A SecY Homologue Is Required for the Elaboration of the Chloroplast Thylakoid Membrane and for Normal Chloroplast Gene Expression

Laura M. Roy and Alice Barkan
Institute of Molecular Biology, University of Oregon, Eugene, Oregon 97403

Abstract. Results of in vitro and genetic studies have provided evidence for four pathways by which proteins are targeted to the chloroplast thylakoid membrane. Although these pathways are initially engaged by distinct substrates and involve some distinct components, an unresolved issue has been whether multiple pathways converge on a common translocation pore in the membrane. A homologue of eubacterial SecY called cpSecY is localized to the thylakoid membrane. Since SecY is a component of a protein-translocating pore in bacteria, cpSecY likely plays an analogous role. To explore the role of cpSecY, we obtained maize mutants with transposon insertions in the corresponding gene. Null cpSecY mutants exhibit a severe loss of thylakoid membrane, differing in this regard from mutants lacking cpSecA. Therefore, cpSecY function is not limited to a translocation step downstream of cpSecA. The phenotype of cpSecY mutants is also much more pleiotropic than that of double mutants in which both the cpSecA- and ΔpH-dependent thylakoid-targeting pathways are disrupted. Therefore, cpSecY function is likely to extend beyond any role it might play in these targeting pathways. CpSecY mutants also exhibit a defect in chloroplast translation, revealing a link between chloroplast membrane biogenesis and chloroplast gene expression.

The majority of chloroplast-localized proteins are associated with the internal membrane system of the organelle, the thylakoid membrane. Many thylakoid membrane proteins are products of nuclear genes and are synthesized in the cytoplasm; others are products of chloroplast genes and are synthesized in the chloroplast stroma. In either case, newly synthesized proteins must be targeted to their correct position with respect to the membrane. Mechanisms of targeting to the thylakoid lumen have been particularly well studied. All known luminal proteins are nuclear-encoded and are synthesized with a bipartite targeting sequence (for review see Cline and Henry, 1996). The NH₂-terminal segment is a stromal targeting sequence that directs the protein across the chloroplast envelope. Adjacent to the stromal targeting sequence is a cleavable luminal targeting sequence that resembles the signal sequences that target proteins for translocation across bacterial cytoplasmic membranes. Two energetically and genetically distinct pathways have been described for the translocation of proteins to the thylakoid lumen (for review see Cline and Henry, 1996). One set of luminal proteins is targeted by a mechanism that requires cpSecA, a chloroplast-localized homologue of the bacterial protein SecA (Yuan et al., 1994; Nohara et al., 1995; Voelker and Barkan, 1995; Voelker et al., 1997). Cytochrome f, a chloroplast-encoded integral thylakoid protein, is also targeted to the membrane via this pathway (Voelker and Barkan, 1995; Mould et al., 1997; Nohara et al., 1997; Voelker et al., 1997). A second set of luminal proteins is targeted via a pathway that relies upon a trans-thylakoidal ΔpH in vitro (for review see Cline and Henry, 1996) and that involves the hcf106 (Voelker and Barkan, 1995) and tha4 genes (our unpublished results) in vivo. A third pathway has been proposed for several integral membrane proteins that appear to integrate spontaneously in vitro (Kim et al., 1996; Robinson et al., 1996). Finally, the integration of the polytopic membrane protein light-harvesting chlorophyll a/b binding protein (LHCP), whose targeting signals lie within the mature portion of the protein (Viitanen et al., 1988), defines a fourth pathway in that it requires neither cpSecA (Voelker et al., 1997) nor hcf106 (Voelker and Barkan, 1995), but does require GTP (Hoffman and Franklin, 1994) and cp54 (Li et al., 1995), a homologue of the signal recognition particle protein SRP54.

1. Abbreviations used in this paper: LHCP, light-harvesting chlorophyll a/b binding protein; OE16, 16-kD subunit of oxygen evolving complex; OE23, 23-kD subunit of oxygen evolving complex; OE33, 33-kD subunit of oxygen evolving complex; PC, plastocyanin; Rubisco, ribulose biphosphate carboxylase; SRP, signal recognition particle; csy1, chloroplast secY 1.
These previous experiments indicated that each targeting pathway involves some unique components. It is possible, however, that the different targeting machineries also have shared components. For example, two or more of the pathways may converge on a common translocation pore in the membrane, in analogy to the convergence of signal recognition particle (SRP)-dependent and -independent pathways on Sec61-containing translocons in the endoplasmic reticulum membrane (for review see Rapoport et al., 1996). One likely component of the translocon for proteins that engage cpSecA is cpSecY, a chloroplast-localized SecY homologue, since SecY forms a component of the translocon in the bacterial plasma membrane (for review see Rapoport et al., 1996). Genes encoding cpSecY have been discovered in algal chloroplast genomes (for review see Vogel et al., 1996) and in the Arabidopsis thaliana and spinach nuclear genomes (Laidler et al., 1995; Berg-hoefer and Kloesgen, 1996). The protein encoded by the Arabidopsis cpSecY cDNA is targeted to chloroplast thylakoid membranes in vitro (Laidler et al., 1995). However, functional studies of cpSecY have not yet been reported.

To gain insight into the roles of cpSecY in vivo, we have used a reverse genetics strategy to obtain maize mutants with transposon insertions in a nuclear gene encoding cpSecY. As shown below, cpSecY mutants exhibit a severe loss of thylakoid membrane. This phenotype is much more severe and global than that of loss of thylakoid membrane. This phenotype is much more severe and global than that of the thylakoid targeting pathway (Fong et al., 1983).

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Cloning of Maize cpSecY cDNAs

A degenerate primer (5′-GAA[GAT][AT][AT][AT][GCAT]-3′) was designed to a conserved amino acid region (met-pro-ile-ile-phe-ser) of cpSecY and then used to prime reverse transcription of maize seedling leaf poly A + RNA by following the manufacturer’s protocol for SuperScript™ II reverse transcriptase (GIBCO BRL, Gaithersburg, MD). PCR amplification was performed on the cDNA template using the reverse transcription primer and a second degenerate primer (5′-GTAAGC/ TGCACAGC/TGTTCAAT/A/TCA/AACTG/C-3′) encoding val-pro-ile-ala-gly. Each 50-μl reaction contained 1 μl of the reverse transcription reaction, 20 pmol of each primer, 2 μM deoxyribonucleoside triphosphate, 1.5 mM MgCl₂, 50 mM KC1, and 10 mM Tris-HCl, pH 8.3. The reactions were incubated at 95°C for 5 min, supplemented with 5 U of Taq DNA polymerase, and then incubated according to the following “touchdown” profile: 10 cycles of 94°C/30 s, 54°C/40 s (+1°C/cycle), 72°C/30 s; 10 cycles of 94°C/30 s, 44°C/40 s (+0.5°C/cycle), 72°C/30 s, 20 cycles of 94°C/30 s, 40°C/40 s, 72°C/60 s. These reactions were subjected to a secondary amplification involving nested degenerate primers that included a 5′ EcoRI site (5′-GGAGAT- TCAT[AA/T][CA][CTG][ATG/G][AG/ATCTG]-3′ encoding ile-asn-ala-ghn-ile-val) and a 5′ BamHI site (5′-CGCGGATCCTG/AAG/ATCG/TCGGAT[AA/T][CTG]-3′ encoding gly-val-val-pro-ile-ala), according to the following conditions: 20 cycles of 95°C/30 s, 48°C/40 s (-0.5°C/cycle), 72°C/60 s; 20 cycles of 95°C/30 s, 39°C/40 s, 72°C/60 s. The amplification product was gel-purified and then subcloned into a Bluescript SK + plasmid. The isolated insert was used to screen a cDNA library constructed from mRNA isolated from the leaves of 2-week-old greenhouse-grown maize, inbred line B73.

Sequence Alignments

Sequences of SecY homologues from Arabidopsis thaliana (EMBL/GenBank/DDBJ accession number U37247), Spinacia oleracea (accession number Z54351), Escherichia coli (accession number 134413), Synechococcus PCC7942 (accession number 401077), cryptomonas (accession number X62348), and Pyrenomonas salina (accession number X74773) were compared to the maize cysl cDNA sequence (accession number AF093904). Alignments were calculated using ClustalW 1.7 (Thompson et al., 1994) and Boxshade (Bioinformatics Group, ISREC, Lausanne, Switzerland), with default parameters.

Isolation and Analysis of Plant DNA and mRNA

DNA was extracted and then analyzed as previously described (Voelker et al., 1997). The genomic Southern blot was probed with a partial cysl cDNA, corresponding to amino acids 205–387 (see Fig. 1) that was radiolabeled by the random hexamer priming method. RNA was extracted and analyzed by Northern blotting as described in Jenkins et al. (1997). The association of chloroplast mRNAs with polysomes was assayed as described previously (Barkan, 1993). The cysl transcript was detected with the full-length cysl cDNA radiolabeled by random hexamer priming. The 16S rRNA and LHCP mRNAs were detected with the radiolabeled DNA probes described in Barkan (1993). The rbcL mRNA was detected with a radiolabeled DNA probe prepared from the cDNA described in Nelson et al. (1984). The atpF probe was described in Jenkins et al. (1997). The rbcL mRNA was detected with a radiolabeled RNA probe prepared by in vitro transcription of a plasmid containing a 570-bp PstI fragment of the maize rbcL gene.

Extraction and Analysis of Protein and Chlorophyll

Leaf proteins were extracted, analyzed on immunoblots, radiolabeled in vivo, and then immunoprecipitated by using the antibodies and methods described previously (Voelker and Barkan, 1995). The relative chlorophyll content of mutant and normal leaves was determined by excising a 35-mg segment from the tip of 10-d-old seedlings, grinding in a mortar and pestle in the presence of 2 ml of 80% acetone, centrifuging at 12,000 g for 1 min, and then measuring the absorbance of the supernatant at 652 nm.

Electron Microscopy

Leaf tissue was fixed overnight at room temperature in 4% glutaraldehyde in 0.05 M potassium phosphate, pH 7.0. After three washes in phos-
Results

The Maize Genome Contains Several Genes That Are Closely Related to secY

We took advantage of the Trait Utility System for Corn developed at Pioneer Hi-Bred (Meeley and Briggs, 1995) to obtain plants with Mu transposons in a nuclear gene encoding cpSecY. A prerequisite for this approach was knowledge of the nucleotide sequence of the maize cpSecY gene(s). To obtain maize cDNAs encoding cpSecY, the deduced Arabidopsis, spinach, Cryptomonas, and Pyrenomonas cpSecY proteins were compared to identify regions of high conservation. A degenerate primer designed against one conserved region was used to prime reverse transcription of maize leaf polyA+ RNA. A second degenerate primer was then used in conjunction with the first to amplify a PCR product. The amplified cDNA fragment was of the expected size (550 bp) and was related in sequence to known cpSecY genes (data not shown). When the PCR fragment was used to screen a maize leaf cDNA library, several full-length cDNA clones (2.1 kbp) were obtained whose DNA sequence revealed a continuous open reading frame encoding a protein with high identity to plastid and eubacterial SecY proteins (Fig. 1).

Arabidopsis and spinach cpSecY are synthesized with an NH2-terminal extension of ~120 amino acids, relative to the bacterial and plastid-encoded SecY homologues (Fig. 1). This region has characteristics typical of chloroplast targeting signals (for review see Cline and Henry, 1996). The maize cDNAs encode a protein with an NH2-terminal extension of a similar length and with features of chloroplast targeting signals. Downstream of this region, the deduced maize protein is highly similar to other chloroplast-localized SecY homologues (e.g., 94% similar/86% identical to Arabidopsis cpSecY). As shown below, the phenotypes associated with mutations in the gene encoding the maize cDNA are restricted to the chloroplast. Together, these observations provide strong evidence that the cDNA encodes maize cpSecY. The gene encoding this cDNA was named csy1.

Genomic Southern blots were probed at high stringency with a cDNA fragment encoding a highly conserved region of cpSecY (Fig. 2). Multiple bands were detected when the genomic DNA was digested with enzymes that do not cut within the genomic sequence corresponding to the probe. These results revealed the existence of at least two and perhaps three genes in the maize genome that are closely related to the cDNA clone. To estimate the relative contributions of these different genes to the pool of cpSecY in leaf tissue, the same probe was used to identify clones in a maize leaf cDNA library. 18 independent cDNA clones were obtained. The nucleotide sequences of the 3′ untranslated regions were identical for all 18 clones, although their sites of polyadenylation varied (Fig. 3 c). A probe made from this 3′ untranslated region detected only one band on genomic Southern blots (data not shown), indicating that all of the cDNAs were derived from the same gene. These results strongly suggest that the csy1 gene contributes the vast majority of cpSecY in leaf tissue.

Mu Insertions Cause Severe Defects in the Expression of csy1

The progeny of maize plants with Mu transposons in csy1 were tentatively identified at Pioneer Hi-Bred in a PCR screen involving csy1-specific primers used in conjunction with a primer corresponding to the Mu terminal inverted repeat (Bensen et al., 1995; Meeley and Briggs, 1995). We confirmed that three of these putative alleles had Mu insertions in csy1. To define the precise position of each insertion, the PCR products resulting from amplification with a Mu primer and a csy1 primer were cloned and then their nucleotide sequences were determined. Fig. 3 shows a partial map of the csy1 gene, including the location of the Mu insertion in each mutant allele. csy1–1 contains a Mu insertion at a 5′ splice junction, in sequences encoding the presumed chloroplast targeting signal. csy1–2 has a Mu insertion in sequences encoding the targeting signal, and csy1–3 has a Mu insertion in sequences encoding the highly conserved portion of cpSecY. As is typical of nuclear-encoded chloroplast proteins, an intron lies between sequences encoding the presumed targeting sequence and sequences encoding the portion of the protein that was derived from the ancestral bacterial gene (refer to Fig. 1).
The insertions in csyl are genetically linked to the pigment-deficient phenotype shown in Fig. 4. This pigment deficiency did not segregate from csyl in >50 meioses. Homozygous mutants are nearly albino, with a distinct yellow-green tint. Heterozygous plants appear entirely normal. All three csyl mutant alleles are similar in phenotype, confirming that the pigment-deficiency results from the Mu insertions in csyl. The chlorophyll content of csyl mutant seedlings grown under relatively low-intensity light (400 μE/m² per s) is ~10% of that of their normal siblings grown in parallel, when normalized to leaf wet weight (data not shown). In contrast, tha1 and hcf106 mutants, with lesions in the cpSecA and ΔpH thylakoid targeting pathways, respectively, accumulate near normal amounts of chlorophyll (Martienssen et al., 1987, and our unpublished observations).

During the development of their first three leaves, csyl mutant plants grow at the same rate as their wild-type siblings (data not shown). The early normal growth rate indicates that mitochondrial function is not significantly disrupted by these mutations and supports the notion that csyl function is limited to the chloroplast compartment. The rate of growth of the mutant seedlings began to lag as seed reserves became depleted and they died after the development of four leaves, as is typical of mutations that disrupt photosynthesis in maize (Miles, 1982).

A single csyl mRNA of ~2.1 kb was detected in RNA prepared from wild-type leaf tissue (Fig. 5). This mRNA

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**Figure 2.** Genomic Southern blot of DNA from the maize inbred line Tx303 probed with the csyl cDNA. The probe consisted of the cDNA fragment encoding amino acids 208–381. BamHI and EcoRI do not recognize sites inside the genomic region corresponding to the probe. The filter was washed at high stringency (65°C in 0.3× SSC).

**Figure 3.** Map and partial nucleotide sequence of csyl. (a) Partial genomic map of csyl. Sites of Mu insertions are indicated. The identity of the Mu family member at each site was deduced based upon the sequence of its terminal-inverted repeat (Chandler and Hardeman, 1992). The insertion sites are shown in the context of the encoded amino acid sequence in Fig. 1. The size of each intron in nucleotides is indicated. (b) Partial nucleotide sequence of csyl. The sequence shown begins 135 nucleotides downstream of the start codon and ends within the third exon. Locations of Mu insertions are shown by boxes marking the nine base pairs that were duplicated upon insertion. Intron sequence is in italics. Only a portion of the sequence of the second intron is shown, with the missing sequence represented by //.

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was not detectable in any of the mutant alleles. A low abundance aberrant transcript of ~3.5 kb in csy1–3 is of the expected size of a chimeric transcript that includes the Mu8 sequences. Two albino mutants, iojap and w3, whose albinism results from different primary defects (Robertson et al., 1978; Walbot and Coe, 1979; Han et al., 1992), and two mutants disrupted in thylakoid protein targeting (tha1 and hcf106), were analyzed in parallel. All of these contain normal levels of csy1 mRNA, indicating that the loss of the mRNA in the csy1 mutants is not a consequence of their pigment deficiency or of a defect in thylakoid protein targeting. The lack of detectable csy1 mRNA in the mutants indicates that the mutations are either null or nearly so.

Chloroplasts in csy1 Mutants Have Little Thylakoid Membrane

The ultrastructure of csy1 mutant chloroplasts was examined by transmission electron microscopy (Fig. 6). Both the bundle sheath and mesophyll chloroplasts in mutant leaves contain little internal membrane. Small internal vesicles were frequently observed (Fig. 6b and data not shown). Mitochondrial morphology is not altered in the mutants (Fig. 6, a–c).

Etioplasts in csy1 mutants are also abnormal in structure (Fig. 6c). Whereas normal etioplasts contain a crystalline prolamellar body with radiating membranes, csy1 mutant etioplasts contain a condensed but noncrystalline prolamellar body that resembles structures observed in maize mutants defective in chlorophyll biosynthesis (Mascia and Robertson, 1978). However, unlike the etioplasts in chlorophyll-biosynthetic mutants, csy1 mutant etioplasts lack the radiating membranes, indicating that their ultrastructural defects are not simply a consequence of the absence of chlorophyll.

The accumulation of representative proteins from each major thylakoid membrane complex was assayed by immunoblot (Fig. 7a). Lesions in either the ΔPH or cpSecA luminal targeting pathway do not significantly reduce the content of the chloroplast ATP synthase or of LHCP (Voelker and Barkan, 1995). In csy1 mutant leaves, however, the abundance of these proteins is ~10-fold lower than normal. The psaD and petD gene products and the D1 protein are reduced to <10% of their normal levels in csy1 mutants. The abundance of these proteins likely reflects the abundance of other subunits of the photosystem I core complex, the cytochrome b6f complex, and the photosystem II core complex, respectively, since subunits of these complexes are generally reduced coordinately in mutants in which the availability of any single subunit is limiting (for review see Barkan et al., 1995).

The accumulation of luminal proteins in csy1 mutant leaves was measured to assess the status of the cpSecA and ΔPH luminal targeting pathways (Fig. 7c). A disruption in luminal targeting is reflected by a decrease in the abundance of the mature protein coupled with an increase in the abundance of its stromal precursor (Voelker and Barkan, 1995). In csy1 mutants, the mature form of three substrates of the cpSecA pathway (the 33-kD subunit of the oxygen evolving complex [OEC33], plastocyanin [PC], and Psaf) accumulate to <10% of their normal levels, and proteins corresponding in size to their stromal intermediates accumulate to unusually high levels. Two substrates of the ΔPH pathway, the 23- and 16-kD subunits of the oxygen evolving complex (OEC23 and OEC16, respectively), behave analogously. These properties are consistent with the possibility that both the ΔPH and cpSecA pathways involve a translocon with a cpSecY subunit. The loss of LHCP is likewise consistent with the idea that cpSecY is required for the integration of proteins that engage cp54. However, it is possible that one or more of these defects is a consequence of the reduction in membrane content, or results because cpSecY is required to target components of other translocans to the membrane.

**cpSecY Mutants Are More Severe in Phenotype Than Mutants Simultaneously Disrupted in Both the cpSecA and ΔPH Targeting Pathways**

Every aspect of the mutant phenotype investigated was more global and severe in csy1 mutants than in tha1 and hcf106 mutants, suggesting that cpSecY function is not limited to either the cpSecA or the ΔPH thylakoid targeting pathway. It remained possible, however, that cpSecY functions in both pathways. If so, and if it has no additional functions, the csy1 mutant phenotype should resemble the phenotype resulting from simultaneous disruption...
of both the cpSecA and ΔpH pathways. To address this possibility, \( \text{tha4/tha1} \) double mutants were constructed. The \( \text{tha1} \) gene encodes cpSecA and the mutation causes at least a 40-fold decrease in cpSecA accumulation (Voelker et al., 1997). The \( \text{tha4} \) mutation disrupts the ΔpH lumenal targeting pathway (our unpublished results and Fig. 8) and its phenotype strongly resembles that of the \( \text{hcf106} \) mutation (Voelker and Barkan, 1995).

Fig. 8 shows that \( \text{tha1/tha4} \) double mutants have a much less severe phenotype than \( \text{csy1} \) mutants. For example, LHCP and RbcL, the large subunit of ribulose bisphosphate carboxylase (Rubisco), accumulate normally in the double mutants but are severely reduced in \( \text{csy1} \) mutants (refer to Fig. 7). The accumulation of the luminal proteins OE16 and OE23 (and their stromal precursors) is similar in \( \text{tha1/tha4} \) double mutants to that in \( \text{tha4} \) single mutants. The accumulation of the luminal proteins OE33 and PC (and their stromal precursors) is similar in \( \text{tha1/tha4} \) double mutants to that in \( \text{tha1} \) single mutants. Components of the ATP synthase, photosystem I, photosystem II, and the cytochrome \( b_{6}f \) complex also accumulate to similar levels in the double and single mutants (Fig. 8, AtpA, PsaD, D1, and PetD, respectively). Finally, the chlorophyll content of double mutant seedlings is ~50% of that in their normal siblings, and is similar to that of \( \text{tha1} \) and \( \text{tha4} \) single mutants (data not shown).

The fact that the \( \text{csy1} \) mutant phenotype is much more global than that of \( \text{tha1/tha4} \) double mutants suggests that \( \text{csy1} \) gene function is not limited to the cpSecA and ΔpH targeting pathways. This finding is consistent with the possibility that cpSecY is part of the translocon for integrating proteins via the cp54 pathway and/or for the proposed

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**Figure 6.** Ultrastructure of \( \text{csy1} \) mutant chloroplasts and etioplasts. (a) Mesophyll chloroplasts. (b) Bundle sheath chloroplasts. (c) Etioplasts. Arrowheads point to mitochondria. Chloroplast ultrastructure was similar in all three mutant alleles (data not shown). Bar, 1 μm.
spontaneous pathway (for review see Cline and Henry, 1996). However, because the thal mutation does not result in the complete loss of thal gene product (Walker, M., and A. Barkan, unpublished results), we cannot eliminate the possibility that a complete disruption of both the ΔpH and cpSecA pathways might mimic the cysI phenotype.

**Chloroplast Translation Is Disrupted in cpSecY Mutants**

We observed that the level of the large subunit of Rubisco is dramatically reduced in cysI mutants grown under day-night cycles (Fig. 7 b). To address the possibility that this was due to photooxidative damage, the RbcL subunit was quantified in leaves that had developed in the absence of light, and in the same leaves after 24 h of exposure to light (Fig. 9). In both cases, RbcL abundance was reduced 10-fold in cysI mutant leaves, indicating that the Rubisco deficiency is not a consequence of photooxidative damage.

The dependence of Rubisco accumulation on cpSecY function was unexpected, given cpSecY’s presumed role in thylakoid membrane biogenesis. Therefore, the basis for its loss was explored in some detail. Rubisco is composed of two types of subunits, a small subunit encoded by nuclear rbcS genes and a large subunit encoded by the chloroplast rbcL gene. A defect in the synthesis or targeting of either the small or large subunit is sufficient to account for the loss of the entire enzyme, since the two subunits accumulate stoichiometrically when the synthesis of one subunit is blocked (Schmidt and Mishkind, 1983; Spreitzer et al., 1985). We considered the possibilities that: (a) cysI is required for normal rates of import of proteins into the chloroplast; (b) cysI is required for normal accumulation of Rubisco mRNAs; and (c) cysI is required for normal chloroplast translation.

Nuclear-encoded proteins that are imported into the chloroplast engage a common translocation machinery that spans the inner and outer envelope membranes (for review see Schnell, 1995; Cline and Henry, 1996). A defect in import is expected to be reflected by a reduced rate of cleavage of stromal targeting sequences. Rates of processing of three nuclear-encoded chloroplast proteins were monitored by pulse labeling leaf proteins in vivo, immunoprecipitation, and then gel fractionation (Fig. 10). There were no detectable differences between wild-type and cysI mutant samples in the rates of accumulation of imported forms of the Rieske protein, PC, and OE33. PC and OE33

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**Figure 7.** Immunoblot analysis of leaf proteins in cysI mutants. 5 μg of total leaf protein (or the indicated dilutions of the wild-type [WT] sample) were fractionated by SDS-PAGE and then transferred to nitrocellulose. (a) Accumulation of integral and membrane extrinsic thylakoid proteins. LHCP, PsaD, D1, and PetD were each detected with monospecific antisera. AtpA, AtpB, and AtpC were detected with sera raised against the CF1 complex. (b) The protein on the blot used in a was visualized by staining with Ponceau S. RbcL is the large subunit of Rubisco. (c) Accumulation of luminal proteins and their precursors. Proteins were detected with the indicated monospecific antisera. iOE23, iOE16, iPC, and iPsaF are of the size predicted for the stromal intermediates of OE23, OE16, PC, and PsaF, respectively. The stromal intermediate of OE33 migrates only slightly slower than the mature form, and can be seen as a doublet in the cysI mutant samples.
To determine whether a disruption of this hypothetical clear genes is dependent upon a signal from chloroplasts, given rise to the notion that the transcription of certain nuclear-encoded mRNAs encoding chloroplast proteins (for review see Taylor, 1989). This has accumulated in several other nuclear-encoded mRNAs encoding chloroplast translation machinery, 16S rRNA in the gradient was less dramatically altered than that of the rbcS mRNA. The distribution of 16S rRNA found in polysomes, although the average polyribosome size is reduced in the mutants. The distribution of the polysome translation machinery, 16S rRNA was quantified by Northern hybridization (Fig. 11). Both mRNAs accumulate to normal levels in csy1 mutants. Therefore, the signal by csy1 mutations could be responsible for the Rubisco deficiency, the LHCP and rbcS mRNAs were quantified by Northern hybridization (Fig. 11). Both mRNAs accumulate to normal levels in csy1 mutants. Since this was not sufficient to account for the 10-fold decrease in Rubisco protein, the status of rbcL mRNA translation was assessed by examining its association with polysomes. Fig. 12 shows that whereas essentially all of the rbcL mRNA is associated with polysomes in normal chloroplasts, only a small proportion is polysome associated in csy1 mutants. The analogous result was obtained for the chloroplast atpF mRNAs (Fig. 12) and psbA mRNA (data not shown). These findings indicate a decreased rate of translation initiation or a stalling of ribosomes at a site early on the mRNAs examined. Previously it was noted that the abundance of the rbcL mRNA is reduced fourfold in maize mutants with defects in chloroplast polysome assembly (Bar- kan, 1993), and it was proposed that this resulted from increased access of ribonucleases to sites that are ordinarily protected by ribosomes. It seems likely, therefore, that the rbcL mRNA deficiency in csy1 mutants is a consequence of its lack of association with ribosomes rather than an independent manifestation of the csy1 defect.

To obtain a more global picture of the status of the chloroplast translation machinery, 16S rRNA was quantified (Fig. 11) and its association with polysomes was examined (Fig. 12). The content of 16S rRNA is reduced approximately fivefold in csy1 mutants, presumably reflecting a similar decrease in the content of 30S ribosomes. PhosphorImager quantification revealed little difference between mutant and wild-type samples in the proportion of 16S rRNA found in polysomes, although the average polyribosome size is reduced in the mutants. The distribution of 16S rRNA in the gradient was less dramatically altered than that of the rbcL, atpF, and psbA mRNAs, suggesting that either the translation of only a subset of chloroplast mRNAs is disrupted in csy1 mutants, or that the translational block is global but that the timing of the block (i.e., initiation, early or late in elongation) varies between mRNAs.
of LHCP and Rubisco (Voelker and Barkan, 1995). Previously, we found that cpSecA, but nonetheless contain normal amounts of thylakoid membrane (Voelker et al., 1997). Should a conservative sorting mechanism exist, it seems plausible that cpSecY mediates the translocation of ΔpH-pathway substrates. Indeed, cysI mutants exhibit a reduced accumulation of mature ΔpH substrates and an increase in the abundance of their stromal precursors. The defects in the ΔpH pathway cannot be attributed to a loss of \( \text{tha}4 \) and \( \text{hcf106} \) gene products because both accumulate to normal levels in cysI mutants (data not shown). On the other hand, the magnitude of the defect in the ΔpH pathway as reflected by the immunoblot assay appears less severe than that in the cpSecA pathway (refer to Fig. 7), suggesting that cpSecY may not function directly in this pathway. The notion that defects in ΔpH substrate processing are only an indirect effect of the absence of cpSecY is supported by the recent discovery of a bacterial secretory mechanism related to the thylakoid–ΔpH pathway that does not require SecY in vivo (Santini et al., 1998).

To address the possibility that cpSecY function does not extend beyond the ΔpH and cpSecA luminal targeting pathways, we compared the phenotype of cpSecY mutants with that of \( \text{tha}1/\text{tha}4 \) double mutants (Fig. 8). The double mutants differ from cysI mutants in that they have a much higher content of chlorophyll and of the core thylakoid membrane complexes, and they are not deficient for LHCP or Ribisco. The much more severe phenotype of cysI mutants suggests that cpSecY performs some function in addition to any role it might play in the SecA and ΔpH pathways. Sec61p, the endoplasmic reticulum homologue of SecY, functions in the translocation of proteins that engage SRP54 (for review see Rapoport et al., 1996). It seems likely, by analogy, that cpSecY mediates the integration of proteins like LHCP that engage cp54. It is also plausible that cpSecY is required for the integration of proteins typified by CFoII, which differ from SecA, ΔpH, and cp54 substrates in that they are capable of integration into protease-treated thylakoid membranes (Robinson et al., 1996). Since cpSecY is predicted to be an integral membrane protein with only limited exposure to the stroma (Laidler et al., 1995; Berghoef er and Kloesgen, 1996), it may be resistant to proteases.

**Discussion**

**cpSecY Function Is Not Limited to the Translocation of Proteins That Engage cpSecA**

In bacteria, secreted proteins that interact with SecA pass through a translocon consisting of SecY, SecE, and SecG (for review see Rapoport et al., 1996). It seems likely that cpSecY is, by analogy, a component of the thylakoid translocon for proteins that engage cpSecA. The cysI mutant phenotype is consistent with this notion. The rate of processing of two cpSecA substrates, PC and OE33, is reduced dramatically in cysI mutants (refer to Fig. 10). In addition, a >10-fold decrease in the accumulation of mature PC and PsaF is accompanied by an increase in the accumulation of their stromal intermediates (refer to Fig. 7). However, we cannot eliminate the possibility that the poor translocation of cpSecA substrates is a consequence of the lack of target membrane. Furthermore, if the signal peptidase itself requires cpSecY to cross the membrane, then the rate of signal-sequence cleavage may not be a reliable indicator of a translocation defect.

It is clear from our results, however, that regardless of whether cpSecY functions in the translocation of proteins that engage cpSecA, it must also play other roles. Previously, we found that \( \text{tha}1 \) mutants contain little, if any, cpSecA, but nonetheless contain normal amounts of thylakoid membrane (Voelker et al., 1997). \( \text{tha}1 \) mutants also differ from cysI mutants in that they contain normal levels of LHCP and Rubisco (Voelker and Barkan, 1995). Therefore, cpSecY function must extend beyond translocating proteins that previously interacted with cpSecA. Our results are consistent with the possibility that cpSecY is an essential component of the thylakoid Sec translocon, whereas cpSecA is merely a facilitator. Alternatively, it seems plausible that cpSecY mediates the translocation of ΔpH-pathway substrates. Indeed, cysI mutants exhibit a reduced accumulation of mature ΔpH substrates and an increase in the abundance of their stromal precursors. The defects in the ΔpH pathway cannot be attributed to a loss of \( \text{tha}4 \) and \( \text{hcf106} \) gene products because both accumulate to normal levels in cysI mutants (data not shown). On the other hand, the magnitude of the defect in the ΔpH pathway as reflected by the immunoblot assay appears less severe than that in the cpSecA pathway (refer to Fig. 7), suggesting that cpSecY may not function directly in this pathway. The notion that defects in ΔpH substrate processing are only an indirect effect of the absence of cpSecY is supported by the recent discovery of a bacterial secretory mechanism related to the thylakoid–ΔpH pathway that does not require SecY in vivo (Santini et al., 1998).
a chloroplast-localized SecY homologue could well be involved. However, the *cysl* gene is clearly not critical for the targeting of all inner envelope proteins because *cysl* mutants are not defective for chloroplast import (refer to Fig. 10), a process that requires several integral components of the inner envelope, including TIC110. In addition, the accumulation of TIC110 protein is unaltered in *cysl* mutants (data not shown). It remains possible that the product of the *cysl* gene functions in the targeting of just a subset of proteins to the inner envelope, or that another of the closely related genes detected by Southern hybridization (refer to Fig. 2) could encode an envelope-localized cpSecY isoform.

**Role of cpSecY in the Elaboration of the Thylakoid Membrane**

The loss of thylakoid membrane in *cysl*− chloroplasts indicates that cpSecY plays a role in membrane accumulation itself. In contrast, the absence of cpSecA in *tha1* mutants does not interfere with the elaboration of the thylakoid membrane (Voelker et al., 1997). Many steps in chloroplast–lipid synthesis are localized to the inner envelope membrane (for review see Joyard et al., 1991). Thus, a loss of thylakoid membrane could result if cpSecY functions in integrating lipid biosynthetic enzymes into the inner envelope. Alternatively, cpSecY may play a role in the transfer of lipids from the inner envelope to the thylakoid membrane. There is evidence that thylakoid membranes initially arise by invagination of the inner envelope membrane, followed by vesicle formation and vesicle fusion (for review see Hoober et al., 1994; Cline and Henry, 1996). A stromal protein has been identified that promotes the fusion of vesicles isolated from chromoplasts (Hugueney et al., 1995). If a similar mechanism exists in chloroplasts, integral membrane proteins are likely to be required to capture these vesicles. cpSecY might be needed for the integration of such proteins. It is interesting in this regard that many of the plastids lacking cpSecY contain small membrane vesicles (refer to Fig. 6 and data not shown), as would be expected if vesicle fusion were disrupted.

**Role of cpSecY in Chloroplast Translation**

An unanticipated aspect of the *cysl* mutant phenotype is the loss of the stromal enzyme Rubisco. This is not a consequence of the chlorophyll deficiency per se, since several maize mutants with more severe chlorophyll deficiencies accumulate near normal levels of Rubisco (Harpster et al., 1984). Likewise, the Rubisco deficiency is not a consequence of photooxidative damage since it is equally severe in dark- and light-grown plastids (refer to Fig. 9). The *rbcS* mRNA accumulates to normal levels in *cysl* mutants (refer to Fig. 11) and the import of nuclear-encoded proteins into the mutant chloroplasts appears normal (refer to Fig. 10). These negative results led us to investigate the possibility that *cysl* mutants have a defect in chloroplast translation. In fact, *cysl*− plastids exhibit a dramatic decrease in the average number of ribosomes associated with the *rbcL* mRNA and with several other chloroplast mRNAs examined (refer to Fig. 12 and data not shown). The fact that the distribution of 16S rRNA in the polysome gradients is not dramatically affected in *cysl* mutants (refer to Fig. 12) suggests that the mutation does not cause a global decrease in the translation initiation rate. Thus, the translation defect in *cysl* mutants differs from that in *cps* mutants, which are defective in chloroplast polysome assembly (Barkan, 1993). The small amount of pre-16S rRNA in *cysl* mutants (refer to Fig. 11) and in all *cps* mutants (Barkan, 1993) is likely the result, rather than the cause, of the translational defects (see Discussion in Barkan, 1993).

This translation defect is intriguing in that it suggests a link between chloroplast membrane biogenesis and chloroplast gene expression. One possibility is that the absence of cpSecY results in the prolonged stalling of ribosomes that are translating membrane proteins, thereby reducing the concentration of free ribosomes available for initiation. A proportion of chloroplast ribosomes is associated with thylakoid membranes (Chua et al., 1973; Margulies and Michaels, 1975; Minami and Watanabe, 1984; Breidenbach et al., 1988), and chloroplast ribosomes pause during the translation of several membrane proteins (Kim et al., 1991; Stollar et al., 1994). It is plausible that the release of some pauses is facilitated by interactions between the ribosome and a proteinaceous receptor in the thylakoid membrane. This would be analogous to the release of SRP-mediated ribosome pausing by interactions with the SRP receptor in the endoplasmic reticulum membrane (for review see Walter and Johnson, 1994). An alternative possibility arises from the recent finding that several chloroplast RNA–binding proteins are associated with internal chloroplast membranes (Zerges and Rochaix, 1998). If translational activators are among these proteins, then the absence or mislocalization of these activators in *cysl* mutants could cause defects in translation. Consistent with this notion are the observations that several translational activators in yeast mitochondria are tightly associated with the mitochondrial inner membrane (for review see Gillham et al., 1994), and that polysomes containing the *rbcL* mRNA have been found in association with thylakoid membranes (Minami and Watanabe, 1984; Breidenbach et al., 1988; Klein et al., 1988).

In summary, the data presented here provide strong evidence that cpSecY functions in more than just the cpSecA-dependent movement of proteins across the thylakoid membrane. Our results suggest further that cpSecY functions in some aspect of thylakoid membrane biogenesis other than translocating proteins to the thylakoid lumen. Finally, analysis of *cysl* mutants has revealed a link between chloroplast translation and the biogenesis of internal chloroplast membranes. A more detailed description of cpSecY function will require biochemical studies and the use of leaky and/or conditional mutations in genes encoding cpSecY.

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