf*::H₈ vector (Choquet et al., 2003) to create plasmid p5f-int. A 965-bp fragment amplified with primers *atpAFW*/*atpA_{FUS}*-RV2

was digested with *EcoRI* and *PstI* and cloned into vector p5f-int, digested with the same enzymes to create plasmid pΔf::α_{Tr}. The *aadA* cassette was then inserted at the *HincII* site, upstream and in reverse orientation with respect to the *petA* CDS, to yield the plasmid pKΔf::α_{Tr}.

p5' *dAfr*

Plasmid p5' *dAfr*, comprising the *petA* CDS expressed under the control of the *atpA* 5' UTR and the *rbcL* 3' UTR, was created from plasmid pFADBE1 (Kuras and Wollman, 1994), which encompasses the *petA* genomic region, but where the whole *petA* gene was replaced by the *aadA* cassette. The *petA* CDS, amplified from plasmid pWF (Kuras and Wollman, 1994) using primers AFRF_FW and AFRF_RV and digested by *NcoI* and *PstI*, replaced the *aadA* CDS, excised with *NcoI* and *PstI*. The 5' *psaA-aadA-3' rbcL* selection cassette (Wostrikoff et al., 2004), excised with *SmaI* and *EcoRV*, was introduced at the *HincII* site upstream and in reverse orientation of the chimeric *petA* gene.

Frame-Shifted *petA* Genes: *pf₃₁St* and *pf₁₃₀St*

To create frame shifts after the 31st and 130th codons of the *petA* gene, plasmid pWF was digested by *HindIII* and *BstEII*, Klenow-treated, and religated on itself to yield plasmids *pf₄₂St* and *pf₁₄₅St*. The 5' *psaA*-driven selection cassette was then inserted at the *HincII* site, upstream and in reverse orientation with respect to the *petA* CDS, to yield plasmids *pKf₄₂St* and *pKf₁₄₅St*.

Chimeric *petD* Genes

To generate the plasmid *ppetD*, which contains a single *Swal* site, plasmid pWQ (Kuras and Wollman, 1994) was digested with *SacI* and *AflIII*, Klenow-treated, and religated on itself. To insert the target of NCC2^M within the *petD* 5' UTR or CDS, a PCR fragment of 462 bp was amplified using primers *petD5*::T2_FW and *petD_{Cod}*::T2_RV and plasmid *ppetD* as a template. This amplicon was digested with *Swal* and *HindIII* and the 88-bp fragment was cloned into the *Swal*-*HindIII*-digested vector *ppetD* to create plasmid p5' *petD*::T2 or with *HindIII* and *PstI* and the 358-bp fragment was cloned into *ppetD* digested by the same enzymes to create plasmid *ppetD_{Cod}*::T2.

These two plasmids were then digested by *AvrII*, Klenow-treated, and ligated to a recycling 5' *psaA*-driven spectinomycin resistance cassette. Because the *ncc1* mutation targets the *atpA* mRNA, we avoided using the 5' *atpA*-driven recycling cassette (Fischer et al., 1996) as a selectable marker and constructed a 5' *psaA-aadA-3' rbcL* recycling cassette. The *atpA* 5' UTR, excised from plasmid *paadA485* (Fischer et al., 1996) by *NcoI* and *NruI*, was replaced by the *psaA* 5' UTR, itself excised from plasmid *pfaAK* (Wostrikoff et al., 2004) by *EcoRV* and *NcoI*, yielding plasmid p5' *aA-aadA₄₈₅*. The recycling cassette was excised from this vector by digestion with *SacI* and *KpnI* and Klenow treatment. In the final plasmids pK'5' *petD*::T2 and pK' *petD_{Cod}*::T2, the *aadA* cassette is transcribed in reverse orientation with respect to the *petD* gene.

Mutation of the NCC1^M Target

We mutated the target of NCC1^M by two-step megaprimer PCR (Higuchi, 1990): Primers *atpA_{Ext}*-FW/*atpAMT1RV* and *atpAMT1FW*/*atpA_{Ext}*-RV allowed the amplification from plasmid *patpA2* of two partially overlapping amplicons that were mixed and used as templates in a third PCR with the external primers *atpA_{Ext}*-FW/*atpA_{Ext}*-RV. The final amplicon, digested by *MfeI* and *PacI*, two restriction sites on both sides of the mutation, was cloned into plasmid pK' *atpA₃₀₀*St (Drapier et al., 2007) digested with the same enzymes, yielding plasmid pK' *atpA^M*.

Mutation of the NCC2^M Target

We destroyed the target of NCC2^M by the same two-step PCR procedure using the mutagenic primers *petAMT2FW* and *petAMT2RV* and the external primers *petA_{Ext}*-FW and *petA_{Ext}*-RV to amplify from plasmid template

pWF a 1053-bp fragment. This amplicon, digested by *Bgl*II and *Ac*cl, was cloned into *Bgl*II/*Ac*cl-digested plasmid pKWF_{Stop} (Boulouis et al., 2011) to create plasmid pKpetA_{Mut}T2.

Transformation Vector for Expression of the Mutated NCC1 and NCC2 Proteins

Because of the high percentage of similarity between the paralogous genes of the NCL cluster, designing specific primers to amplify the mutant *ncc1* and *ncc2* genes turned out to be difficult. We thus ordered the synthetic DNA sequences shown in the supplemental data section (Genscript). They were digested by *Eco*RI and *Bgl*II (*ncc1*-HA) or *Eco*RI and *Bam*HI (*ncc2*-HA) and cloned into the vector pJHL (kindly provided by Jae-Hyeok Lee, University of British Columbia) digested by *Eco*RI and *Bam*HI.

All DNA constructs were sequenced before transformation in *Chlamydomonas*. RNA gel blot analyses were performed as described by Drapier et al. (2002) using ³³P-labeled probes described by Eberhard et al. (2002). Transcript accumulation was quantified from phosphor imager scans of the blots, as described by Choquet et al. (2003).

Chloroplast translation was arrested by supplementing cells grown in TAP medium (2×10^6 cells mL⁻¹) with lincomycin (final concentration 500 μ g mL⁻¹) at $t = 0$. Aliquots, taken at the indicated time points, were briefly chilled on ice before RNA extraction.

Map-Based Localization of the *ncc2* Mutation

To localize the *ncc2* mutation, a first set of ~50 *ncc2* progeny (based on their fluorescence phenotype) was selected out of independent meiosis from the cross *ncc2* \times S1-D2. After Chelex-based DNA extraction (Werner and Mergenhagen, 1998), amplified fragment length polymorphism markers (Kathir et al., 2003; Rymarquis et al., 2005) allowed us to determine the proportion of each parental version of the marker on each chromosome arm by PCR. Once linkage (96%) to the ZYS3 marker was established, new amplified fragment length polymorphism markers, based on identified polymorphisms in S1-D2 ESTs or on putative differences in tandem repeat copy numbers, were designed along chromosome 15 (Supplemental Table 2) to define the region containing the *ncc2* mutation. To observe rare crossing-over events, this analysis was performed on 500 independent meioses and allowed us to restrict the location of the mutation to a 405-kb region.

Genomic DNA Preparation, Whole-Genome Sequencing, and Data Analysis

DNA from wild-type and {FAFA} *ncc1 ncc2* double mutant strains was extracted with the DNAeasy Plant Maxi Kit (Qiagen), according to the manufacturer's protocol, starting from 100 mL of stationary culture.

Genome sequencing was performed using the high-throughput, short-read, Illumina technology. Sequencing was done at the Tufts University Core Facility, Boston, MA, on a HiSeq2000 instrument in paired-end mode, 2×100 nucleotides. Libraries had insert sizes of ~300 bp. About 180 million read pairs were generated. Reads were mapped simultaneously onto the nuclear, chloroplast, and mitochondrial genomes of *Chlamydomonas* using BWA 0.6.0-r85 (Li and Durbin, 2009). The organelle genomes were taken from GenBank (accession numbers FJ423446 and CRU03843), and Phytozome v.5 was used for the nuclear genome. The BWA "aln" and "sampe" commands were run with default parameter values, except for the following options: "-q -1 -R 10 -o 2 -e 0 -l 30 -t 20" for "aln" and "-a 600 -n 9999 -N 9999" for "sampe." The alignment files created by BWA in SAM format were then converted to BAM format and indexed, and duplicate read pairs were removed using Samtools 0.1.16 (Li et al., 2009). Read mapping data were visualized in IGV 2.0 (Robinson et al., 2011; Thorvaldsdóttir et al., 2013).

The duplicate-filtered alignment files were then fed to the SVMerge 1.1r32 pipeline (Wong et al., 2010) to detect single nucleotide polymorphisms, short insertions and deletions (indels), and large structural variants. SVMerge included the following tools: breakdancer 1.1 (Chen et al., 2009), pindel 0.2.4q (Ye et al., 2009), cnD 1.3 (Simpson et al., 2010), and SECluster (bundled with SVMerge). The results from InGAP-sv 2.8.1 (Qi et al., 2010; Qi and Zhao, 2011) were also integrated in the SVMerge pipeline. Breakdancer was run with the following options: "-q 35 -c 3 -r 3 -y 40 -m 10000000"; pindel was run with the following options: "-T 1 -x 9 -e 0.02 -u 0.05 -a 1 -m 3 -n 50 -v 50 -d 30 -A 35 -M 6"; cnD was run with the following options: "-threshold=0.5-window=5"; SECluster was run with the following options: "-q 35 -m 6 -c 6 -r {1} -x 10000"; inGAP-sv was run with default parameters, but a maximum coverage of 1000. In the SVMerge configuration file, the following cutoff scores were set: "BDscore=40," "BDrs=3," and "PDscore=200." Velvet 1.1.05 (Zerbino and Birney, 2008) and exonerate 2.2.0 (Slater and Birney, 2005) were run for local assemblies of structural variants in SVMerge. The following Velvet parameters were specifically set: "hashlen=25, exp_cov=auto, cov_cutoff=3." SVMerge requires the BEDtools package (Quinlan and Hall, 2010); version 2.16.2 was used.

Based on version 5.0 of the gene predictions (obtained from Phytozome), single nucleotide polymorphisms and insertions and deletions were annotated using SHOREmap_annotate from the SHOREmap 1.2 package (Schneeberger et al., 2009) to determine in which features they were located (gene, UTR, CDS, and intron) and whether they would cause synonymous or nonsynonymous changes.

The phylogenetic tree was constructed on the Phylogeny.fr platform (Dereeper et al., 2008), including the following steps: Sequences were aligned with MUSCLE (v3.7) (Edgar, 2004) configured for highest accuracy (MUSCLE with default settings; see Supplemental Data Set 2 for the alignment) and cured with Gblocks (v0.91b), using relaxed parameters. The phylogenetic tree was reconstructed using the maximum likelihood method implemented in the PhyML program (v3.0) (Guindon et al., 2010) with reliability for internal branch assessed using the aLRT test (SH-Like) (Anisimova and Gascuel, 2006) and visualized using TreeDyn (v198.3) (Chevenet et al., 2006).

Bioinformatic Analysis of Positive (Diversifying) Selection

Multiple sequence alignment of the NCL proteins was generated with Muscle (Edgar, 2004) and manually refined to optimize conservation of the repeats. Pal2NAL v.14 was used to align the CDS based on this alignment (Suyama et al., 2006) and PhyML to generate the phylogenetic tree using maximum likelihood. A text file of the alignment is provided in Supplemental Data Set 2. Diversifying selection was analyzed using PAML v4.7a (routine codml) as described (Yang, 2007), using the graphic interface in PAMLX (Xu and Yang, 2013). The likelihood of neutral selection models (M1 and M7) was compared with that of models allowing an additional class with $\omega > 1$ (M2 and M8, positive selection). Model 1 allows two classes of codon with $\omega = 0$ (negative selection) and $\omega = 1$ (neutral selection), while model 7 allows a continuous β -distribution of ω values < 1 (Yang and Swanson, 2002). Posterior probability of positive selection at each site was calculated using the Bayes-Empirical Bayes method (Anisimova et al., 2002).

Transformation Experiments

Chloroplast transformation by tungsten particle bombardment (Boynton et al., 1988) was conducted as described (Kuras and Wollman, 1994). Transformants were selected on TAP-Spec (100 mg mL⁻¹) and subcloned on TAP-Spec (500 μ g mL⁻¹) until they reached homoplasmy, assessed by restriction fragment length polymorphism analysis. At least three independent transformants were analyzed for each transformation.

Phenotypic variations between independent transformants proved negligible.

Nuclear transformation of the wild type was performed by electroporation, as described by Raynaud et al. (2007), with the following parameters: 25 μF and 1000 V cm^{-1} . Transformants were selected on plates supplemented with paromomycin (5 $\mu\text{g}\cdot\text{mL}^{-1}$).

Protein Analysis

Pulse-labeling experiments, protein electrophoresis, and immunoblots were performed on exponentially growing cells (2×10^6 cells $\cdot\text{mL}^{-1}$) according to Kuras and Wollman (1994). Cell extracts, loaded on equal chlorophyll basis, were analyzed by SDS-PAGE (12 to 18% acrylamide and 8 M urea). Anti-OEE2 and -cytochrome *f* antibodies, used for [^{35}S] protein A detection, were raised in the laboratory against proteins isolated from *Chlamydomonas* and have been described previously (de Vitry et al., 1989; Kuras and Wollman, 1994). PsbB and HA-tagged NCC1^M and NCC2^M proteins were detected by ECL using the monoclonal antibody anti HA.11 (Covance) and horseradish peroxidase-conjugated antibody against mouse IgG (Promega).

Fluorescence Measurements

Fluorescence measurements were performed on dark-adapted liquid cultures using a home-built spectrofluorimeter according to Zito et al. (1997).

Accession Numbers

Sequence data for *NCC1*, *NCC2*, and *NCL* genes from this article can be found in the Phytozome database, as indicated in Supplemental Data Set 1. Other sequence data used in this article can be found in the GenBank/EMBL databases under the following accession numbers: *petA* (cytochrome *f*), FJ423446.1; *petD*, X72919.1; *psaB*, X05848.1; *psaC*, U43964.1; *psbA*, CAA25670; *psbB* (CP47), X64066.1; *psbD*, X04147.1; *OEE2*, M15187.1; *atpA*, X60298.1; and *rbcl*, J01399.1.

Supplemental Data

Supplemental Figure 1. Phenotype of *ncc1 ncc2* {FAFA} double mutants.

Supplemental Figure 2. Alignment of NCL proteins.

Supplemental Figure 3. Comparison of NCC1 RAP domain model with known structures of endonucleases.

Supplemental Figure 4. I-TASSER alignment used for threading of the NCC1 RAP domain.

Supplemental Table 1. Genetic independence of the *ncc2* mutation from *MCA1* and *TCA1* genes.

Supplemental Table 2. Markers designed to map the *ncc2* mutation.

Supplemental Table 3. Oligonucleotides used in this study.

Supplemental Table 4. BLAST search-based identification of NCL proteins in Phytozome v5.5.

Supplemental References.

Supplemental Material and Methods.

Supplemental Data Set 1. Description of NCL proteins, improved gene models for *NCL7*, *8*, *21*, *30*, and *35*, and intracellular targeting of NCL proteins.

Supplemental Data Set 2. Text file of alignment corresponding to the phylogenetic analysis in Figure 6D.

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AUTHOR CONTRIBUTIONS

A.B., D.D., F.-A.W., and Y.C. designed research. A.B., D.D., H.R., K.W., J.G.-B., and Y.C. performed research. N.J.T., K.P., and O.V. contributed to analytic tools and bioinformatics analysis. A.B., D.D., F.-A.W., and Y.C. analyzed data. A.B., D.D., F.-A.W., and Y.C. wrote the article.

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