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PNAS 2007;104;5686-5691; originally published online Mar 19, 2007;
doi:10.1073/pnas.0700774104

This information is current as of March 2007.

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Chloroplast iron-sulfur cluster protein maturation requires the essential cysteine desulfurase CpNifS

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Edited by Bob B. Buchanan, University of California, Berkeley, CA, and approved February 12, 2007 (received for review January 27, 2007)

NifS-like proteins provide the sulfur (S) for the formation of iron-sulfur (Fe-S) clusters, an ancient and essential type of cofactor found in all three domains of life. Plants are known to contain two distinct NifS-like proteins, localized in the mitochondria (MtNifS) and the chloroplast (CpNifS). In the chloroplast, five different Fe-S cluster types are required in various proteins. These plastid Fe-S proteins are involved in a variety of biochemical pathways including photosynthetic electron transport and nitrogen and sulfur assimilation. *In vitro*, the chloroplastic cysteine desulfurase CpNifS can release elemental sulfur from cysteine for Fe-S cluster biogenesis in ferredoxin. However, because of the lack of a suitable mutant allele, the role of CpNifS has not been studied thus far *in planta*. To study the role of CpNifS in Fe-S cluster biogenesis *in vivo*, the gene was silenced by using an inducible RNAi (interference) approach. Plants with reduced CpNifS expression exhibited chlorosis, a disorganized chloroplast structure, and stunted growth and eventually became necrotic and died before seed set. Photosynthetic electron transport and carbon dioxide assimilation were severely impaired in the silenced plant lines. The silencing of CpNifS decreased the abundance of all chloroplastic Fe-S proteins tested, representing all five Fe-S cluster types. Mitochondrial Fe-S proteins and respiration were not affected, suggesting that mitochondrial and chloroplastic Fe-S assembly operate independently. These findings indicate that CpNifS is necessary for the maturation of all plastidic Fe-S proteins and, thus, essential for plant growth.

Fe-S proteins | inducible RNAi | photosynthesis | *Arabidopsis thaliana*

NifS-like proteins have cysteine desulfurase activity, which releases elemental sulfur (S) from the amino acid cysteine for the formation of iron-sulfur (Fe-S) clusters. First discovered in the nitrogen-fixing bacterium *Azotobacter vinelandii* (1), NifS-like proteins now represent a conserved group of proteins that are found in all of the domains of life (2). In bacteria, NifS-like proteins can be broadly classified on the basis of primary structure as either group I, which have a proposed general housekeeping role, or group II, which are likely required during oxidative stress. *Escherichia coli* contains 3 NifS-like proteins: IscS (group I), CsdA (group II), and SufS (group II) (for reviews see refs. 2 and 3). It is noteworthy that deletion of IscS is not lethal in *E. coli*; this mild phenotype is attributed to complementation by SufS (4).

In plants, Fe-S proteins are known to exist in the mitochondria, cytosol, and chloroplasts. The chloroplast contains five Fe-S cluster types: ferredoxin (Fd)-type 2Fe-2S, Rieske-type 2Fe-2S, 3Fe-4S, 4Fe-4S, and the siroheme 4Fe-4S (5, 6). These Fe-S clusters are used by an assortment of chloroplastic proteins that are involved in a diverse range of functions including protein import, sulfur and nitrogen reduction, chlorophyll synthesis, and photosynthetic electron transport (5).

Mitochondria and chloroplasts are thought to be the result of separate endosymbiotic events during the evolution of eukaryotes. Plants contain two distinct NifS-like proteins, one localized to mitochondria (MtNifS) (7) and the other localized to chloroplasts (CpNifS) (8, 9). MtNifS (group I) is most similar

to bacterial IscS. The MtNifS homologue in yeast is required for Fe-S cluster formation in the mitochondria and cytosol, and MtNifS may function similarly in plants (7). CpNifS is a group II NifS-like protein (9). *In vitro*, CpNifS can provide the S for Fe-S cluster insertion into apo-Fd to form functional holo-Fd (10). Similar to the bacterial SufS, CpNifS cysteine desulfurase specific activity is low but greatly stimulated by a SufE protein (11, 12).

It may be hypothesized that CpNifS provides the sulfur for all five types of Fe-S proteins that occur in plastids. However, the specific function of CpNifS in chloroplastic Fe-S cluster assembly and its significance for plant development and survival has not been established *in planta* because of the lack of a suitable mutant. To determine the role of CpNifS in the synthesis of chloroplastic Fe-S clusters, we silenced the *Arabidopsis CpNifS* gene by using an ethanol-inducible RNAi construct and investigated the effects on levels of Fe-S proteins and photosynthesis.

Results

CpNifS Is Essential for Plant Growth and Maintenance of Chloroplast Structure. Initially, *CpNifS* was silenced constitutively by RNAi using the cauliflower mosaic virus 35S promoter. Lines in which CpNifS expression was significantly reduced displayed severely chlorotic cotyledons and died as seedlings. Although plants with milder phenotypes could be propagated, we found these constitutively driven *CpNifS* RNAi lines to be unstable for the trait. To avoid these problems, and to be able to study the effects of complete *CpNifS* silencing at a later developmental stage, an ethanol-inducible RNAi construct was used to silence *CpNifS* in *Arabidopsis* plants. Eleven transgenic inducible *CpNifS* RNAi lines were obtained and bred to homozygosity. Two of these lines, *CpNifS-6* and *CpNifS-9*, were selected for further studies. To determine the efficacy of the inducible construct, wild-type (WT) and the two selected transgenic RNAi lines were grown for 2 weeks and then induced by ethanol treatment; control plants were not treated with ethanol. WT and transgenic plants that were not induced with ethanol typically did not show any signs of stress (Fig. 1A) and had normal development and seed production. Indeed, cytosolic ascorbate peroxidase I, a marker of oxidative stress in plants (13), was not induced in ethanol-

Author contributions: M.P. and E.A.H.P.-S. contributed equally to this work; D.V.H., M.P., and E.A.H.P.-S. designed research; D.V.H., S.E.A.-G., C.M.C., S.K.H., and P.K. performed research; S.K.H. and P.K. contributed new reagents/analytic tools; D.V.H. and C.M.C. analyzed data; and D.V.H., M.P., and E.A.H.P.-S. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

Abbreviations: Fd, ferredoxin; MtNifS, mitochondrial NifS-like protein; CpNifS, chloroplastic NifS-like protein; PSI/II, photosystem I/II; ETR, electron-transport rate; RRF, ribosome-recycling factor; Rubisco, ribulose biphosphate carboxylase oxygenase; SiR, sulfite reductase; NiR, nitrite reductase; GOGAT, 2-oxoglutarate aminotransferase.

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This article contains supporting information online at www.pnas.org/cgi/content/full/0700774104/DC1.

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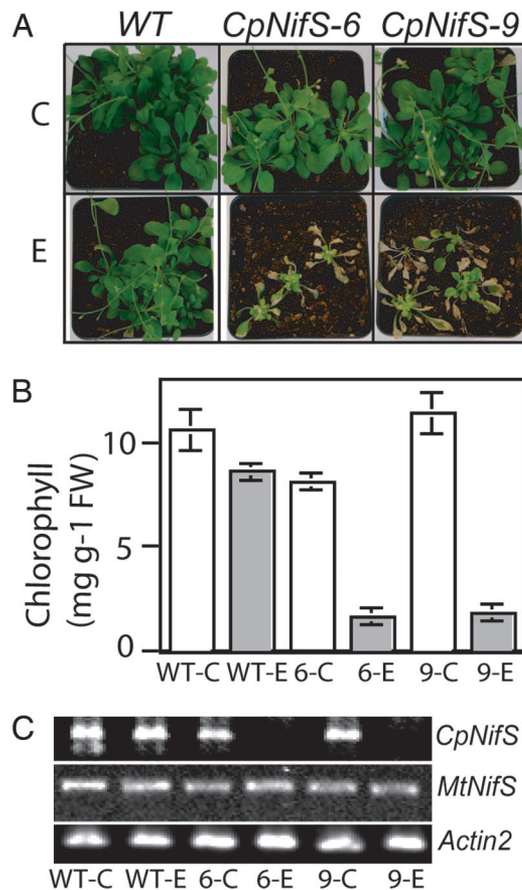


Fig. 1. Phenotypes of *CpNifS*-silenced plants. (A) Two-week old WT, *CpNifS-6*, and *CpNifS-9* plants were treated with 2% ethanol (E) every 4 days for 3 weeks. Control plants (C) were treated with water. (B) Chlorophyll content in plants treated with ethanol or control plants. $P < 0.05$ for ethanol-treated *CpNifS-6* and *CpNifS-9*. (C) *CpNifS*, *MtNifS*, and *Actin2* transcript detection by RT-PCR 1 week after the start of ethanol treatment. FW, fresh weight.

treated plants [see supporting information (SI) Fig. 6]. Only in rare cases was slight chlorosis observed in uninduced transgenics, possibly because of leakiness of the ethanol-inducible promoter. After 3 weeks of ethanol treatment, the *CpNifS-6* and *CpNifS-9* transgenics showed severely stunted growth, chlorosis, and leaf necrosis. Ethanol-treated WT plants showed no such symptoms, albeit that ethanol-treated WT plants were sometimes slightly smaller after 3 weeks of treatment (Fig. 1A). At this stage, the leaves of ethanol-treated *CpNifS-6* and *CpNifS-9* plants had at least 5-fold lower chlorophyll content compared with WT and untreated control plants (Fig. 1B). When ethanol treatment was stopped at this stage, the *CpNifS* plants recovered and were able to set seed. However, continued ethanol treatment ultimately resulted in irreversible damage and death of the RNAi transgenics, whereas it had no visible or only a marginal effect on the WT.

To determine whether the stunted growth of ethanol-treated transgenic plants coincided with the loss of *CpNifS* mRNA, leaf samples were taken from treated and untreated plants at week 3 of plant growth (1 week after the start of ethanol treatment) for RT-PCR analysis. The *CpNifS* transcript was not detected in leaves of chlorotic *CpNifS-6* and *CpNifS-9* plants that were treated with ethanol (Fig. 1C). The mRNAs for *Actin2* and *MtNifS* (NFS1) were not affected by ethanol treatment, showing that the RNAi ethanol-inducible construct is specific for the *CpNifS* gene product.

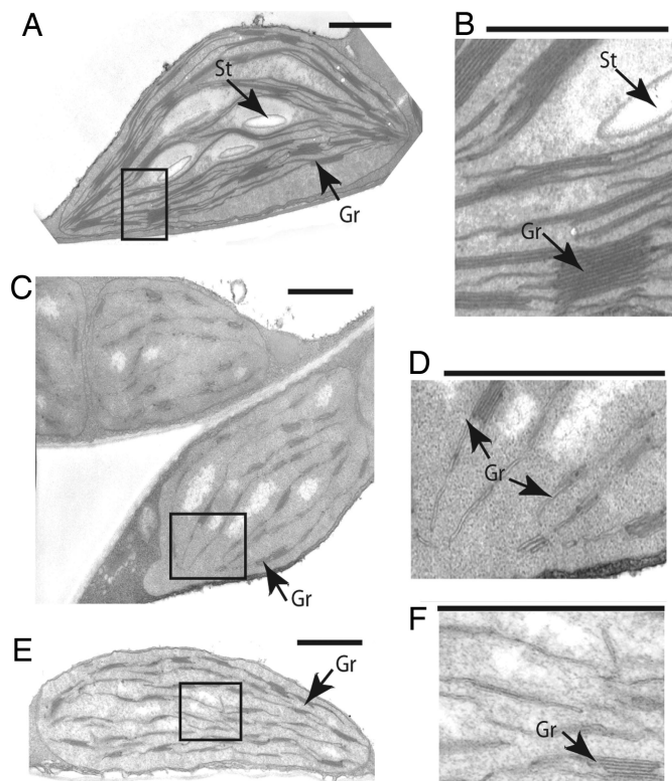


Fig. 2. Chloroplast ultrastructure is altered in *CpNifS*-silenced plants. Leaf samples were fixed 10 days after ethanol treatment of WT (A and B), *CpNifS-6* (C and D), and *CpNifS-9* (E and F) plants. Thin sections were examined by transmission electron microscopy. B, D, and F show magnifications of the boxed areas in A, C, and E, respectively. St, starch granules; Gr, grana. (Scale bars, 1 μ m.)

The severe chlorosis displayed by the ethanol-induced *CpNifS*-silenced plants prompted us to analyze their ultrastructure by transmission electron microscopy to determine the effects of *CpNifS* deficiency on leaf cell and plastid structure. The overall cell shape and the morphology of the nucleus appeared the same in ethanol-treated WT, *CpNifS-6*, and *CpNifS-9* plants; it is also noteworthy that the structure of the mitochondria was unchanged in *CpNifS* plants, suggesting that *MtNifS* and mitochondrial processes were not affected (data not shown). However, chloroplast structure was drastically changed in plants in which *CpNifS* was silenced (Fig. 2). Compared with the discrete and stacked thylakoid membrane grana displayed by WT, the grana in silenced *CpNifS-6* and *CpNifS-9* plants were hypertrophied and dissociated from each other. Also of interest was the absence of starch granules in *CpNifS-6* and *CpNifS-9*, likely caused by a disruption of photosynthesis.

Taken together, these results strongly suggest that *CpNifS* is an essential protein in *Arabidopsis*. *CpNifS* loss of function causes pleiotropic phenotypes and eventually plant death. To analyze the primary cause of these phenotypes and to insight into the direct function of *CpNifS* in plants before pleiotropic phenotypes were apparent, further experiments with the inducible RNAi lines were performed by using plants in which ethanol treatment started 3 weeks after germination, analyzing the plants 10 days after induction.

***CpNifS* Mutants Have Impaired Photosynthesis but Unaltered Respiration.** We hypothesized that *CpNifS* silencing would affect photosynthesis, because photosynthetic electron transport requires many Fe-S proteins, particularly in photosystem I (PSI). To determine how loss of *CpNifS* affects photosynthetic electron

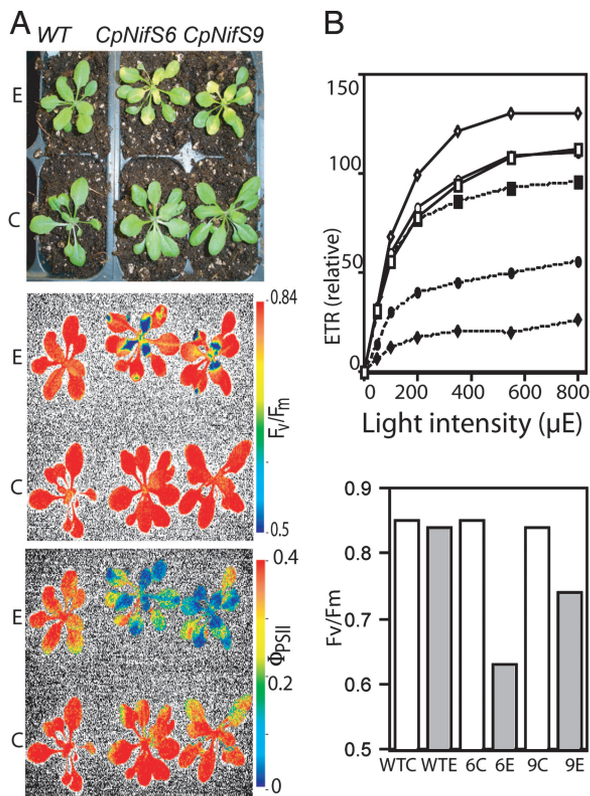


Fig. 3. Chlorophyll fluorescence analysis. (A) Chlorophyll fluorescence imaging. Shown are a regular-color photograph (Top) and false-color images for F_v/F_m (Middle) and Φ_{PSII} (Bottom) of the same plants. False-color scales for fluorescence parameters are shown to the right (red color represents the highest values and blue represents the lowest values). (B Upper) Relative ETR at varying light intensities. Squares, diamonds, and circles represent WT, *CpNifS-6*, and *CpNifS-9*, respectively. Open symbols correspond to untreated controls, and closed symbols signify ethanol treatment. (B Lower) F_v/F_m . Note: standard errors were too small to be plotted. $P < 0.05$ for ethanol-treated *CpNifS-6* and *CpNifS-9*. C, control plants; E, ethanol-treated plants.

transport in intact plants, a chlorophyll fluorescence imaging system was used. For silenced plants and controls, images were captured of two chlorophyll fluorescence parameters: F_v/F_m , which indicates the maximum photochemical efficiency (intactness) of PSII, and Φ_{PSII} , which indicates the fraction of PSII complexes available for photochemistry and indirectly measures the flux of electrons out of PSII (14).

Transgenic plants that were ethanol-treated showed chlorosis, particularly in younger leaves (Fig. 3A). However, the older leaves that had already fully expanded before induction had not become chlorotic at this stage (10 days after induction). Accordingly, when treated with ethanol, F_v/F_m was strongly reduced in the younger leaves of *CpNifS-6* and *CpNifS-9* transgenics compared with WT and untreated controls, but older leaves were less affected (Fig. 3A). The electron flow, Φ_{PSII} , was reduced in both the young and the older leaves of the transgenics, compared with WT.

To gain more quantitative insight into how *CpNifS* silencing affects photosynthesis, Φ_{PSII} was measured and used to estimate the electron-transport rate (ETR) over a range of light intensities in expanded leaves in which chlorosis was absent or minimal. At all light intensities the Φ_{PSII} and, as a consequence, ETR were reduced; ETR saturated at much lower light intensities in *CpNifS-6* and *CpNifS-9* plants induced with ethanol, compared with WT and untreated plants (Fig. 3B). The reduced ETR correlated with a deficiency in *CpNifS* mRNA in the same leaves

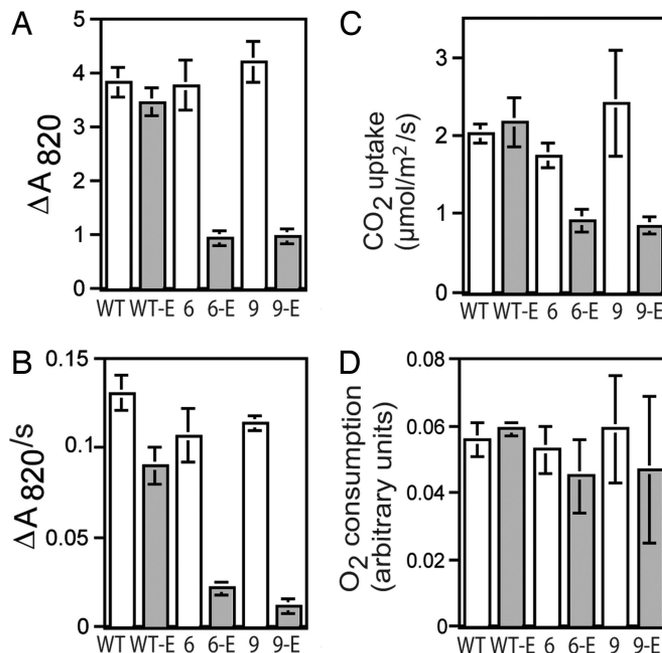


Fig. 4. PSI activity and CO_2 assimilation. WT, 6, and 9 indicate control plants for WT, *CpNifS-6*, and *CpNifS-9* respectively, and WT-E, 6-E, and 9-E indicate the respective ethanol-treated plants. (A) PSI activity. Shown is the extent of the ΔA_{820} (absorbance change at 820 nm, indicative of the P700 activity of PSI) induced with a flash of saturating light, in relative units. (B) Rate of reduction of PSI by upstream electron donors. Values, in relative units, represent the initial rate of the absorbance change at 820 nm ($\Delta A_{820}/s$) in darkness after complete photooxidation of P700, which results mainly from reduction of $P700^+$ by upstream electron donors. (C) Leaf photosynthetic CO_2 uptake. (D) Leaf respiratory O_2 consumption. $P < 0.05$ for ethanol-treated 6-E and 9-E for A–C. All values are the average and standard error of five independent measurements.

(Fig. 1C). The *CpNifS*-silenced plants also showed a decrease in F_v/F_m (Fig. 3C), suggesting photoinhibition or some other effect that reduced the maximum photochemical efficiency of PSII.

All of the Fe-S proteins involved in photosynthetic electron transport are found downstream of PSII: one 2Fe-2S cluster is in the Rieske protein of the cytochrome *b₆f* complex, and three 4Fe-4S clusters are found in PSI (5). The photochemical activity of PSI was analyzed in leaf disks by measuring the light-induced absorbance change at 820 nm (ΔA_{820}), which occurs with photooxidation of the P700 reaction center of PSI (15, 16). Ethanol-treated *CpNifS-6* and *CpNifS-9* plants had less than one third of the ΔA_{820} exhibited by ethanol-treated WT or untreated plants (Fig. 4A), which indicates a substantial loss of photochemically active P700. The rate of dark reduction of P700 after the oxidizing flash was also reduced in the *CpNifS*-silenced plants, indicating that the flow of electrons into PSI from upstream donors was slower (Fig. 4B). To test whether the reduced electron-transport activity limited plant productivity, we measured photosynthetic CO_2 fixation. Indeed, the CO_2 fixation rate in the light was reduced by 50% in the ethanol-treated *CpNifS-6* and *CpNifS-9* plants compared with WT and untreated plants (Fig. 4C). However, oxygen consumption in the dark, which is indicative of mitochondrial respiration, was not affected (Fig. 4D). Thus, *CpNifS* silencing does not appear to disrupt mitochondrial function, while severely affecting chloroplast function.

Levels of Chloroplastic Fe-S Proteins Are Reduced by *CpNifS* Silencing.

We next investigated the direct effect of *CpNifS* silencing on the abundance of chloroplast proteins with or without Fe-S clusters (Fig. 5A). Consistent with an absence of *CpNifS* mRNA, very little CpNifS protein was detected in transgenic plants treated

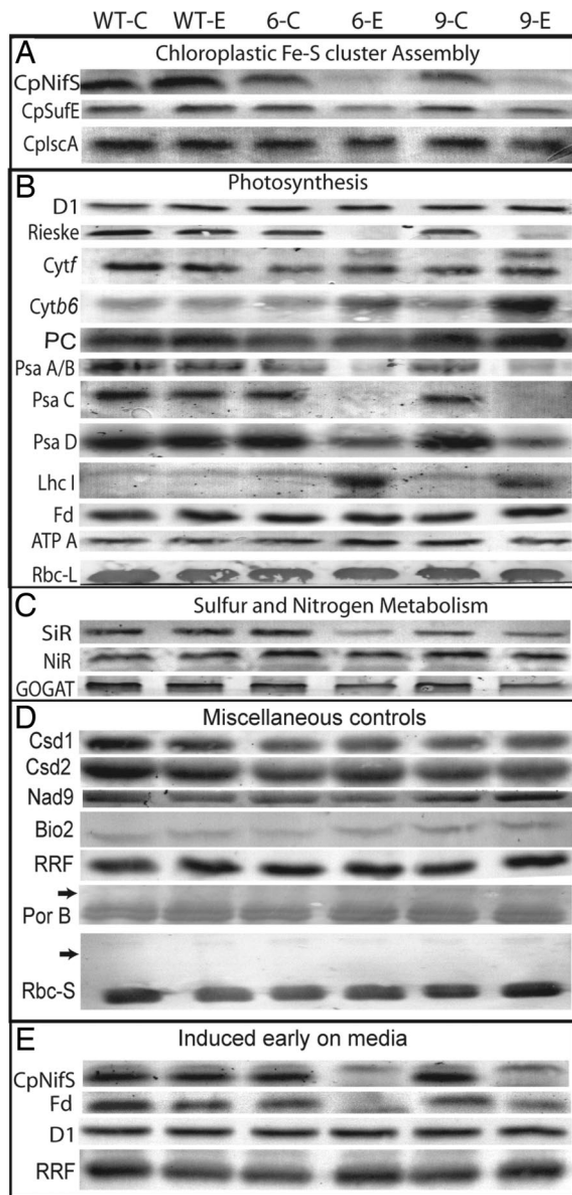


Fig. 5. Effect of *CpNifS* silencing on protein levels. Immunoblot analysis for selected proteins is shown. Lanes labeled WT-C, 6-C, and 9-C indicate control plants for WT, *CpNifS-6*, and *CpNifS-9* respectively, and lanes labeled WT-E, 6-E, and 9-E indicate the respective ethanol-treated plants. (A–D) Samples from plants grown on soil either under control conditions or after treatment with 2% ethanol for 10 days: (A) Chloroplastic Fe-S cluster assembly; (B) photosynthesis; (C) sulfur and nitrogen metabolism; (D) miscellaneous controls. (E) Protein extracts from plants grown from germination on 0.5 Murashige and Skoog medium + 1% sucrose plates with or without ethanol. Total protein extracts (25 μ g of protein) were separated by SDS/PAGE and blotted to nitrocellulose before immunostaining with the indicated antibodies. RRF was used as a loading control. D1, D1 subunit of PSII; PC, plastocyanin; ATP A, ATP synthase subunit A; LhcI, light-harvesting complex proteins of PSI; Rbc-L, large subunit of Rubisco; Csd, Cu/Zn superoxide dismutase; Nad9, subunit 9 of mitochondrial complex 1; Bio2, mitochondrial biotin synthase; PsaA–D, PSI subunits A–D; Por B, protochlorophyllide oxidoreductase; Rbc-S, small subunit of Rubisco. Specific bands were identified by immunostaining and by their molecular weight. Arrows indicate where precursor proteins would have been predicted to accumulate. For each protein, the relevant section of one representative blot of three independent repeats is shown.

with ethanol. The abundance of the *CpNifS* activator, *CpSufE*, was decreased in the absence of *CpNifS*, compared with WT and untreated controls. In contrast, the relative abundance of

CpIscA, a putative scaffold protein for chloroplastic Fe-S cluster formation (17), was hardly affected by *CpNifS* silencing.

To identify which specific steps of photosynthesis may be affected by the silencing of *CpNifS*, we examined the abundance of a variety of proteins involved in photosynthesis (Fig. 5B). *CpNifS* silencing did not affect the D1 protein, a component of the PSII reaction center in which no Fe-S clusters are present (18), suggesting that most PSII was intact. In contrast, components of the cytochrome *b₆f* complex were affected by *CpNifS* silencing. Most noteworthy is the near absence of the 2Fe-2S Rieske protein. Although the abundance of cytochrome *f* was not affected, cytochrome *b₆* was more abundant in *CpNifS*-silenced plants. Both cytochrome *f* and cytochrome *b₆* are heme (Fe) proteins and chloroplast-encoded. A very profound effect of *CpNifS* silencing was observed in PSI. PsaA, PsaB, and PsaC, all 4Fe-4S proteins, were strongly reduced or absent in *CpNifS*-silenced plants. The PsaD protein, which is not an Fe-S cluster protein but thought to be involved in Fd docking (19), was reduced also. This reduction in PsaD abundance may reflect the overall lack of integrity of PSI in *CpNifS*-silenced plants. Perhaps to compensate for the very low abundance of PSI reaction-center proteins in *CpNifS*-silenced plants, there was an increased abundance of subunits of the light-harvesting complex proteins of PSI (LhcI).

Surprisingly, no effect was seen on the abundance of the 2Fe-2S protein Fd in soil-grown *CpNifS*-silenced plants. One possible explanation is that the Fd already present in the 3-week-old plants before ethanol induction was very stable, so that Fd levels did not go down once further production of holo-Fd ceased. To test this possibility, we induced *CpNifS* silencing during germination on media plates. After 1 week, ethanol-treated transgenic plants had extreme chlorosis and stunted growth compared with WT and untreated plants. Western blotting showed that *CpNifS* and Fd were absent in these ethanol-treated transgenics, indicating that Fd levels do decrease if *CpNifS* is silenced at an early stage in plant development (Fig. 5E). Therefore, holo-Fd may be very stable and remain abundant even when the production of its cofactor is impeded. D1, a plastid-encoded subunit of PSII, and ribosome-recycling factor (RRF), which is a nuclear-encoded plastid protein, were not affected by *CpNifS* silencing from germination.

In the soil-grown plants that were ethanol-induced at the 3-week-old stage, the ATP-A subunit of chloroplastic ATP synthase and the large subunit of ribulose biphosphate carboxylase oxygenase (Rubisco), involved in photosynthetic ATP production and CO₂ fixation, respectively, were not affected; neither of these are Fe-S proteins (Fig. 5B). There was also no major effect on plastocyanin, the copper protein that is active in electron transport between cytochrome *b₆f* and PSI (Fig. 5B). Two other copper-containing proteins, Cu/ZnSOD1 and Cu/ZnSOD2, involved in free radical scavenging, were also not affected by *CpNifS* silencing (Fig. 5D).

In addition to the photosynthetic proteins described above, other plastidic Fe-S cluster proteins are expected to depend on *CpNifS* for their maturation. Indeed, the 3Fe-4S cluster protein Fd-glutamine-2-oxoglutarate aminotransferase (GOGAT) was slightly less abundant compared with WT and untreated controls. Moreover, sulfite reductase (SiR) and, to a lesser extent, nitrite reductase (NiR) were reduced in *CpNifS*-silenced plants (Fig. 5C). SiR and NiR contain the siroheme 4Fe-4S cluster type unique to plants. The polypeptide level of NiR was reduced only moderately (30%) compared with WT and untreated plants. In contrast, the enzyme activity of NiR was significantly reduced, by 70% in *CpNifS-6* and 45% in *CpNifS-9*, compared with untreated transgenic plants (SI Fig. 7). The observation that the activity of NiR was reduced to a greater extent than NiR protein levels suggests that apo-NiR, lacking the Fe-S cofactor, had accumulated in *CpNifS*-deficient plants.

The levels of CpNifS and several Fe-S proteins were monitored over a 10-day period after ethanol induction to determine how quickly CpNifS decreased in *CpNifS-6* plants and how quickly it was followed by a decrease in chloroplastic Fe-S proteins. CpNifS began to diminish in transgenic plants 1 day after ethanol induction and had almost completely disappeared by day 10 (SI Fig. 8). Protein abundance of PSI subunits PsaA/B began to decrease on day 2 after ethanol induction, whereas decline of PsaC levels started on day 7. By day 10, a slight reduction in GOGAT, NiR, and SiR was seen. Levels of the control protein RRF remained constant over the course of 10 days.

To determine whether the decrease in chloroplastic Fe-S proteins was caused by a decrease in mRNA, a Northern blot was performed to detect the transcript abundance of Rieske, GOGAT, and SiR. A decrease in mRNA was not observed (SI Fig. 6) despite a large reduction in Rieske, GOGAT, and SiR protein levels (Fig. 5 B and C), which suggests that these Fe-S proteins need their appropriate cofactor to be stable and supports the hypothesis that CpNifS is critical for Fe-S cluster formation in the chloroplast.

In view of the decrease in several abundant Fe-S proteins in the *CpNifS*-silenced lines, leaf nutrient status was investigated (SI Table 1). We did not see a reduction in total leaf iron content on the basis of dry mass; in fact, a modest increase (+25%, nonsignificant) was seen in the ethanol-induced transgenics, compared with ethanol-induced WT. Sulfur and phosphorous levels were significantly increased (by 25% and 50%, respectively) in these same plants.

To rule out the possibility that *CpNifS* plants suffered from a defective chloroplast-import machinery, protein levels of the nuclear encoded protochlorophyllide reductase B and the small subunit of Rubisco were tested to determine whether their mature protein levels are reduced together with accumulation of precursors, as reported for chloroplast-import mutants (20, 21). Both proteins were present in mature size at equal levels in all plant types and treatments. Protochlorophyllide reductase B and the small subunit of Rubisco precursor-sized proteins did not accumulate, suggesting that protein import was not affected by *CpNifS* silencing. Finally, the abundance of two mitochondrial Fe-S proteins, Nad9 (a component of respiratory complex I) and biotin synthase, were not affected by *CpNifS* silencing (Fig. 5D). In summary, *CpNifS* silencing seems to specifically affect the maturation of Fe-S proteins in plastids.

Discussion

Silencing of *CpNifS* severely affected levels of chloroplastic Fe-S proteins and photosynthesis, and prolonged silencing resulted in a pleiotropic-stressed phenotype and eventually plant death. These results suggest that *CpNifS* is an essential protein that functions in plastid Fe-S cluster assembly and cannot be bypassed or complemented by MtNifS. When silencing was induced after the seedling stage, specific and reversible defects could be observed that yielded information about the functions of *CpNifS*. Silencing of *CpNifS* caused a defect in the accumulation of all eight chloroplastic Fe-S cluster proteins that were tested. The Fe-S proteins affected by *CpNifS* silencing together represent all five types of Fe-S clusters found in plastids, supporting the hypothesis that the cysteine desulfurase activity of CpNifS is required for the maturation of all Fe-S proteins in this organelle. Two mitochondrial Fe-S proteins were not affected, lending evidence that Fe-S cluster assembly in the mitochondria can operate independently of the chloroplastic cysteine desulfurase.

Fd did not exhibit any decrease when ethanol was initiated at week 3. However, Fd was absent when *CpNifS* silencing was induced with ethanol from germination, suggesting that holo-Fd is very stable once formed. In contrast to Fd, several other chloroplastic Fe-S proteins had decreased significantly 10 days after *CpNifS* silencing at week 3. These different delays in reduction of Fe-S proteins after *CpNifS* silencing may reflect protein stability or priority of Fe-S cofactor delivery.

The observed defects in photosynthetic electron transport and carbon fixation likely were a consequence of the lack of thylakoid Fe-S proteins, particularly in the cytochrome *b₆f* complex and PSI. Indeed, PSI function was severely compromised after *CpNifS* silencing. At the same time, PSII was only marginally affected in comparison, as evidenced by the presence of the D1 protein and functional heat dissipation, measured as nonphotochemical quenching of chlorophyll fluorescence (data not shown). A comparison of the Φ_{PSII} and Fv/Fm images suggests that Φ_{PSII} was affected before Fv/Fm, which may imply that damage to PSII could be a secondary consequence of a downstream defect in photosynthetic electron transport.

The altered chloroplast ultrastructure observed in *CpNifS*-silenced lines is reminiscent of the ultrastructure reported for an *APO1* mutant (22) that affects PSI accumulation, as well as a mutant in *Hcf101* and other mutations that affect PSI (23, 24). Therefore, the dilated stromal lamellae and absence of grana may be a consequence of a lack of PSI.

Mitochondria and chloroplasts originated from separate endosymbiotic events during the evolution of eukaryotes, and the two organelles have separate NifS-like proteins. It is likely that these two NifS-like proteins with their different properties each evolved to function optimally in their respective environments. The main function of mitochondria is to carry out the oxygen-consuming process of respiration, whereas chloroplasts perform the oxygen-generating process of photosynthesis. Thus, although both organelles contain an electron-transport chain that depends on Fe-S protein assembly, they contrast in redox conditions. Moreover, photosynthesis is known to produce reactive oxygen species, which can lead to oxidative stress. Fe-S cluster biosynthesis is particularly sensitive to oxygen. Therefore, it is not surprising that the chloroplastic NifS that has to operate under high-oxygen conditions is most similar to the bacterial SufS, which is thought to function under oxidative stress (3). MtNifS is most similar to bacterial IscS, the housekeeping NifS-like protein that is more sensitive to oxygen. Therefore, MtNifS likely would not function properly in an oxygen-producing compartment.

Another difference between the chloroplast and mitochondrion is that the chloroplast is the main site of cysteine synthesis in plant cells. It may be important to tightly control the cysteine desulfurase activity of the chloroplastic NifS, to avoid futile cycling. CpNifS may be particularly suited for the chloroplast because its cysteine desulfurase activity is extremely low in the absence of its activator CpSufE, in contrast to group I NifS-like proteins such as IscS and MtNifS (11). In summary, the two Fe-S cluster biosynthesizing machineries in the chloroplast and mitochondrion likely have different evolutionary origins and display properties that fit their function and environmental conditions.

After these two Fe-S biosynthesis machineries came together in the same plant cell, have they shared or transferred some of their functions? As shown here, *CpNifS* silencing is lethal in *Arabidopsis* and affects all five chloroplast Fe-S cluster types. Thus, MtNifS cannot complement the function of CpNifS in the biogenesis of any of these cluster types. At this point, it has not been reported whether MtNifS is essential as well, as was shown to be the case in yeast (25). In our studies, *CpNifS* silencing had no effects on mitochondrial Fe-S protein levels or respiration, suggesting that the mitochondrial Fe-S biogenesis machinery does not depend on CpNifS. Together, these results indicate that in plants, mitochondria and chloroplasts still have separate, essential cysteine desulfurases and Fe-S cluster assembly machineries.

Materials and Methods

Generation of *CpNifS*-Silencing Constructs and Induction. Standard cloning techniques were used to make the plant-transformation constructs and to generate transgenic *Arabidopsis thaliana*. For a detailed description of the cloning steps, see SI Text, Supporting Information on the Cloning and Plant Transformation. The primers

used in plasmid construction and verification are listed in SI Table 2. Ten constitutive RNAi lines and 11 inducible RNAi lines were obtained, which were selfed and propagated to homozygosity. Two of these lines were used for further functional characterization: *CpNifS-RNAi-6* and *CpNifS-RNAi-9* (denoted as *CpNifS-6* and *CpNifS-9*, respectively). To induce the RNAi construct, plants grown in soil were sprayed and soil-drenched every 4 days with a 2% ethanol solution, a concentration that was reported not to induce stress (26, 27); untreated control plants were sprayed with water. For RNAi induction on agar medium, plants were germinated on 0.5 strength Murashige and Skoog medium + 1% sucrose (28) solidified with 0.4% Agargel (Sigma) in 15-cm-diameter Petri dishes with 50 μ l of 100% ethanol (or water for controls) placed in the center at the time of germination.

RT-PCR and Immunoblotting. The presence of transcripts in plants was detected by using RT-PCR (29). Protein extraction, SDS/PAGE, and immunoblot analysis were performed essentially as described (30). Leaf tissue for protein analysis was collected 10 days after induction. Antibodies for CpNifS (9), CpSufE (11), CpIscaA (17), Fd and SiR (31), cytochrome *f*, light-harvesting complex of PSI, and PsaA/B (32), Fd-GOGAT and NiR (33), cytochrome *b₆* and the Rieske subunit (34), PsaC and PsaD (35), protochlorophyllide reductase B and the small subunit of Rubisco (36), and the chloroplastic RRF (37) have been described. Specific antibodies for the D1 subunit of PSII and subunit A of ATP synthase were generous gifts from Alice Barkan (University of Oregon, Eugene) and Anna Sokolenko (Ludwig-Maximilians University, München, Germany), respectively. Specific antibodies for mitochondrial biotin synthase (38) and Nad9 (39) have been described. The intensity of mRNA and protein bands was quantified by using Image J imaging software [National Institutes of Health, Bethesda (<http://rsb.info.nih.gov/ij/>)].

Electron Microscopy. Leaves were sampled from soil-grown plants 10 days after the start of ethanol treatment. Fixation and sectioning

before analysis by transmission electron microscopy were performed as described (40).

Photosynthesis and Respiration Measurements. Chlorophyll content was assayed as described (41). Chlorophyll fluorescence images of Fv/Fm and Φ_{PSII} were captured from control and ethanol-treated dark-adapted soil-grown plants by using a Photon System Instruments imaging system (Photon System Instruments, Brno, Czech Republic). Default protocol settings were used with an actinic light intensity of 110 μ E. A fluorescence monitoring system chlorophyll fluorometer (Hansatech, Cambridge, U.K.) was used for quantitative chlorophyll fluorescence analysis on detached, fully expanded leaves taken from dark-adapted plants. Fv/Fm, Φ_{PSII} , and ETR were calculated as described (14).

Photooxidation and dark-reduction kinetics of P700 (PSI) were measured in leaf disks by determining the light-induced absorbance change at 820 nm (ΔA_{820}) (16, 42) and at a saturating light intensity of 1600 μ E, determined empirically. Carbon assimilation was assayed in detached leaves at 770 μ E by using a Qubit Systems analyzer according to manufacturer instructions (Qubit Systems, Kingston, Ontario, Canada). Oxygen consumption by dark respiration was monitored in leaf tissue with a Hansatech LD2/3 leaf-disk O₂ electrode system maintained at 26°C.

Elemental Analysis, Enzyme Activity, and Statistics. Elemental composition was measured as described previously (43). Enzyme activity of NiR was measured according to ref. 44. All statistical analyses (ANOVA, *t* tests) were performed by using the Jmp-In software package (SAS Institute, Cary, NC).

We acknowledge Syngenta for supplying the ethanol-inducible construct vectors. We are also grateful to the following colleagues for kindly providing antibodies used in this study: Drs. Iwona Adamska, Alice Barkan, Geraldine Bonnard, Antonio Marquez Cabeza, Toshiharu Hase, Norbert Rolland, Henrik Sheller, and Anna Sokolenko. This work was supported by U.S. Department of Agriculture grants USDA-NRI 2003-35318-13758 and 2005-35318-16212 (to M.P. and E.A.H.P.-S).

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