Protein lipoylation in mitochondria requires Fe–S cluster assembly factors NFU4 and NFU5

Jonathan Przybyla-Toscano, Andrew E. Maclean, Marina Franceschetti, Daniela Liebsch, Florence Vignols, Olivier Keech, Nicolas Rouhier and Janneke Balk

1 Université de Lorraine, INRAE, IAM, F-54000 Nancy, France
2 Department of Biological Chemistry, John Innes Centre, Norwich NR4 7UH, UK
3 School of Biological Sciences, University of East Anglia, Norwich NR4 7TJ, UK
4 Department of Plant Physiology, Umeå Plant Science Centre, Umeå University, S-90187 Umeå, Sweden
5 BPMP, Université de Montpellier, CNRS, INRAE, SupAgro, F-34060 Montpellier, France

Abstract

Plants have evolutionarily conserved NifU (NFU)-domain proteins that are targeted to plastids or mitochondria. “Plastid-type” NFU1, NFU2, and NFU3 in Arabidopsis (Arabidopsis thaliana) play a role in iron–sulfur (Fe–S) cluster assembly in this organelle, whereas the type-II NFU4 and NFU5 proteins have not been subjected to mutant studies in any plant species to determine their biological role. Here, we confirmed that NFU4 and NFU5 are targeted to the mitochondria. The proteins were constitutively produced in all parts of the plant, suggesting a housekeeping function. Double nfu4 nfu5 knockout mutants were embryonic lethal, and depletion of NFU4 and NFU5 proteins led to growth arrest of young seedlings. Biochemical analyses revealed that NFU4 and NFU5 are required for lipoylation of the H proteins of the glycine de-carboxylase complex and the E2 subunits of other mitochondrial dehydrogenases, with little impact on Fe–S cluster-containing respiratory complexes or aconitase. Consequently, the Gly-to-Ser ratio was increased in mutant seedlings and early growth improved with elevated CO2 treatment. In addition, pyruvate, 2-oxoglutarate, and branched-chain amino acids accumulated in nfu4 nfu5 mutants, further supporting defects in the other three mitochondrial lipoate-dependent enzyme complexes. NFU4 and NFU5 interacted with mitochondrial lipoyl synthase (LIP1) in yeast 2-hybrid and bimolecular fluorescence complementation assays. These data indicate that NFU4 and NFU5 have a more specific function than previously thought, most likely providing Fe–S clusters to lipoyl synthase.
Introduction

Iron–sulfur (Fe–S) clusters are inorganic cofactors with redox or regulatory functions. These labile metal cofactors are essential for the function of more than hundred different proteins involved in electron transfer, catalysis, or regulatory processes in plants (Przybyla-Toscano et al., 2021a). The biosynthesis of Fe–S clusters requires dedicated assembly proteins that are conserved from bacteria to eukaryotes. Plants contain three Fe–S cluster assembly pathways, the so-called iron–sulfur cluster (ISC) pathway in the mitochondria, the SUF (sulfur mobilization) pathway in the plastids, and the CIA (cytosolic iron–sulfur protein assembly) pathway for the maturation of Fe–S proteins in the cytosol and nucleus (Couturier et al., 2013; Balk and Schaedler, 2014). The mitochondrial ISC pathway provides Fe–S clusters for the respiratory complexes I, II, and III of the electron transport chain, aconitase in the tricarboxylic acid cycle, and several enzymes in biosynthetic pathways (Przybyla-Toscano et al., 2021b).

The proposed working mechanism of the ISC pathway is based on mutant studies in Baker's yeast, Saccharomyces cerevisiae, as well as extensive in vitro studies using orthologs from other eukaryotes (Lill and Freibert, 2020; Przybyla-Toscano et al., 2021b). In brief, and using nomenclature of the plant proteins, the first step of Fe–S cluster assembly is carried out by the cysteine desulfurase NifS-like1 (NFS1), which acquires a protein-bound persulfide (S0) from cysteine. The persulfide is transferred to the scaffold protein ISCU1 (ISU1), where it is reduced to sulfide (S2–) and combined with Fe to form a Fe2S2 cluster. Next, a chaperone–cochaperone couple helps move the cluster from ISU1 to one of several so-called transfer proteins which facilitate incorporation of the correct type of cluster into client Fe–S proteins. The precise molecular roles of these transfer proteins are not known, and research has focused on establishing their sequence of action and whether they are required for specific proteins that are conserved from bacteria to eukaryotes. Plants contain three Fe–S cluster assembly pathways, the so-called iron–sulfur cluster (ISC) pathway in the mitochondria, the SUF (sulfur mobilization) pathway in the plastids, and the CIA (cytosolic iron–sulfur protein assembly) pathway for the maturation of Fe–S proteins in the cytosol and nucleus (Couturier et al., 2013; Balk and Schaedler, 2014). The mitochondrial ISC pathway provides Fe–S clusters for the respiratory complexes I, II, and III of the electron transport chain, aconitase in the tricarboxylic acid cycle, and several enzymes in biosynthetic pathways (Przybyla-Toscano et al., 2021b).

The proposed working mechanism of the ISC pathway is based on mutant studies in Baker’s yeast, Saccharomyces cerevisiae, as well as extensive in vitro studies using orthologs from other eukaryotes (Lill and Freibert, 2020; Przybyla-Toscano et al., 2021b). In brief, and using nomenclature of the plant proteins, the first step of Fe–S cluster assembly is carried out by the cysteine desulfurase NifS-like1 (NFS1), which acquires a protein-bound persulfide (S0) from cysteine. The persulfide is transferred to the scaffold protein ISCU1 (ISU1), where it is reduced to sulfide (S2–) and combined with Fe to form a Fe2S2 cluster. Next, a chaperone–cochaperone couple helps move the cluster from ISU1 to one of several so-called transfer proteins which facilitate incorporation of the correct type of cluster into client Fe–S proteins. The precise molecular roles of these transfer proteins are not known, and research has focused on establishing their sequence of action and whether they are required for specific Fe–S proteins. The mitochondrial glutaredoxin (GRX) S15 is likely to accept the Fe2S2 cluster from ISU1, as shown for yeast GRX5 (reviewed in Lill and Freibert, 2020). In vitro studies demonstrated that Arabidopsis GRXS15 can bind an Fe2S2 cluster and transfer it directly to ferredoxin 1 (Moseler et al., 2015; Ströher et al., 2016; Azam et al., 2020b). These GRXs also mediate the reductive fusion of two Fe2S2 clusters to form a Fe3S4 cluster on the heterodimeric ISCA1/2 proteins (Brancaccio et al., 2014; Azam et al., 2020a; Weiler et al., 2020). Further in vitro studies showed that ISCA1/2 can transfer the Fe3S4 cluster to NFU4 or NFU5, which can donate the cluster to aconitase 2 (Azam et al., 2020a). All mitochondrial NFU proteins studied to date form homodimers with a Fe3S4 cluster bound to the CxxC motifs of each protomer. The Fe–S cluster binding CxxC motif is also found in INDH (ISC protein required for NADH dehydrogenase) which has been proposed to specifically transfer Fe–S clusters to respiratory complex I (Wydro et al., 2013).

The NFU proteins are named after the nifU gene in the nitrogen fixation (nif) regulon in Azotobacter vinelandii (Yuvaniyama et al., 2000). In fact, AvNifU has three protein domains that usually exist as separate proteins in other organisms: the N-terminal domain of AvNifU is homologous to ISU1, the central domain is a functional ferredoxin, and the C-terminal domain is similar to what are generically called NFU proteins (Mühlenhoff and Lill, 2000; Py et al., 2012). Based on phylogenetic analysis, four different types of NFU proteins have been identified in bacteria, of which type II in alpha-proteobacteria has been inherited by mitochondria (Py et al., 2012). In the green lineage, NFU proteins are also present in plastids. They fall outside the bacterial classification system and are characterized by a duplicated NFU sequence, of which the second sequence is degenerate and lacks the CxxC motif. Of the five NFU proteins in Arabidopsis (Arabidopsis thaliana), NFU1, NFU2, and NFU3 belong to the “plastid-type” and are indeed targeted to plastids (León et al., 2003). NFU4 (AT3G20970) and NFU5 (AT1G51390) are type-II NFU proteins. Mitochondrial localization of Arabidopsis NFU4 was supported by transient expression of a GFP fusion protein (León et al., 2003). Furthermore, it was shown that NFU4 and NFU5 are able to complement the yeast nfu1Δ and nfu1Δisu1Δ strains with respect to growth and biochemical phenotypes (León et al., 2003; Uzarska et al., 2018), demonstrating that the plant genes are functional orthologs of yeast NFU1. However, the physiological role of mitochondrial NFU proteins has not been investigated in plants.

Deletion of NFU1 in yeast impairs growth on acetate as carbon source and affects specific Fe–S enzymes, such as aconitase, respiratory complex II (succinate dehydrogenase, SDH), and also lipoyl-dependent enzymes (Melber et al., 2016). The severity of the nfu1Δ phenotype depends on the yeast strain but is relatively mild in contrast to the essential nature of NFU1 in mammals. A small number of mitochondrial disorders in human patients has been associated with rare mutations in the coding sequence of NFU1 (Cameron et al., 2011; Navarro-Sastre et al., 2011; Mayr et al., 2014). NFU1 patients commonly display infantile encephalopathy with symptoms such as hyperglycinemia and lactic acidosis, which in severe cases is fatal in the first year of life. Biochemical tests on muscle biopsies or fibroblasts showed decreased activities of lipoylated enzymes and, in some but not all cases, a decrease in aconitase and SDH activity.

In eukaryotes, there are four mitochondrial lipoyl-dependent enzyme systems, pyruvate dehydrogenase (PDH), α-ketoglutarate dehydrogenase (KGDH), branched-chain α-ketoacid dehydrogenase (BCKDH), and glycine decarboxylase complex (GDC), also referred to as the glycine cleavage system. The PDH, KGDH, and BCKDH complexes consist of three different subunits (E1–E3), of which E2 is lipoylated. The structure of GDC complex differs, being composed of four proteins named L, P, T, and H, with lipoyl bound to the H protein. The lipoyl cofactor (6,8-dithiooctanoic acid) mediates the oxidative decarboxylation reactions carried out by those enzyme complexes (Mayr et al., 2014). The 8-
carbon fatty acid is covalently bound to proteins via a lysine residue, forming a lipoaide, and contains two sulfur atoms at C6 and C8 that are inserted by lipoyl synthase. The sulfur atoms are provided by the auxiliary Fe₄S₄ cluster of lipoyl synthase. Therefore, this cluster needs to be reassembled after every catalytic cycle. In vitro studies showed that the bacterial NfuA protein enabled catalytic rates of lipoyl formation by its ability to reconstitute the auxiliary cluster of lipoyl synthase (McCarthy and Booker, 2017).

Here, we show that the Arabidopsis NFU4 and NFU5 proteins perform an essential function in mitochondria. Analysis of several mutant lines showed that low levels of the NFU proteins are sufficient for normal growth, but their near-complete depletion revealed a defect in protein lipoylation, affecting substrate turnover by lipoate-dependent enzyme complexes such as GDC, KGDH, and BCKDH.

Results

NFU4 and NFU5 are soluble mitochondrial matrix proteins

A previous study showed that transient expression of a fusion protein of the first 116 amino acids of NFU4 with GFP was targeted to mitochondria in Arabidopsis protoplasts (Léon et al., 2003). However, the localization of NFU5 was not experimentally tested. Some evidence for the presence of NFU5 in mitochondria has been obtained by proteomics studies (Ito et al., 2006; Tan et al., 2010; Fuchs et al., 2020), but has so far not been confirmed by other approaches. Therefore, we cloned the predicted mitochondrial targeting sequences (MTS) of NFU4 and NFU5 in frame with the GFP coding sequence. The length of the MTS of each protein was determined by combining several in silico analyses such as target peptide prediction algorithms, the distribution of acidic amino acids, and alignment with eukaryotic and prokaryotic homologs. Following this, amino acids 1–79 for NFU4 and 1–74 for NFU5 were assigned as MTS. The fusion genes were placed under the control of a double CaMV 35S promoter and transformed into Arabidopsis protoplasts. Both NFU4MTS-GFP and NFU5MTS-GFP showed a punctate pattern of GFP which co-localized with MitoTracker dye, but was distinct from the chloroplast autofluorescence (Supplemental Figure S1, A). Protein blot analysis of total and mitochondrial protein fractions confirmed the enrichment of the native NFU4 and NFU5 proteins in mitochondria (Supplemental Figure S1, B). Furthermore, fractionation of the mitochondria into soluble matrix and membrane fractions indicated that NFU4 and NFU5 are matrix proteins, similar to ISU1 and GRXS15 (Supplemental Figure S1, C).

NFU4 and NFU5 proteins are abundant in all plant organs

Antibodies raised against NFU4 cross-reacted with NFU5 and vice versa, which is not surprising as the mature proteins share 90% amino acid identity. To determine which of the two immune signals with similar gel mobilities belongs to NFU4 and NFU5, and for subsequent mutant studies, we obtained three mutant alleles each for nfu4 and nfu5 (Figure 1, A).

Quantitative reverse transcription PCR (RT-qPCR) showed a virtual absence of NFU4 transcripts in homozygous nfu4-2 and nfu4-4 single mutant plants, whereas ~10% transcript remained in the nfu4-1 mutant (Figure 1, B). Expression of NFU5 was strongly diminished in the nfu5-1 line (Figure 1, B). The nfu5-2 mutant had approximately 15% less NFUS transcript compared with its respective wild type (WT), consistent with insertion of the T-DNA in the promoter, 251 nucleotides upstream of the ATG start codon. For the nfu5-3 mutant, RT-qPCR analysis suggested that the transcript levels of NFUS were strongly increased, but this result is likely due to the presence of an outward facing promoter sequence at the right T-DNA border and the RT-qPCR primers being downstream of this. Probing for full-length NFUS transcript by RT-PCR showed that the transcript is lacking in the nfu5-3 mutant (Figure 1, B, right panel), as expected from the position of the T-DNA in exon 2 (Figure 1, A).

Protein blot analysis using anti-NFU4 serum showed that the upper immune signal is absent in the nfu4-2 and nfu4-4 mutants, and that the lower signal is missing in the nfu5-1 and nfu5-3 mutant alleles (Figure 1, C). The native protein products of NFU4 and NFU5 could thus be assigned and their electrophoretic mobilities match the calculated molecular weights of 22.1 kDa for NFU4 and 21.7 kDa for NFU5 without their predicted MTS. The levels of NFU5 protein produced from the nfu5-2 allele were assessed by semi-quantitative protein blot analysis in plants homozygous for the nfu4-2 allele, to rule out signal contribution from NFU4. The NFU5 protein level was ~15% less in the nfu4-2 nfu5-2 double mutant compared with WT (Figure 1, D and Supplemental Figure S2), matching the decrease in NFUS transcripts (Figure 1, B).

Publicly available RNA-seq data (bar.utoronto.ca) indicated that NFU4 and NFUS are expressed throughout the plant's life cycle and in all organs. Protein blot analysis using protein extracts from roots, leaves, stems, and reproductive organs confirmed that NFU4 and NFU5 proteins are expressed in all organs, with slightly lower levels in stems and slightly higher levels in flower buds and flowers, based on total protein (Figure 2, A). Densitometry and quantification of the immune signals relative to known amounts of recombinant protein indicated that NFU4 was approximately two-fold more abundant than NFUS (Figure 2, B), in agreement with a recent quantitative analysis of all mitochondrial proteins (Fuchs et al., 2020).

NFU4 and NFUS are redundant genes, but together are essential for embryo development

In order to evaluate the physiological importance of NFU4 and NFUS, the available T-DNA insertion lines were grown on soil under long-day conditions. The three different mutant lines for NFU4 or NFUS showed normal growth and development of the rosette leaves compared with their respective WT plants (Figure 3, A). The growth rate of...
primary roots in young seedlings, which is particularly sensitive to mitochondrial defects, was decreased by 8% in the nfu4-2 mutant but a decrease in the nfu4-4 mutant was not statistically significant (Supplemental Figure S3, A and B). Root growth in the nfu5-1 mutant was /C24 30% decreased, whereas nfu5-2 root length was similar to WT (Supplemental Figure S3, C and D), as expected based on the weak nature of this mutant allele. In addition, the seedlings were challenged with Paraquat (methyl viologen) in the medium, which strongly inhibited growth of an E. coli nfuAΔ strain (Angelini et al., 2008) and is a well-known inducer of oxidative stress in mitochondria (Cochemé and Murphy, 2008). A concentration of 10 nM Paraquat was chosen based on experimental calibration, resulting in 20% inhibition of WT root growth. However, Paraquat treatment did not specifically affect the mutants more than the WT controls (Supplemental Figure S3). Taken together, these observations indicate that aside from a role in root elongation, individually neither NFU4 nor NFU5 perform a critical function under normal growth conditions.

To generate double mutants, we made reciprocal crosses between nfu5-1 and nfu4-2 or nfu4-4 plants. In the

**Figure 1** Genetic analysis of Arabidopsis mutants in NFU4 and NFU5. A, Gene models of NFU4 and NFU5 and the positions of T-DNA insertions. Black bars represent exons, gray bars are the 5’ and 3’ untranslated regions of the transcript. Triangles represent T-DNA insertions, their orientation is marked with an arrow to indicate the outward facing left border primer. The position of the T-DNA relative to the ATG start codon is indicated by the number of the nucleotide next to the left-border sequence. B, Transcript levels of NFU4 and NFU5 in leaf tissue of WT (Col-0, Col-4, or Ler) and the indicated T-DNA insertion lines, determined by RT-qPCR (graphs) or standard RT-PCR (right). For RT-qPCR, values are the average of three biological samples ± se. C, Protein blot analysis of NFU4 and NFU5 in mitochondria isolated from seedlings. Blots were labeled with antibodies against NFU4. Ponceau S stain was used to confirm equal loading and transfer. D, Decrease in NFU5 protein as a consequence of the nfu5-2 allele, quantified in the nfu4-2 mutant background. See Supplemental Figure S2 for more details of the quantification.
F2 generation, double mutants could not be isolated despite extensive screening by PCR. Therefore, the siliques of nfu4-2−/− nfu5-1−/− and nfu4-4−/− nfu5-1−/− plants were dissected to analyze embryo development. Approximately one quarter of the immature seeds were white (Figure 3, B and C), containing embryos arrested at the globular stage (Figure 3, D). Based on these numbers it is reasonable to assume that the white seeds are double knockout nfu4 nfu5, and that a complete lack of both NFU4 and NFU5 protein is detrimental for embryo development.

Functional relationship between ISU1, NFU4, and NFU5

In yeast, the mild growth defect as a result of NFU1 deletion (nfu1Δ) in an isu1Δnfu1Δ double mutant, underscores the functional relationship of the ISU1 and NFU1 proteins (Schilke et al., 1999). It should be noted that the yeast isu1Δ mutant is viable because of a second ISU gene, ISU2, but evidently this cannot fully replace the function of ISU1. Arabidopsis has three ISU paralogs, each of which can functionally substitute for the yeast ISU1 homolog in the isu1Δnfu1Δ strain (León et al., 2005). The Arabidopsis ISU1 gene is thought to be the main Fe–S scaffold protein in mitochondria, because ISU2 and ISU3 have very low expression levels and are almost exclusively expressed in pollen grains (León et al., 2005; Frazzon et al., 2007). ISU1 protein levels in the isu1-1 line are decreased to ~20% of WT (Supplemental Figure S4, A). ISU1 protein levels in the isu1-1 line were decreased to ~20% of WT (Supplemental Figure S4, B). In cell culture generated from the roots of isu1-1 seedlings, the levels of NFU4, NFU5, INDH, and aconitase were normal, and the activity of respiratory complex I was also comparable to WT (Supplemental Figure S4, B). Interestingly, GRXS15 protein levels were strongly decreased in the isu1-1 mutant.

Rosettes of isu1-1 plants were phenotypically indistinguishable from WT, but seedlings had a 20% decrease in root elongation, which was exacerbated to 40% shorter than WT in the presence of Paraquat (Supplemental Figure S4, D and E). This phenotype may be associated with decreased GRXS15 levels, since grxs15 mutants have strongly impaired root growth (Ströher et al., 2016). To investigate genetic interactions between ISU1 and NFU4 or NFU5, the isu1-1 line was crossed with nfu4 and nfu5 mutant alleles. Crosses involving any of the nfu5 alleles and isu1-1 were unsuccessful, despite multiple attempts in reciprocal combinations. In contrast, double isu1-1 nfu4 mutants were isolated from the F2, and had a shorter root length but not an enhanced growth defect (Supplemental Figure S4, D).

Thus, mild growth impairment of the primary root in seedlings with less than 20% ISU1 is not enhanced by deletion of NFU4. This is in agreement with the model that NFU proteins function downstream from ISU1 in Fe–S cluster transfer and not as an alternative assembly scaffold.

Low amounts of NFU5 alone are sufficient for normal mitochondrial function in vegetative tissues

The normal rosette growth of single nfu4 and nfu5 mutants does not preclude defects in Fe–S cluster enzyme activities. For
example, up to 80% of complex I activity can be lost in Arabidopsis without major growth penalties (Meyer et al., 2011). To analyze the levels of other ISC maturation factors and activities of mitochondrial Fe–S enzymes as a consequence of loss of NFU4 and/or NFU5, mitochondria were purified from 2-week-old seedlings. Protein blot analyses showed no differences in the levels of ISU1, GRXS15, INDH, and aconitase between nfu4-2, nfu4-4 and nfu5-1 and WT (Supplemental Figure S5, A). Labeling with antibodies against lipoate also showed no differences in the abundance of lipo-oylated proteins (Supplemental Figure S5, B). To analyze the levels of mitochondrial respiratory complexes, we generated cell cultures from primary roots, then purified mitochondria for blue-native PAGE. Protein complexes were stained with Coomassie Blue (Supplemental Figure S5, C), and complex I and complex II were additionally visualized by in-gel activity staining (Supplemental Figure S5, D and E). Again, no differences in the levels or the activities of these major Fe–S cluster-dependent complexes were detected in single mutants compared with WT plants.

To obtain viable plants with strongly diminished NFU4 and NFU5 protein levels, mutant alleles were combined via crossing. nfu4-2 nfu5-2 plants lack NFU4 protein and have ~85% NFU5 compared with WT, whereas hemizygous nfu4-4 nfu5-1/nfu5-2 plants have only ~42.5% NFU5 (Figure 4, A). The size and fresh weight of the rosettes of hemizygous

![Figure 3 Phenotypes of nfu4 and nfu5 single and double mutants. A, Growth phenotype of 4-week-old plants of the indicated genotype. Scale bar: 1 cm. B, Images of open silique with immature seeds in WT (Col-0) and the indicated mutant lines. Black arrowheads point at aborted seeds. Scale bars: 0.5 mm. C, Frequency of normal and aborted embryos in nfu4 nfu5-1/+ plants. ***P < 0.0001 for 1:3 segregation ratio (χ² test). D, An aborted and healthy embryo from the silique of a nfu4-2 nfu5-1/+ plant. Plant tissue was cleared with Hoyer’s solution and imaged with DIC microscopy. Scale bars: 50 μm.](image-url)
plants were similar to WT (Figure 4, B and Supplemental Figure S6). In root growth assays, the double mutant seedlings performed similarly to the Col-4 parent. However, in the presence of Paraquat, there was a small but significant decrease in root growth in the hemizygous mutant (Supplemental Figure S3, E). Because of a lipoate biosynthesis defect in yeast and mammalian nfu mutants, we further tested growth of the Arabidopsis nfu4-2 nfu5-2 double mutants under low CO₂ (150 ppm) which should bring out defects in GDC. However, no obvious differences in either fresh or dry weight biomass were observed between mutant and WT plants (Supplemental Figure S6).

To follow up the growth assays, biochemical analyses were carried out on mitochondria purified from cell culture of hemizygous nfu4-2 nfu5-1/nfu5-2 and homozygous nfu4-2 nfu5-2 seedlings. Protein blot analyses showed no differences in the levels of the mitochondrial Fe–S cluster assembly proteins ISU1, GRXS15, and INDH and the Fe–S enzyme aconitase (Figure 4, A). The only observed difference was a decrease in the levels of the H2 protein of GDC. The levels of respiratory complexes I and III in the viable nfu4 nfu5 double mutants were comparable to WT (Figure 4, C). Taken together, these biochemical studies show that NFU4 and NFU5 are largely redundant during vegetative development; and suggest that small amounts of NFU5 are sufficient to sustain key mitochondrial functions.

Depletion of NFU4 and NFU5 causes a pleiotropic growth defect and accumulation of substrates of lipoate-dependent enzyme complexes

To uncover the phenotypic effects of a complete lack of NFU4 and NFU5 in vegetative tissues and circumvent embryo lethality, NFU4 was placed under the control of the ABI3 promoter (Rohde et al., 1999; Despres et al., 2001) and transformed into hemizygous nfu4-2 nfu5-1/nfu5-2 plants (Figure 5, A). The ABI3 promoter drives expression in developing and germinating seeds and can be used to study essential genes (Despres et al., 2001). Positive transformants were selected on hygromycin-containing plates, and three seedlings carrying the nfu5-1 knockout allele...
were identified by PCR for further study (independent lines 1, 4, and 10). In the next generation (T2), seedlings segregated in approximately 21%–25% with a severe growth phenotype and 75%–79% with normal growth. Leaves of the mutant seedlings turned white upon emergence, and growth was arrested at the second pair of true leaves (Figure 5, B). PCR analysis confirmed that the mutant (m) seedlings lacked a functional copy of **NFU5**, whereas the WT-like (wtl) siblings carried the **nfu5-2** allele with an intact **NFU5** open-reading frame (ORF) (Figure 5, C). In agreement with the genotyping results, the mutant seedlings lacked NFU5 protein, whereas the NFU5-specific protein band was present in the wtl seedlings (Figure 5, D). In both mutant and wtl segregants, the protein levels of NFU4 were either undetectable or very low, reflecting the combined effect of the **nfu4-2** allele and **ABI3** promoter-driven NFU4 expression.

In an attempt to overcome growth arrest, the segregating T2 generation was grown under elevated CO₂. We reasoned that the observed photobleaching of the mutant seedlings could be due to a defect in photorespiration. Under high CO₂, photorespiration is prevented and impairment of GDC is tolerated, except when it is so low as to affect one-carbon metabolism (Peterhansel et al., 2010). Indeed, when seedlings were germinated and grown under high CO₂, segregating **nfu4 nfu5** mutant seedlings were more difficult to distinguish from wtl siblings in the first 2 weeks, as double mutants remained green and started to develop a third pair of true leaves (Figure 6, A). However, soon after transferring the seedlings from agar plates to soil in the high CO₂ cabinet, the **nfu4 nfu5** mutants turned white and their growth was arrested before the inflorescence was formed (Supplemental Figure S7).

To investigate if GDC activity was decreased in the mutants, we measured the concentration of glycine and serine as well as other free amino acids using LC–MS. Glycine accumulated eight-fold in the mutant compared with wtl seedlings (Figure 6, B and Supplemental Table S2). The Gly:Ser ratio of 3.5 in the mutant versus a ratio of 1 in wtl is indicative of impaired GDC activity in the mutant (Timm et al., 2012; Reinholdt et al., 2021). It should be noted that the serine levels we measured in control (wtl) seedlings were approximately two-fold lower than routinely measured in rosette leaves of Arabidopsis but the levels of other amino acids were comparable to published values (reviewed in Hildebrandt et al., 2015).

Additionally, the branched-chain amino acids leucine and valine were significantly increased in concentration compared with wtl, and the isoleucine concentration was elevated although not significant (Supplemental Table S2). This suggests that the lipooyl-dependent enzyme complex BCKDH has decreased functionality. Of the other amino acids, alanine and phenylalanine levels were on average more than seven-fold increased in the mutant, but the levels were very variable in the three biological replicates of the mutant. Similarly, arginine, asparagine, lysine, and tryptophan were more than two-fold increased in the mutant, and aspartic acid was more than two-fold decreased, but not statistically significant because of large variation.

To identify changes in pyruvate and TCA cycle intermediates, we measured the concentrations of selected organic acids by LC–MS/MS. The **nfu4 nfu5** mutant accumulated 10-fold more α-ketoglutarate than the wtl segregants and

---

**Figure 5** Seedlings depleted of NFU4 and NFU5 proteins have a pleiotropic phenotype. A, Scheme for obtaining a mutant line expressing NFU4 under the control of the ABI3 promoter in a nfu4 nfu5 knockout background. The ABI3 promoter is active during embryogenesis but switched off after seed germination. The observed segregation numbers in T2 seedling from three independent lines were: 76 chlorotic/small and 260 WT appearance (total n = 336). B, Representative images of a WT seedling and the two classes of segregants, mutant (m) and WT-like (wtl), grown for 21 d on 0.5 MS medium in 8-h light/16-h dark cycles. Scale bars: 0.5 cm. C, PCR genotyping results of mutant (m) and WT-like (wtl) seedlings from three independent lines, showing the absence or presence of a functional NFU5 sequence using primers AM84 and AM85. D, Protein blot analysis of NFU4 and NFU5 in plants lines as in (C).
WT, indicating that the activity of KGDH is impaired (Figure 6, B and Supplemental Table S3). The concentration of pyruvate in the mutant was two-fold higher than in WT seedlings and 1.5-fold higher than in wtl. Citrate and malate concentrations were lower in the *nfu4 nfu5* mutant seedlings, whereas succinate was elevated in both the mutant and wtl segregant compared to WT seedlings (Figure 6, B and Supplemental Table S3). Together, these results indicate
that NFU4 and NFU5 are required for the function of GDC as well as the other major mitochondrial enzyme complexes that depend on lipoate as a cofactor.

NFU4 and NFU5 are required for lipoylation of GDC H proteins and E2 subunits of mitochondrial dehydrogenase enzyme complexes

To investigate a possible decrease in protein-bound lipoyl cofactor in the nfu4 nfu5 mutant, immunoblots of total plant protein extracts were labeled with antibodies against lipoate. This showed a pattern similar to those previously published and proteins were assigned accordingly (Ewald et al., 2014; Ströher et al., 2016; Guan et al., 2017). We observed that nfu4 nfu5 mutant seedlings lacked a signal with an electrophoretic mobility of 15 kDa corresponding to the lipoylated H1 + H3 proteins of GDC (Figure 6, C, upper panel). The mutants also showed a decreased signal at ~65 kDa, assigned as the E2b subunit isoform of the PDH complex, compared with WT and WT-like (wtl) segregants. The signal corresponding to the 82-kDa E2a subunit of PDH was slightly decreased in the mutant compared with WT, but increased in wtl seedlings. The 45 kDa band was assigned to the E2 subunit proteins of plastid-localized PDH, which had similar levels of lipoylation in the mutant and WT plants.

To probe whether non-lipoylated H proteins were present in the mutant lines, a parallel blot of the same samples was probed with an antibody that recognizes H isoforms from a range of plant species including the three isoforms in Arabidopsis (Ströher et al., 2016). The signal from the H1 and H3 isoforms at 15 kDa was strongly decreased in the nfu4 nfu5 mutant, pointing to destabilization of these subunits due to a lack of bound lipoate. The mature H2 protein has a theoretical mass similar to H1 and H3, but as noted before, it has a lower electrophoretic mobility in SDS-PAGE (Figure 6, C, middle panel), possibly because of its unusual low pI (Lee et al., 2008; Ströher et al., 2016). The two signals assigned as H2 likely correspond to the lipoylated and non-lipoylated forms (Ströher et al., 2016). The major H2 isoform present in wtl and WT was barely detectable in nfu4 nfu5 seedlings. These data indicate that NFU4 and NFU5 are required for protein lipoylation in mitochondria, strongly affecting the H subunits of GDC and to a lesser extent the E2 subunits of PDH.

Aconitase and complex II activities are normal in seedlings depleted of NFU4 and NFU5

Since high CO2 conditions only partially rescued the severe phenotype of nfu4 nfu5 mutants, we assessed the overall mitochondrial respiration rate by measuring O2 consumption in intact mutant and wt seedlings in the dark. The nfu4 nfu5 double mutant showed a five-fold decrease in respiration compared with WT seedlings and segregating wt segregants in the dark. The nfu4 nfu5 mutant, in line with defects in key mitochondrial enzyme complexes. In the yeast nfu1Δ mutant, the activities of the Fe–S enzymes aconitase and SDH/complex II were decreased by ~25% (Melber et al., 2016; Uzarska et al, 2018). Measurements of aconitase activity in total plant extracts showed a decrease of ~50% in the nfu4 nfu5 mutant compared with WT seedlings and a decrease of 30% compared with wtl segregants, although the latter was not statistically significant (Figure 7, B). However, in plants grown under high CO2, aconitase activity was two-fold higher than under ambient CO2 levels and similar in the mutant and wtl segregants (Figure 7, B). This suggests that any decrease in aconitase activity may be a secondary defect, caused by, for example, limited carbon flux through the TCA cycle. This is in agreement with the observed decrease in citrate and malate (Figure 6, B).

To measure complex II activity, mutant and wtl segregants were pooled for small-scale mitochondrial preparations. Of the different enzyme assays for complex II, succinate to ubiquinone reduction (SQR) using 2,6 dichloroindophenol as electron acceptor gave the most robust results in relatively impure mitochondrial preparations (León et al., 2007). The activity was measured before and after addition of the complex II inhibitor 2-thenoyltrifluoroacetone (TTFA) to distinguish complex II activity from other succinate reduction reactions. The TTFA-sensitive succinate turnover was similar in all three genotypes, that is nfu1 Δ mutant, wtl segregants and true WT (Figure 7, C). Thus, both aconitase and SDH had a normal operational capacity in nfu4 nfu5 mutant seedlings, in contrast to the decrease in activities of these enzymes in yeast nfu1Δ mutants.

NFU4 and NFU5 proteins interact with the mitochondrial lipoyl synthase LIP1

Previously, Arabidopsis NFU4 and NFU5 were shown to interact with the late-acting ISC proteins ISCA1a and ISCA1b (Azam et al., 2020a). To test if NFU4 and NFU5 can interact with downstream client Fe–S proteins, we carried out yeast two-hybrid assays with mitochondrial lipoyl synthase (LIP1), biotin synthase (BIO2), and the major aconitase protein localized to mitochondria (ACO2). Biotin synthase is similar to lipoyl synthase in that an auxiliary Fe–S cluster is destroyed to donate a sulfur atom in the last step of biotin synthesis, except that this cluster is a Fe2S2 cluster rather than a Fe3S4 cluster. MTS were removed from the coding sequences, which were cloned behind the activation domain (AD) or DNA binding domain (BD) of the Gal4 transcriptional activator. Yeast growth on medium without histidine indicated that the HIS3 reporter gene is transcribed as a consequence of a direct protein interaction between NFU4/5 and LIP1 (Figure 8, A). The growth persisted under more stringent conditions with 3-amino-1,2,4-triazole (3AT), a competitive inhibitor of the HIS3 gene product, suggesting that the interaction between NFU4, or NFU5, with LIP1 is relatively strong. However, no interactions between NFU4/5 and BIO2, nor with ACO2, were detected in these assays.

To assess NFU4/5-LIP1 interactions in plant cells, we additionally performed bimolecular fluorescence complementation (BiFC) assays in Arabidopsis protoplasts using the LIP1 coding region fused upstream of the N-terminal domain of the YFP
protein, and NFU proteins fused upstream of the C-terminal region of YFP (LIP1-N and NFU-C, respectively, in Figure 8, B and Supplemental Figure S9). Positive BiFC signals indicated that LIP1 is in close proximity to NFU4 and NFU5 within plant cells, and that the LIP1 interaction with NFU4 or NFU5 occurs in the mitochondria as highlighted by the overlap of the YFP fluorescence with MitoTracker dye (Figure 8, B, merged panels). None of the proteins expressed alone could restore YFP fluorescence as previously shown (Azam et al., 2020a). Together, these results indicate that NFU4 and NFU5 interact equally well with LIP1 in the mitochondria, and likely assist with the insertion or repair of Fe-S clusters in LIP1.

**Discussion**

The physiological function of the mitochondrial NFU proteins has thus far not been studied in plants, except for one study in sweet potato which found that the gene encoding a mitochondrial NFU was upregulated under salt stress in a salt-tolerant variety (Wang et al., 2013). Here, we show that the mitochondrial NFU proteins perform a key function in lipoate-dependent metabolism in the model plant Arabidopsis, most likely by providing Fe–S clusters to lipoate synthase as shown for bacterial NfuA.

Arabidopsis has two paralogous genes, NFU4 and NFU5, each encoding a functional mitochondrial NFU protein of relatively high abundance (Figures 1–3). While no obvious growth phenotypes were observed in nfu4 or nfu5 single mutants, some of our findings suggest that Arabidopsis NFU4 and NFU5 may not be fully redundant. Single nfu4 and nfu5 mutants had impaired primary root growth in young seedlings (Supplemental Figure S3). Moreover, during reproductive development, nfu5 mutants could not be crossed with the isu1-1 mutant as male or female parent, despite a functional copy of NFU4 (Supplemental Figure S4). Possibly, cell-specific expression of NFU4 and NFU5 in some specialized cell types, or differences in protein stability, could underlie these observations. Related to this is the question of whether NFU4 and NFU5 can form heterodimers or exist exclusively as homodimers. The yeast two-hybrid assay testing for NFU4–NFU5 interaction was negative (Azam et al., 2020a), which is not due to technical issues since other
mitochondrial proteins such as ISCA1/2 (Azam et al., 2020a) and LIP1 (Figure 8, A). Thus, NFU4 and NFU5 are likely functioning as homodimers, with mostly overlapping but perhaps not identical physiological functions, which will be interesting to delineate in future studies.

The double knockout mutant of NFU4 and NFU5 in Arabidopsis is embryonic lethal (Figure 3, B–D), demonstrating that mitochondrial NFUs are essential in plants. The single NFU1 homolog is also essential in mammals but the yeast gene is non-essential. How could this difference be explained? Lipooyl synthase is the main client Fe–S protein in all three types of organisms and an essential enzyme in higher eukaryotes, but not in yeast (Cronan, 2016). This is because lipooyl enzyme complexes can be bypassed, especially when yeast is grown on glucose and switches to fermentation. In bacteria, loss of lipooyl synthase (LipA) can be overcome by supplementation with lipoate in the medium (McCarthy and Booker, 2017), but this cannot rescue yeast lip5 mutants (Sulo and Martin, 1993). Whether plant mutants in lipooyl synthase can be chemically complemented by external lipoate has not been investigated to our knowledge, but we found that addition of lipoate to the medium did not rescue the growth of nfu4 nfu5 mutant seedlings in any way (Supplemental Figure S10). The relatively high

Figure 8 NFU4 and NFU5 proteins interact with LIP1. A, Yeast 2-hybrid analysis to test direct interaction between NFU4/NFU5 and mitochondrial lipooyl synthase (LIP1), biotin synthase (BIO2), and the main mitochondrialaconitase (ACO2). AD-, Gal4 AD; BD-, DNA BD, both at the N-terminal position. Images were taken after 5 d and are representative of at least three independent transformations. B, BiFC to test interaction between NFU4/NFU5 and LIP1. The coding sequences were placed upstream of the N-terminal or C-terminal region of YFP, and the plasmids transformed into Arabidopsis protoplasts. Results are representative of at least two independent transfection experiments and ≥20 fluorescent cells per transformation event. Images are provided with (Figure 8, B) and without (Supplemental Figure S7) maximal Z-stack intensity projections. Scale bars: 10 μm.
abundance of NFU4 and NFU5 compared with other Fe–S cluster maturation proteins would fit with a role of NFU proteins in reassembling the auxiliary cluster of LIP1, which is constantly turned over to insert two sulfurs into lipoyl cofactor. Particularly mitochondria in leaf mesophyll cells have a very high demand of lipote for the H proteins of GDC.

To obtain nfu4 nfu5 mutant lines with a clear phenotype, we used a strategy based on the transcriptional depletion of NFU4 in a nfu4 nfu5 genetic background (Figure 5), after other approaches gave non-viable plants or no visible growth defects (e.g. Figure 4). While the depletion approach was stringent and robust, the seedlings did not develop further than the two-leaf stage, as reported for other essential genes (Rohde et al., 1999; Despres et al., 2001). This limited biochemical assays, especially since it was not possible to isolate mitochondria from the double mutant for enzyme assays. However, protein blot analysis showed a striking decrease in lipoylated proteins, especially the H proteins, and metabolite analysis provided further evidence of blocks in specific mitochondrial processes. While some variation between biological replicates of the mutant, for example in amino acid levels, could have been due to the precise timing of sampling relative to growth arrest, several highly significant changes were noted that have also been observed in other mutants in the lipoate biosynthesis pathway. Specifically, accumulation of glycine and elevated serine has so far been observed in all mutants affected in lipoyl cofactor biosynthesis and its octanoyl precursor (Ewald et al., 2014; Guan et al., 2017; Fu et al., 2020; Guan et al., 2020). Weaker mutant alleles (e.g. RNAi lines or non-essential genes) are primarily affected in the lipoylation of H proteins, whereas stronger alleles also showed decreased lipoylation of E2 subunits of PDH and KGDH. Interestingly, mutation of GRXS15, the mitochondrial GRX associated with the Fe–S cluster assembly pathway, also primarily affects lipoyl biosynthesis but not other Fe–S cluster-dependent processes (Ströher et al., 2016; Moseler et al., 2021). In-depth metabolomics analysis of a grxs15 K83A mutant line showed accumulation of glycine (but not a higher GlySer ratio), accumulaton of branched-chain amino acids, and their ketoacids, a five-fold increase in pyruvate but no significant difference in α-ketoglutarate (Moseler et al., 2021). Decreased lipoylation of the H proteins coincided with lower abundance of the polypeptides, as seen in nfu4 nfu5 mutants (Figure 6, C).

NFU1 mutations in human cells and yeast are known to cause defects in specific Fe–S enzymes, namely complex II and aconitase (Mayr et al., 2014; Melber et al., 2016). In Arabidopsis we saw no marked impairment of the activities of these enzymes in either single mutants or the nfu4 nfu5 double mutant (Figures 4, 7 and Supplemental Figure S5). However, metabolite analysis of young leaves showed accumulation of succinate to similar levels in both nfu4-2 and nfu4-2 nfu5-1 seedlings, suggesting that NFU4 may have a specific role in supporting SDH function. Similarly, aconitase activity was slightly but significantly decreased in the double mutant compared with WT when grown under standard conditions. By contrast, yeast two-hybrid assays were negative for a protein interaction between NFU4 and ACO2 (Figure 8). Interestingly, in vitro, ACO2 can receive an Fe–S cluster from NFU4 or NFU5 (Azam et al., 2020a), but this may simply reflect a possible thermodynamic transition, and not a physiological event. Alternatively, if ACO2 can receive its cluster directly from ISCA1/2 in vivo, this would bypass NFU4/S. It is also possible that the decreased aconitase and complex II activities represent an indirect effect of strong impairment of PDH. Further investigations are necessary to establish if NFU proteins do play a minor role in Fe–S cluster assembly on other proteins than lipoyl synthase.

The mitochondrial NFU proteins differ from their plastid counterparts in both structure and function. The plastid NFU proteins have an extra C-terminal domain which is a copy of NFU but lacking the CxxC motif. The function of this second domain is as yet unclear. Mutant analysis has shown that NFU1, NFU2, and NFU3 have partially overlapping functions required for the stability of Photosystem I and other plastid Fe–S proteins (Touraine et al., 2004; Yabe et al., 2004; Touraine et al., 2019; Berger et al., 2020; Roland et al., 2020). Interestingly, one of the proteins found to interact with NFU2 using yeast two-hybrid and BiFC assays was the plastid isofrom of lipoyl synthase LIP1p (Berger et al., 2020). The levels of lipoyl cofactor on the plastid PDH E2 subunits have not been analyzed to date. Plastid PDH is an essential enzyme in fatty acid synthesis, and defects in this pathway have pleiotropic effects on photosynthesis (Bao et al., 2000; Ke et al., 2000).

In summary, our functional study of the plant mitochondrial NFU proteins reveals their importance for lipoyl cofactor biosynthesis and narrows down the position of these proteins in the downstream pathway of Fe–S cluster assembly.

Materials and methods

Plant material and growth conditions

The following T-DNA insertion mutants were obtained from Arabidopsis (A. thaliana) stock centers (ecotype in brackets, see Supplemental Table S1 for details): isu1-1, SALK_006332 (Col-0); isu1-2, GK_424D02 (Col-0); nfu4-1, SALK_035493 (Col-0); nfu4-2, SALK_061018 (Col-0); nfu4-4, SAIL_1233_C08 (Col-0); nfu5-1, WiscDsloxHs609_06B (Col-0); nfu5-2, SK24394 (Col-4); nfu5-3, GT_3_2834 (Ler). Homozygous plants were selected using genotyping PCR and the T-DNA insertion site was verified by sequencing from the left T-DNA border. For isu1-1 and isu1-2, the right border and contingent genomic DNA was also sequenced. Primers are listed in Supplemental Table S4. The nfu5-1 allele needed to be outcrossed to remove an unrelated leaf phenotype.

Seeds were sown onto Levington’s F2 compost, or they were surface sterilized using chlorine gas and spread on 0.5-strength Murashige and Skoog (MS) medium containing 0.8% (w/v) agar. After vernalization for 2 d at 4°C, plants were grown under long-day conditions (16-h light, 8-h dark) at 22°C and light intensity of 180–200 μmol m−2 s−1.
unless otherwise indicated. The generation and propagation of callus cell culture was performed as previously described (May and Leaver, 1993).

Gene expression analysis
Total RNA was extracted using the Plant RNeasy kit (Qiagen), followed by DNase treatment (Turbo DNase kit, Agilent). The integrity of RNA in all samples was verified using agarose gels and RNA purity was analyzed by comparing 260/230 nm and 260/280 nm absorbance ratios (Nanodrop 2000, Thermo Fisher). RNA was quantified using a Qubit 2.0 fluorometer (Thermo Fisher). RNA (4 μg) was reverse transcribed to cDNA using Superscript III (Thermo Fisher). RT-qPCR reactions were made using SensiFAST master-mix (Bioline), in 20 μL volumes, each with 20 ng of cDNA. Reactions were measured in a Bio-Rad CFX-96 real-time PCR system and cycled as per the Bioline protocol. Data were analyzed using the Bio-Rad CFX Manager 3.1 software, and were normalized using primer efficiency. Primers are listed in Supplemental Table S4. The house-keeping gene SAND (AT2G28390) was used as reference gene (Han et al., 2013).

Protein extraction, gel electrophoresis and protein blot analysis
To extract proteins, plant tissues were ground in cold buffer containing 50 mM Tris–HCl pH 8.0, 50 mM 2-mercaptoethanol, and 1 mM phenylmethylsulfonyl fluoride, then centrifuged for 10 min to remove debris. Mitochondrial proteins from seedlings or cell culture were isolated according to Sweetlove et al. (2007). Blue-native polyacrylamide gel electrophoresis (BN-PAGE) was carried out as described by Wydro et al. (2013). For protein blot analysis, total protein extracts from seedlings or cell culture were isolated according to Winter et al. (1992), with minor modifications. Approximately 30 mg of whole seedling were homogenized in 60 μL of 20 mM HEPES pH 7.0, 5 mM EDTA and 10 mM NaF. After addition of 250 μL of chloroform:methanol (1:5:3.5 volumes) and incubation on ice for 30 min, the water-soluble amino acids were extracted twice with 300 μL of HPLC-grade H₂O. The aqueous phases were combined and stored at −80°C until further analysis. Samples were filtered and diluted 50× in water. Ten microliters of each sample was derivatized using the AccQ tag kit following manufacturer’s instructions (Waters, UK) and 2 μL was used for injection. Derivatized amino acids were separated on a 100 mm × 2.1 mm, 2.7 μm Kinetex XB-C18 column (Phenomenex, USA) in an Acquity UPLC using a 20-min gradient of 1%–20% (v/v) acetonitrile versus 0.1% (v/v) formic acid in water, run at 0.58 mL min⁻¹ at 25°C. The UPLC was attached to a TQS tandem mass spectrometer (Waters, UK) instrument for detection of the correct masses and quantitive measurement.

The organic acids citrate, α-ketoglutarate, malate, pyruvate, and succinate were measured using a recently developed method (Marquis et al., 2017). In brief, 30 mg of whole seedlings were homogenized in 900 μL methanol:H₂O (50:50 by volume, ice cold). After centrifugation at 15,000 × g for 10 min at 4°C, the supernatant was evaporated in a GeneVac EZ-2 Plus (SP Industries, USA) and stored at −80°C until further analysis. Samples were then resuspended in 15 μL of water and derivatized with 50 μL of 10 mM 4-bromo-N-methylbenzamidine in acetonitrile and 25 μL of 1 M 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride in acetonitrile:water (90:10 by volume) for 90 min at 37°C. Samples were run on an Acquity UPLC equipped with a Xevo TQS tandem mass spectrometer (Waters, UK) and a 100 × 2.1 mm 2.6 μm Kinetex EVO C18 column (Phenomenex). The following gradient of 1% (v/v) formic acid in acetonitrile versus 1% (v/v) formic acid in water, at 0.6 mL/min, 40°C was used: (0 min) 40%, (1 min) 40%, (2 min) 45%, (6 min) 95%, (7 min) 95%, (7.5 min) 100%, (8.5 min) 100%, (8.6 min) 40%, and (12 min) 40%.

Oxygen electrode and enzyme assays
Oxygen consumption rates of intact seedlings were measured using an Oxygraph (Hansatech, King’s Lynn, UK). Seedlings were transferred from 0.5 MS agar plates to 2 mL water in the oxygen electrode chamber and kept dark during the measurement. Aconitase activity was determined in total cell extracts in a coupled reaction with isocitrate, as described (Ströher et al., 2016). Succinate:ubiquinone reductase (SQR) activity of complex II, including preparation of mitochondria-enriched fractions from seedlings, was essentially as described by León et al. (2007), except that 10 mM TPTA instead of 1 mM was used as specific complex II inhibitor. Addition of 10 mM TPTA in dimethyl sulfoxide (10 μL in a 1-mL reaction) inhibited SQR activity by 50%–60%, but there was no inhibition with dimethyl sulfoxide alone.

Free amino acids and organic acids
Unbound, water soluble amino acids were extracted according to Winter et al. (1992), with minor modifications. Approximately 30 mg of whole seedling were homogenized in 60 μL of 20 mM HEPES pH 7.0, 5 mM EDTA and 10 mM NaF. After addition of 250 μL of chloroform:methanol (1:5:3.5 volumes) and incubation on ice for 30 min, the water-soluble amino acids were extracted twice with 300 μL of HPLC-grade H₂O. The aqueous phases were combined and stored at −80°C until further analysis. Samples were filtered and diluted 50× in water. Ten microliters of each sample was derivatized using the AccQ tag kit following manufacturer’s instructions (Waters, UK) and 2 μL was used for injection. Derivatized amino acids were separated on a 100 mm × 2.1 mm, 2.7 μm Kinetex XB-C18 column (Phenomenex, USA) in an Acquity UPLC using a 20-min gradient of 1%–20% (v/v) acetonitrile versus 0.1% (v/v) formic acid in water, run at 0.58 mL min⁻¹ at 25°C. The UPLC was attached to a TQS tandem mass spectrometer (Waters, UK) instrument for detection of the correct masses and quantitive measurement.

The organic acids citrate, α-ketoglutarate, malate, pyruvate, and succinate were measured using a recently developed method (Marquis et al., 2017). In brief, 30 mg of whole seedlings were homogenized in 900 μL methanol:H₂O (50:50 by volume, ice cold). After centrifugation at 15,000 × g for 10 min at 4°C, the supernatant was evaporated in a GeneVac EZ-2 Plus (SP Industries, USA) and stored at −80°C until further analysis. Samples were then resuspended in 15 μL of water and derivatized with 50 μL of 10 mM 4-bromo-N-methylbenzamidine in acetonitrile and 25 μL of 1 M 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride in acetonitrile:water (90:10 by volume) for 90 min at 37°C. Samples were run on an Acquity UPLC equipped with a Xevo TQS tandem mass spectrometer (Waters, UK) and a 100 × 2.1 mm 2.6 μm Kinetex EVO C18 column (Phenomenex). The following gradient of 1% (v/v) formic acid in acetonitrile versus 1% (v/v) formic acid in water, at 0.6 mL/min, 40°C was used: (0 min) 40%, (1 min) 40%, (2 min) 45%, (6 min) 95%, (7 min) 95%, (7.5 min) 100%, (8.5 min) 100%, (8.6 min) 40%, and (12 min) 40%.

Oxygen electrode and enzyme assays
Oxygen consumption rates of intact seedlings were measured using an Oxygraph (Hansatech, King’s Lynn, UK). Seedlings were transferred from 0.5 MS agar plates to 2 mL water in the oxygen electrode chamber and kept dark during the measurement. Aconitase activity was determined in total cell extracts in a coupled reaction with isocitrate, as described (Ströher et al., 2016). Succinate:ubiquinone reductase (SQR) activity of complex II, including preparation of mitochondria-enriched fractions from seedlings, was essentially as described by León et al. (2007), except that 10 mM TPTA instead of 1 mM was used as specific complex II inhibitor. Addition of 10 mM TPTA in dimethyl sulfoxide (10 μL in a 1-mL reaction) inhibited SQR activity by 50%–60%, but there was no inhibition with dimethyl sulfoxide alone.

Yeast 2-hybrid assays
ORFs corresponding to the presumed mature forms of NFU4, NFU5, LIP1, BIO2, ACO2 (primers in Supplemental Table S4)
were cloned into the pGADT7 and pGBK7 vectors (Clontech) between the NdeI or Ncol and BamHI restriction sites in order to express N-terminal fusion proteins with the GAL4 DNA BD or with the GAL4 AD, respectively, under the control of the ADH1 promoter. The plasmids were cotransformed by heat shock treatment into the GAL4-based yeast 2-hybrid reporter strain CY306, which is deficient for cytosolic thioredoxins (MATα, ura3-52, his3-200, ade2-101, lys2-801, leu2-3, leu2-112, trp1-901, gal4-542, gal80-538, lys2::UASGal1-TATAGAL1-HIS3, URA3::UASGal4 17MERS(x3)-TATACYC1-LacZ, trz1::KanMX4, trz2::KanMX4 (Vignols et al., 2005). Transformants were selected on yeast nitrogen base (YNB) medium (0.7% (w/v) yeast extract without amino acids, 2% glucose, 2% agar) without tryptophan and leucine (-Trp, -Leu). Two clones were selected and interacted acids, 2% glucose, 2% agar) without tryptophan and leucine (-Trp, -Leu). Two clones were selected and interacted}

were transformed by heat shock treatment into the GAL4-based yeast 2-hybrid reporter strain CY306, which is deficient for cytosolic thioredoxins (MATα, ura3-52, his3-200, ade2-101, lys2-801, leu2-3, leu2-112, trp1-901, gal4-542, gal80-538, lys2::UASGal1-TATAGAL1-HIS3, URA3::UASGal4 17MERS(x3)-TATACYC1-LacZ, trz1::KanMX4, trz2::KanMX4 (Vignols et al., 2005). Transformants were selected on yeast nitrogen base (YNB) medium (0.7% (w/v) yeast extract without amino acids, 2% glucose, 2% agar) without tryptophan and leucine (-Trp, -Leu). Two clones were selected and interacted

absence of histidine (-His-Trp-Leu). Two clones were selected and interacted acids, 2% glucose, 2% agar) without tryptophan and leucine (-Trp, -Leu). Two clones were selected and interacted

 Confocal microscopy
For localization of GFP-fusion proteins, full-length ORFs of NFU4 and NFU5, or the predicted MTS (amino acids 1–74 for NFU5 and 1–79 for NFU4) were amplified from Arabidopsis leaf cDNA and cloned between Ncol and BamHI sites of pCK-GFP-C65T (2 × CaMV 35S promoter, C-terminal GFP (Reichel et al., 1996)). For BiFC, full-length ORFs of NFU4, NFU5, and LIP1 were amplified and cloned upstream of the C-terminal and N-terminal regions of the mVenus YFP protein into the pUC-SPYC and pUC-SPYNE vectors (Walter et al., 2004, abbreviated -C and -N in figures, respectively) using the restriction sites XhoI and XhoI or Smal. Primers for amplification are listed in Supplemental Table S4.

 Leaf protoplasts were prepared from 21- to 28-d-old Arabidopsis seedlings, transfected according to Yoo et al. (2007) and imaged after 20–24 h. Prior to confocal analyses, the protoplasts were incubated with 20 nM MitoTracker Orange CMXRos (ThermoFisher) to fluorescently label the mitochondria.

 Image acquisition for NFU-GFP localization was performed using a Zeiss LSM780 confocal laser scanning microscope, and a water-corrected C-Apochromat 40 × objective with a numerical aperture (NA) of 1.2. Signals were detected according to the following excitation/emission wavelengths: GFP (488 nm/494–534 nm), MitoTracker (561 nm/569–613 nm) and chloroplast autofluorescence (488 and 561 nm/671–720 nm). Pictures were analyzed using ImageJ software (https://imagej.nih.gov/ij/).

 Image acquisition for BiFC was performed on a Leica TCS SP8 confocal laser-scanning microscope using a water × 40 objective. Waveforms for excitation/emission were: mVenus (514 nm/520–550 nm), MitoTracker (560 nm/580–620). Images were obtained using LAS X software and processed with Adobe Photoshop CS3 at high resolution. All confocal images shown here were captured without Z-stack intensity projection.

 Statistical analysis
Graphs and statistical analyses were done in MS Excel or in R. Details regarding the number of replicates and the statistical tests are given in figure legends.

 Accession numbers
Accession numbers are as follows: AT3G20970, NFU4; AT1G51390, NFU5; AT4G22220, ISU1; AT2G20860, LIP1.

 Supplemental data
The following materials are available in the online version of this article.

 Supplemental Figure S1. NFU4 and NFU5 are localized in the mitochondrial matrix.
Supplemental Figure S2. Quantification of NFU5 protein levels from the nfu5-2 allele.
Supplemental Figure S3. Root elongation in seedlings grown on vertical plates.
Supplemental Figure S4. Characterization of the isu1-1 mutant allele alone and combined with nfu4-2.
Supplemental Figure S5. Protein blot and BN-PAGE analysis of mitochondria from nfu4 and nfu5 single mutants.
Supplemental Figure S6. Growth of nfu4-2 nfu5 mutants under low CO2.
Supplemental Figure S7. Elevated CO2 cannot rescue growth of nfu4 nfu5 mutants to produce seeds.
Supplemental Figure S8. Protein blot analysis of lipoic-labeled proteins.
Supplemental Figure S9. BiFC to test interaction between LIP1 and NFU4/5, continued.
Supplemental Figure S10. Lipoic acid in growth medium does not rescue the nfu4 nfu5 mutant phenotype.
Supplemental Table S1. isu1, nfu4, and nfu5 T-DNA insertion lines used in this study.
Supplemental Table S2. Free, water-soluble amino acid concentrations in nfu4-2 nfu5-1 seedlings and WT-like segregants.
Supplemental Table S3. Organic acid concentrations in WT, nfu4-2 nfu5-1 and WT-like segregants.
Supplemental Table S4. Primers used in this study.

 Acknowledgments
We thank Brigitte Touraine, Cyril Magno, and Frédéric Gaymard (INRA Montpellier) for mutant selection, seed propagation, and initial phenotyping; Delphine Bernard (Department of Plant Sciences, University of Cambridge) for initial studies on the isu1-1 and nfu4-2 mutants; Rob Green...
and Lucy Anderson (John Innes Centre, JIC) for technical assistance; Markus Dräger (JIC) for root growth measurements and cloning of the ABI3:NFU4 construct; Baldekar Kular (JIC Metabolomics Platform) for amino acid analysis; and the Montpellier PHIV Platform for assistance with confocal microscopy.

Funding

This work was supported by a studentship from the John Innes Foundation to A.E.M.; the Biotechnology and Biological Sciences Research Council, grant awards BB/P012523/1 and BB/P012574/1 to M.F. and J.B.; funding from the T4F Trees for the Future and the Kempe Foundations (GÖF) to O.K. and D.L.; the French National Research Agency, grant awards ANR-13-BSV6-0002-01, ANR-11-LABX-0002-01 (Lab of Excellence ARBRE) and ANR-15-IDEX-04-LUE (Lorraine Université d’Excellence) to N.R. and J.P.-T.

Conflict of interest statement

There are no conflicts of interest for the work reported here.

References

Angelini S, Gerez C, Choudens SO De, Sanakis Y, Fontecave M, Barras F, Py B (2008) NfuA, a new factor required for maturing Fe/S proteins in Escherichia coli under oxidative stress and iron starvation conditions. J Biol Chem 283: 14084–14091.

Azam T, Przybyla-Toscano J, Vignols F, Couturier J, Rouhier N, Johnson MK (2020a) [4Fe–4S] cluster trafficking mediated by Arabidopsis mitochondrial ISCA and NFU proteins. J Biol Chem 295: 18367–18378.

Azam T, Przybyla-Toscano J, Vignols F, Couturier J, Rouhier N, Johnson MK (2020b) The Arabidopsis mitochondrial glutaredoxin GRXS15 provides [2Fe–2S] clusters for ISCA-mediated [4Fe–4S] cluster maturation. Int J Mol Sci 21: 9237.

Balk J, Schaedler TA (2014) Iron cofactor assembly in plants. Annu Rev Plant Biol 65: 125–153.

Bao X, Focke M, Pollard M, Ohrogge J (2000) Understanding in vivo carbon precursor supply for fatty acid synthesis in leaf tissue. Plant J 22: 39–50.

Berger N, Vignols F, Przybyla-Toscano J, Roland M, Rofidal V, Touraine B, Zienkiewicz K, Couturier J, Feussner I, Santoni V, et al. (2020) Identification of client iron–sulfur proteins of the chloroplastic NFU2 transfer protein in Arabidopsis thaliana. J Exp Bot 71: 4171–4187.

Brancaccio D, Gallo A, Nikolajczyk M, Zovo K, Palumaa P, Novellino E, Piccioni M, Cioffi-Baffoni S, Banci L (2014) Formation of [4Fe–4S] clusters in the mitochondrial iron-sulfur cluster assembly machinery. J Am Chem Soc 136: 162–150.

Cameron JM, Jager A, Levandovskiy V, Mackay N, Rouault TA, Tong WH, Ogilvie I, Shoubridge EA, Robinson BH (2011) Mutations in iron-sulfur cluster scaffold genes NFU1 and BOLA3 cause a fatal deficiency of multiple respiratory chain and 2-oxoacid dehydrogenase enzymes. Am J Hum Genet 89: 486–495.

Carrie C, Kühn K, Murcha M, Duncan O, Small I, O’Toole N, Whelan J (2009) Approaches to defining dual-targeted proteins in Arabidopsis. Plant J 57: 1128–1139.

Cochemé HM, Murphy MP (2008) Complex I is the major site of mitochondrial superoxide production by paraquat. J Biol Chem 283: 1786–1798.

Couturier J, Touraine B, Briat J-F, Gaymard F, Rouhier N (2013) The iron-sulfur cluster assembly machineries in plants: current knowledge and open questions. Front Plant Sci 4: 259.

Cronan JE (2016) Assembly of lipoyc acid on its cognate enzymes: an extraordinary and essential biosynthetic pathway. Microbiol Mol Biol Rev 80: 429–450.

Despres B, Delseny M, Devic M (2001) Partial complementation of embryo defective mutations: a general strategy to elucidate gene function. Plant J 27: 149–159.

Ewald R, Hoffmann C, Florian A, Neuhaus E, Fernie AR, Bauwe H (2014) Lipoyc-protein ligase and octanoyltransferase are essential for protein lipoylation in mitochondria of arabidopsis. Plant Physiol 165: 978–990.

Frazzon A, Ramirez M, Warek U, Balk J, Frazzon J, Dean D, Winkel B (2007) Functional analysis of Arabidopsis genes involved in mitochondrial iron–sulfur cluster assembly. Plant Mol Biol 64: 225–240.

Fu X, Guan X, Garlock R, Nikolaou BJ (2020) Mitochondrial fatty acid synthase utilizes multiple acyl carrier protein isoforms. Plant Physiol 183: 547–557.

Fuchs P, Rugen N, Carrie C, Elsässer M, Finkemeier I, Giese J, Hildebrandt TM, Köhn K, Maurino VG, Ruberti C, et al. (2020) Single organelle function and organization as estimated from Arabidopsis mitochondrial proteomes. Plant J 101: 420–441.

Guan X, Okazaki Y, Lithio A, Li L, Zhao X, Jin H, Nettleton D, Saito K, Nikolaou BJ (2017) Discovery and characterization of the 3-hydroxyacyl-ACP dehydratase component of the plant mitochondrial fatty acid synthase system. Plant Physiol 173: 2010–2028.

Guan X, Okazaki Y, Zhang R, Saito K, Nikolaou BJ (2020) Dual-localized enzymatic components constitute the fatty acid synthase systems in mitochondria and plastids. Plant Physiol 183: 517–529.

Han B, Yang Z, Samma M, Wang R, Shen W (2013) Systematic validation of candidate reference genes for qRT-PCR normalization under iron deficiency in Arabidopsis. BioMetals 26: 403–413.

Hildebrandt TM, Nunes Nesi A, Araújo WL, Braun HP (2015) Amino acid catabolism in plants. Mol Plant 8: 1563–1579.

Ito J, Heazlewood J, Millar A (2006) Analysis of the soluble ATP-binding proteome of plant mitochondria identifies new proteins and nucleotide triphosphate interactions within the matrix. J Proteome Res 5: 3459–3469.

Ke J, Behal R, Back S, Nikolaou B, Wuurtele E, Oliver D (2000) The role of pyruvate dehydrogenase and acetyl-coenzyme A synthetase in fatty acid synthesis in developing Arabidopsis seeds. Plant Physiol 123: 497.

Lee CP, Eubel H, O’Toole N, Millar AH (2008) Heterogeneity of the mitochondrial proteome for photosynthetic and non-photosynthetic Arabidopsis metabolism. Mol Cell Proteomics 7: 1297–1316.

León G, Holiugue L, Jordana X (2007) Mitochondrial complex II is essential for gametophyte development in Arabidopsis. Plant Physiol 143: 1534–1546.

León S, Touraine B, Briat JF, Lobréaux S (2005) Mitochondrial localization of Arabidopsis thaliana lue Fe–S scaffold proteins. FEBS Lett 579: 1930–1934.

León S, Touraine B, Ribot C, Briat JF, Lobréaux S (2003) Iron–sulfur cluster assembly in plants: Distinct NFU proteins in mitochondria and plastids from Arabidopsis thaliana. Biochem J 371: 823–830.

Lill R, Freibert S (2020) Mechanisms of mitochondrial iron–sulfur protein biogenesis. Annu Rev Biochem 89: 471–499.

Marquis B, Louks H, Bose C, Wolfe R, Singh S (2017) A new derivatization reagent for HPLC-MS analysis of biological organic acids. Chromatographia 80: 1723–1732.

May M, Leaver C (1993) Oxidative stimulation of glutathione synthesis in Arabidopsis thaliana suspension cultures. Plant Physiol 103: 621–627.

Mayr JA, Feichtinger RG, Tort F, Ribes A, Sperl W (2014) Lipoic acid biosynthesis defects. J Inherit Metab Dis 37: 553–563.

McCarthy EL, Booker SJ (2017) Destruction and reformation of an iron–sulfur cluster during catalysis by lipoyl synthase. Science 358: 373–377.
Mehler A, Na U, Vashisht A, Weiler BD, Lill R, Wohlschlegel JA, Winge DR (2016) Role of Nfu1 and Bo3 in iron-sulfur cluster transfer to mitochondrial clients. Elife 5: 1–24

Meyer E, Solheim C, Tantz S, Bonnard G, Millar A (2011) Insights into the composition and assembly of the membrane arm of plant complex I through analysis of subcomplexes in Arabidopsis mutant lines. J Biol Chem 286: 26081–26092

Moseler A, Aller I, Wagner S, Nietzel T, Przybyla-Toscano J, Mühlenhoff U, Lill R, Berndt C, Rouhier N, Schwarzländer M, et al. (2015) The mitochondrial monothiol glutaredoxin S15 is essential for iron–sulfur protein maturation in Arabidopsis thaliana. Proc Natl Acad Sci USA 112: 13735–13740

Moseler A, Kruse I, Maclean AE, Pedroletti L, Franceschetti M, Wagner S, Wehler R, Fischer-Schrader K, Poschet G, Wirtz M, et al. (2021) The function of glutaredoxin GRXS15 is required for lipoil-dependent dehydrogenases in mitochondria. Plant Physiol 186: 1507–1525

Mühlenhoff U, Lill R (2000) Biogenesis of iron–sulfur proteins in eukaryotes: A novel task of mitochondria that is inherited from bacteria. Biochim Biophys Acta Bioenerg 1459: 370–382

Navarro-Sastre A, Tort F, Stehling O, Uzarska MA, Arranz JA, Del Toro M, Labayru MT, Landa J, Font A, Garcia-Villoria J, et al. (2011) A fatal mitochondrial disease is associated with defective NFU1 function in the maturation of a subset of mitochondrial Fe–S proteins. Am J Hum Genet 89: 656–667

Peterhansel C, Horst I, Niessen M, Blume C, Kebeish R, Kürkciöglu S, Kreuzaler F (2010) Photorespiration. The Arabidopsis Book e0130

Przybyla-Toscano J, Boussardon C, Law S, Rouhier N, Keech O (2021a) Gene atlas of iron-containing proteins in Arabidopsis thaliana. Plant J 106: 258–274

Przybyla-Toscano J, Christ L, Keech O, Rouhier N (2021b) Iron–sulfur proteins in plant mitochondria: Roles and maturation. J Exp Bot 72: 2014–2044

Py B, Gerez C, Angelini S, Planal R, Vinella D, Liseau L, Talla E, Brochier-Armanet C, Garcia Serres R, Latour J, et al. (2012) Molecular organization, biochemical function, cellular role and evolution of NfuA, an atypical Fe–S carrier. Mol Genet Genomics 86: 155–171

Reichel C, Mathur J, Eckes P, Langenkemper K, Koncz C, Schell J, Reiss B, Maas C (1996) Enhanced green fluorescence by the exogenous expression of an Aequorea victoria green fluorescent protein mutant in mono- and dicotyledonous plant cells. Proc Natl Acad Sci USA 93: 5888–5893

Reinholt O, Schwab S, Zhang Y, Reichheld J, Fernie A, Hagemann M, Timm S (2021) Redox-regulation of photosorption through mitochondrial thioredoxin o1. Plant Physiol 181: 442–457

Rohde A, Van Montagu M, Boerjan W (1999) The ABSCISIC ACID-SENSITIVE 3 (AB13) gene is expressed during vegetative quiescence processes in Arabidopsis. Plant Cell Environ 22: 261

Roland M, Przybyla-Toscano J, Vignols F, Berger N, Azam T, Christ L, Santoni V, Wu HC, Dallestein T, Johnson MK, et al. (2020) The plastidial Arabidopsis thaliana NFU1 protein binds and delivers [4Fe–4S] clusters to specific client proteins. J Biol Chem 295: 1727–1742

Schilke B, Voisine C, Beinert H, Craig E (1999) Evidence for a conserved system for iron metabolism in the mitochondria of Saccharomyces cerevisiae. Proc Natl Acad Sci USA 96: 10206–10211

Ströher E, Grasli J, Carrie C, Fenske R, Whelan J, Millar AH (2016) Glutaredoxin S15 is involved in Fe–S cluster transfer in mitochondria influencing lipoic acid-dependent enzymes, plant growth, and arsenic tolerance in Arabidopsis. Plant Physiol 170: 1284–1299

Sulo P, Martin N (1993) Isolation and characterization of LIPS. A lipase biosynthesis locus of Saccharomyces cerevisiae. J Biol Chem 268: 17634–17639

Sweetlove LJ, Taylor NL, Leaver CJ (2007) Isolation of intact, functional mitochondria from the model plant Arabidopsis thaliana. Methods Mol Biol 372: 125–36

Tao Y, O’Toole N, Taylor N, Millar A (2010) Divalent metal ions in plant mitochondria and their role in interactions with proteins and oxidative stress-induced damage to respiratory function. Plant Physiol 152: 7477–61

Timm S, Florian A, Arrivault S, Stitt M, Fernie A, Bauwe H (2012) Glycine decarboxylase controls photosynthesis and plant growth. FEBS Lett 586: 3692–3697

Touraine B, Boutil J, Marion-Poll A, Briat J, Peltier G, Lobréaux S (2004) Nfu2: a scaffold protein required for [4Fe–4S] and ferredoxin iron–sulphur cluster assembly in Arabidopsis chloroplasts. Plant J 40: 101–111

Touraine B, Vignols F, Przybyla-Toscano J, Ischebeck T, Dhalleine T, Wu HC, Magno C, Berger N, Couturier J, Dubs C, et al. (2019) Iron–sulfur protein NFU2 is required for branched-chain amino acid synthesis in Arabidopsis roots. J Exp Bot 70: 1875–1889

Uzarska MA, Przybyla-Toscano J, Spantgar F, Zannini F, Lill R, Mühlenhoff U, Rouhier N (2018) Conserved functions of Arabidopsis mitochondrial late-acting maturation factors in the trafficking of mitochondrial FeS clusters. Biochim Biophys Acta Mol Cell Res 1865: 1250

Vignols F, Bréhelin C, Surdin-Kerjan Y, Thomas D, Meyer Y (2005) A yeast two-hybrid knockdown strain to explore thioredoxin-interacting proteins in vivo. Proc Natl Acad Sci USA 102: 16729–16734

Walter M, Chaban C, Schütze K, Batistic O, Weckermann K, Nägele C, Blazevic D, Grefen C, Schumacher K, Oecking C, et al. (2004) Visualization of protein interactions in living plant cells using bimolecular fluorescence complementation. Plant J 40: 428–438

Wang L, He S, Zhai H, Liu D, Wang Y, Liu Q (2013) Molecular cloning and functional characterization of a salt tolerance-associated gene IbNFU1 from sweet potato. J Integr Agric 12: 27–35

Weiler B, Brück M, Kothe I, Bill E, Lill R, Mühlenhoff U (2020) Mitochondrial [4Fe–4S] protein assembly involves reductive [2Fe–2S] cluster fusion on ISCA1–ISCA2 by electron flow from ferredoxin FDX2. Proc Natl Acad Sci USA 117: 20555–20565

Winter H, Lohaus G, Heldt H (1992) Phloem transport of amino acids in relation to their cytosolic levels in barley leaves. Plant Physiol 99: 996–1004

Wydro MM, Sharma P, Foster JM, Borch K, Meyer EH, Balick J (2013) The evolutionarily conserved iron–sulfur protein INDH1 is required for complex I assembly and mitochondrial translation in Arabidopsis. Plant Cell 25: 4014–4027

Yabe T, Morimoto K, Kikuchi S, Nishio K, Terashima I, Nakai M (2004) The Arabidopsis chloroplastic Nfu-like protein Cnfu, which can act as an iron–sulfur cluster scaffold protein, is required for biogenesis of ferredoxin and photosystem I. Plant Cell 16: 993–1007

Yoo SD, Cho YH, Sheen J (2007) Arabidopsis mesophyll protoplasts: a versatile cell system for transient gene expression analysis. Nat Protoc 2: 1565–1572

Yuvanisamy P, Agar JN, Cash VL, Johnson MK, Dean DR (2000) NiFS-directed assembly of a transient [2Fe–2S] cluster within the NiF protein. Proc Natl Acad Sci USA 97: 599–604

Zannini F, Roret T, Przybyla-Toscano J, Dhalleine T, Rouhier N, Couturier J (2018) Mitochondrial Arabidopsis thaliana TRXo isoforms bind an iron-sulfur cluster and reduce NFU proteins in vitro. Antioxidants (Basel) 1: 142