Dual Location of the Mitochondrial Preprotein Transporters B14.7 and Tim23-2 in Complex I and the TIM17:23 Complex in Arabidopsis Links Mitochondrial Activity and Biogenesis

Yan Wang, a Chris Carrie, a Estelle Giraud, a Dina Elhafez, a Reena Narsai, a,b Owen Duncan, a James Whelan, a and Monika W. Murcha a,1

aAustralian Research Council Centre of Excellence in Plant Energy Biology, University of Western Australia, Crawley 6009, Western Australia, Australia
bCentre for Computational Systems Biology, University of Western Australia, Crawley 6009, Western Australia, Australia

Interactions between the respiratory chain and protein import complexes have been previously reported in Saccharomyces cerevisiae, but the biological significance of such interactions remains unknown. Characterization of two mitochondrial preprotein and amino acid transport proteins from Arabidopsis thaliana, NADH dehydrogenase B14.7 like [B14.7 encoded by At2g42210] and Translocase of the inner membrane subunit 23-2 [Tim23-2 encoded by At1g72750], revealed both proteins are present in respiratory chain complex I and the Translocacon of the Inner Membrane 17:23. Whereas depletion of B14.7 by T-DNA insertion is lethal, Tim23-2 can be depleted without lethality. Subtle overexpression of Tim23-2 results in a severe delayed growth phenotype and revealed an unexpected, inverse relationship between the abundance of Tim23-2 and the abundance of respiratory complex I. This newly discovered relationship between protein import and respiratory function was confirmed through the investigation of independent complex I knockout mutants, which were found to have correspondingly increased levels of Tim23-2. This increase in Tim23-2 was also associated with delayed growth phenotypes, increased abundance of other import components, and an increased capacity for mitochondrial protein import. Analysis of the Tim23-2–overexpressing plants through global quantitation of transcript abundance and in-organelle protein synthesis assays revealed widespread alterations in transcript abundance of genes encoding mitochondrial proteins and altered rates of mitochondrial protein translation, indicating a pivotal relationship between the machinery of mitochondrial biogenesis and mitochondrial function.

INTRODUCTION

A variety of multisubunit protein complexes are present on the mitochondrial outer membrane (Translocase of the Outer Membrane [TOM] and Sorting and Assembly Machinery) (Stroud et al., 2010) and on the inner membrane (Translocase of the Inner Membrane 17:23 [TIM17:23] and TIM22) (Rehling et al., 2004; Mokranjac and Neupert, 2010), which act together to import the wide variety of mitochondrial proteins synthesized in the cytoplasm. Whereas the protein import apparatus is generally well conserved across phylogenetic groups (Hewitt et al., 2011), particularly the pore-forming subunits, which display high levels of orthology (Dolezal et al., 2006; Carrie et al., 2010a), significant differences have been observed between the outer mitochondrial membrane protein import receptors of plants compared with yeast and mammals (Perry et al., 2006; Lister et al., 2007). Further differences include the location of the mitochondrial processing peptidase (MPP) in the cytochrome bc 1 complex (Braun et al., 1992; Glaser et al., 1994) in plants compared with yeast and mammals. Similarly, protein import complexes have also been reported in plants through global quantitation of transcript abundance and in-organelle protein synthesis assays revealed widespread alterations in transcript abundance of genes encoding mitochondrial proteins and altered rates of mitochondrial protein translation, indicating a pivotal relationship between the machinery of mitochondrial biogenesis and mitochondrial function.

1 Address correspondence to monika.murcha@uwa.edu.au.
The author responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (www.plantcell.org) is: Monika W. Murcha (monika.murcha@uwa.edu.au).

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Figure 1. Overview of the PRAT Proteins in Arabidopsis.

(A) Diagrammatic representation of the 10 PRAT proteins previously shown to have a mitochondrial localization. The complex that these proteins have been identified in is indicated according to previous studies (Meyer et al., 2008; Klodmann et al., 2010, 2011). aa, amino acids.

(B) Comparison of sequences of Tim23-2 and B14.7 from Arabidopsis and Tim23 from Saccharomyces cerevisiae (Sc). The predicted transmembrane domains are highlighted, and the PRAT domain (G/A)(X2)F/Y)(X10RX3DX6(G/A/S))GX3(G) is indicated. Percentage identity (in gray) and similarity between Tim23-2, B14.7, and ScTim23.
Figure 2. Interaction between TIM17:23 and Complex I.
The PAM complex is located on the matrix side of the inner membrane and pulls precursor proteins through the TIM17:23translocate into the matrix or inner membrane. Tim21, Tim23, and Tim50 play a role in facilitating interactions with the outer membrane TOM complex to form a dynamic TOM-TIM supercomplex (van der Laan et al., 2006; Chacinskas et al., 2010). In fact, two distinct forms of TIM17:23 have been identified, TIM17:23motor and TIM17:23sortor (Schmidt et al., 2010). TIM17:23sortor contains Tim17, 23, 50, and 21 and is involved in the lateral insertion of preproteins into the inner membrane (Schmidt et al., 2010). TIM17:23motor translocates proteins into the matrix and does not contain Tim21 but does contain the PAM subunits Tim44, HSP70, Mge1, and Pam 16, 17, and 18 (Schmidt et al., 2010).

There are also a number of reports of respiratory chain components and protein import complexes interacting or converging via two types of interactions. First, protein subunits in a respiratory chain complex have been reported to be involved in protein import, this was initially reported with the finding that a subunit of the matrix-located MPP (β-MPP) also functions as one of the core subunits of the cytochrome bc₁ complex in Neurospora crassa (Schulte et al., 1989). Further studies in plants revealed a unique evolutionary history for MPP in mitochondria (Braun et al., 1992; Glaser et al., 1994; Eriksson et al., 1996; Brumme et al., 1998; Glaser and Dessi, 1999), where the core subunits of plant cytochrome bc₁ complexes are bifunctional, acting as both a peptidase and as integral subunits of the cytochrome bc₁ complex (Braun et al., 1992; Eriksson et al., 1996). More recently, it has been reported that the succinate dehydrogenase subunit 3 in yeast is also a component of the Tim22 translocase and has an active role in the import of proteins via the carrier import pathway (Gebert et al., 2011). The second type of interaction is where the multisubunit protein complexes interact to form supercomplexes; for example, current evidence indicates that Tim21 mediates the interaction between the TIM17:23 and complex III in yeast (van der Laan et al., 2006).

Arabidopsis thaliana contains 17 genes encoding proteins that belong to the Preprotein and Amino acid Transport family (PRAT) (Rassow et al., 1999), of which 10 proteins are located in mitochondria, six are located in plastids, and one protein is dual targeted to mitochondria and plastids (Murcha et al., 2007). Of the 10 mitochondrial PRAT proteins, phylogenetic analysis suggested that three genes encode Tim17, three genes encode Tim23, two identical genes encode Tim22, and three genes encode proteins of unknown function (Murcha et al., 2007). One of these genes (At2g42210) encoding a protein of unknown function was subsequently shown to encode the complex I subunit B14.7 (Meyer et al., 2008; Klodmann et al., 2010; Klodmann et al., 2011). Complex I from N. crassa, humans, and Chlamydomonas reinhardtii have also been shown to contain a subunit that is similar to the PRAT family proteins, namely, N. crassa 21.3b (GenBank/P25710), human NDUFA11, and C. reinhardtii v2.0/C_180167 (GenBank/AAS58499) (Nehls et al., 1991; Carroll et al., 2002; Murray et al., 2003). In Arabidopsis, this protein has been identified to be a single peripheral subunit of complex I in two independent studies, both of which identified B14.7 in stoichiometric amounts by proteomic analysis (Meyer et al., 2008; Klodmann et al., 2010).

To investigate the extent and implications of any interactions between the respiratory chain and protein import complexes on the inner mitochondrial membrane, the locations and the interactions of various components of the TIM17:23 complex in Arabidopsis were investigated. The biological implications of interactions between respiratory chain complexes and the mitochondrial protein import machinery are unclear in plants. While yeast is an excellent simple eukaryotic model system, it differs from plants in many aspects with reference to mitochondrial biology, most notably that it lacks a complex I. In this study, we describe an interaction between TIM17:23 and respiratory complex I. Furthermore, the stoichiometric amounts of these two complexes are linked in vivo and may provide a link between protein import and respiration to regulate mitochondrial biogenesis in plants.

**RESULTS**

**Interaction between Tim23 and Complex I**

Of the 10 mitochondrial PRAT proteins (Murcha et al., 2007), two have been reported to be located in complex I of the respiratory chain (Figure 1A). Tim22 was shown to migrate with complex I,
Figure 3. Overexpression of Tim23-2 Results in a Retarded Growth Phenotype.

(A) Growth phenotypes for Arabidopsis plants that contain T-DNA insertions annotated to be located in Tim23-2 (At1g72750) at 2 weeks old.
The complex I subunit B14.7 (Meyer et al., 2008; Klodmann et al., 2010), migrates with both complex I and the TIM17:23 complex (Klodmann et al., 2011; Rode et al., 2011). A comparison of B14.7, Tim23-2, and yeast Tim23 shows that B14.7 is similar to yeast Tim23 in terms of overall characteristics and contains an Arg residue in the correct position of the PRAT consensus domain, G/AXF/YX(−2)RX(−6)G/A/SGX/G, compared with Tim23-2 (Figure 1B). It has been shown that this Arg residue must be inserted into Tim23-2 from Arabidopsis before it can complement a tim23 yeast deletion strain (Murcha et al., 2003). To confirm that B14.7 was present in both complex I and TIM17:23, a variety of assays were used both to test for interactions of B14.7 with TIM17:23 and to determine whether B14.7 was a subunit of TIM17:23.

Proteins of the TIM17:23 complex from Arabidopsis were used to probe for protein–protein interactions using yeast two-hybrid assays. Tim23-2 was tested for interactions against various components of the TIM17:23 complex and complex I and complex III subunits (Figure 2A). Proteins known to interact with Tim23-2, such as Tim23-2 itself, Tim17-2, and Tim50, displayed positive interactions, evidenced by growth on −Leu −Trp −Ade −His with X-α-D-galactoside and a blue phenotype (Figure 2A). Tim23-2 was found to interact with the complex III subunit, MPPx, and the complex I subunit, B14.7 (Figure 2A) (Meyer et al., 2008; Klodmann et al., 2010). The empty vector alone exhibited growth only on −Leu −Trp media, confirming that the growth observed on selection media was due to specific interactions of Tim23-2 and (Tim17-2) with partner proteins.

Whereas the yeast two-hybrid assays showed that B14.7 can interact with Tim23-2 (Figure 2A), this does not directly indicate that B14.7 is a subunit of TIM17:23 or confirm the previous report that B14.7 comigrates with TIM17:23 (Klodmann et al., 2011; Rode et al., 2011). This was determined in two ways: by probing protein complexes separated by Blue-Native PAGE (BN-PAGE) with a variety of antibodies (Figure 2B) and by analyzing the ability of radiolabeled proteins to be incorporated into protein complexes (Figure 2C). Antibodies raised against Tim23-2 detected two protein bands with apparent molecular masses of −100 and −200 kD (Figure 2B). In addition to several non-specific interactions (indicated by an asterisk), a band was consistently detected at −720 kD, which represents the mobility of monomeric complex I. Probing with Tim17-2 antibody detected the single TIM17:23 complex at −200 kD, and no cross-reaction was observed at −720 kD, which corresponds with the location of complex I (Figure 2B). The antibody raised against B14.7 similarly cross-reacted with the Tim17:23 complex at −100 and −200 kD in addition to the monomeric complex I (Figure 2B). The specificity of the antibody to B14.7 was determined against recombinant full-length Tim23-1, Tim23-2, and Tim23-3 proteins, and no cross-reactivity was observed (see Supplemental Figure 1 online). An antibody raised against Tim50 also detected the upper band of Tim17:23 complex, but not complex I (Figure 2B). To confirm further the dual locations of both Tim23-2 and B14.7, in vitro protein uptake assays of radiolabeled proteins were performed and analyzed by BN-PAGE. The uptake of radiolabeled Tim23-2 into the Tim17:23 complexes was observed as expected, but uptake of Tim23-2 into the monomeric complex I was also distinctly observed, in a manner that increased over time (Figure 2C). In the case of Tim17-2, whereas import into the TIM17:23 complexes was similarly observed, there was no incorporation of radiolabeled protein into complex I (Figure 2C). For B14.7, uptake was observed into the TIM17:23 complex and complex I in a time-dependant manner, whereas another complex I subunit, encoded by At1g47260, was shown to incorporate into both monomeric complex I and the supercomplex I+III (Figure 2C). Incorporation of B14.7 was also observed into an unknown complex, located between complex III and complex IV (Figure 2C, indicated with a +). In addition, nonspecific labeling of complex V was observed with all precursor proteins (indicated by an asterisk), which likely represents the incorporation of radiolabeled Met into subunit 9 and the α-subunit of the ATP synthase complex that are encoded in the mitochondrion. While unlabeled Met is added prior to the import assay, it is not unexpected that some incorporation can still occur (Carrie et al., 2010b). These results suggest that both Tim23-2 and B14.7 are imported into and accumulate in both complex I and the TIM17:23 complex.

The interaction of Tim23-2 with complex I was further confirmed by immunoprecipitation and bimolecular fluorescence complementation (BiFC) (Figures 2D and 2E). Mitochondria isolated from Columbia-0 (Col-0) plants were lysed and immunoprecipitation was performed with antibodies raised against Tim23-2. In addition to Tim23-2 and B14.7 being detected in the precipitated products (Figure 2D), the complex I subunit Ndufs4 and complex III subunit Rieske Iron-Sulfur protein (RISP) were also detected, while cytochrome c was not, indicating that antibodies to Tim23-2 can specifically pull down complex I and complex III subunits. Using a split yellow fluorescent protein BiFC assay, the interaction of Tim23-2 with the B14.7 was confirmed and found to be mitochondrial in location by biolistic transformation in onion (Allium cepa) epidermal cells (Figure 2E).

Finally, it has been reported previously that in yeast, Tim21 interacts with complex III (van der Laan et al., 2006). Using a similar set of assays, interaction between Tim21 (encoded by At4g00026) and the cytochrome bc1 complex was revealed in Arabidopsis (see Supplemental Figure 2 online). However, as yeast mitochondria lack complex I, and instead use a type II NADH dehydrogenase (Melo et al., 2004), this interaction between...
complex I and TIM17:23 has not been previously shown in other systems. Therefore, we further investigated this interaction using genetic approaches to determine the biological significance.

Genetic Characterization of the Interactions between Tim17-2, Tim23-2, and B14.7

To characterize the interactions between Tim17-2, Tim23-2, and B14.7, T-DNA insertion lines were screened for each protein. In the case of Tim17-2 and B14.7, no homozygous lines could be obtained, indicating these are essential proteins in Arabidopsis. As tim17 is an essential gene in yeast, it is not unexpected that such lethality was also observed in Arabidopsis (Dekker et al., 1993; Maarse et al., 1994). Interestingly, previous studies have characterized complex I-deficient mutants from tobacco (Nicotiana tabacum) and Arabidopsis and revealed that these plants are viable but exhibit a retarded growth phenotype (Dutilleul et al., 2003; Meyer et al., 2009). Therefore, the lethality of the knockout for B14.7 suggests that it may have other essential roles, in addition to being a complex I subunit.

Two viable T-DNA insertion lines (SALK_143656 and GABI_689C11) were obtained for Tim23-2 (Figure 3A). However, while one line displayed an apparently normal growth phenotype (GABI_689C11), the other line (SALK_143656) displayed a severe delayed growth phenotype throughout the life cycle (Figure 3A). This apparent discrepancy was resolved by mapping and sequencing of the T-DNA insertion in both lines, which revealed that whereas one line (GABI_689C11) represented an insertion in the gene at position 43 bp 3' (downstream) of the translational start site, the insertion for the other line (SALK_143656) was in fact located 135 bp 5' (upstream) of the translational start site (Figure 3B). To determine the effect of both insertions on the abundance of Tim23-2 protein, antibodies raised against Tim23-2 were used in quantitative immunoblot analysis of mitochondria isolated from each line (Figure 3C). In SALK_143656, in which the T-DNA was inserted 135 bp upstream of the translation start site and which exhibited a retarded growth phenotype (Figures 3A and 3B), there was a significant increase (ranging from 1.7- to 2.25-fold; P < 0.05) in abundance of Tim23-2 protein (Figure 3C). By contrast, the same Tim23-2 protein quantitation in the T-DNA insertion line GABI_689C11, where the T-DNA insert is located at a position 43 bp 5' of the gene, resulted in no Tim23-2 protein detected with an apparent molecular mass of 20 kD; instead, a smaller unique band of 18 kD was detected (indicated by an asterisk, Figure 3C). Thus, these lines are referred to as knockout (tim23-2 KO) for GABI_689C11 and overexpressor (tim23-2 OE) for SALK_143656.

The specificity of the Tim23-2 antibody, which was raised against amino acid regions 3 to 32 and 103 to 143, was verified using recombinant full-length overexpressed protein for all three Tim23 isoforms (see Supplemental Figure 3A online). The Tim23-2 antibody cross-reacted against both recombinant Tim23-1 and Tim23-2. Thus, it was not possible to distinguish between Tim23-1 and Tim23-2 using the Tim23-2 antibody alone. An additional antibody raised against Tim23-3 (amino acids 6 to 32 and 64 to 132) detected both recombinant Tim23-1 and Tim23-3 proteins but not Tim23-2 (see Supplemental Figure 3A online). Using the Tim23-3 antibody, no signal was detected in immunoblot analysis of mitochondria isolated from Col-0 plants or any of the Tim23 T-DNA lines (see Supplemental Figure 3A online), implying that Tim23-1 and Tim23-3 are not present in detectable amounts in mitochondria under the conditions tested in this study.

Notably, probing mitochondria isolated from the tim23-2 KO line with the Tim23-2 antibody detected a weak band, corresponding...
Figure 5. Growth Stage Analysis of *tim23-2 OE, tim23-2 KO, Complex I Mutant Lines, rug, ndufs4, and rpotmp and tom20 Triple Mutant.
to a smaller protein of ~18 kD in size, which is not evident in Col-0 or tim23-2 OE blots (Figure 3C). Analysis of the transcript abundance for Tim23-2 revealed that for the tim23-2 KO line, the T-DNA insertion at position 43 bp results in elevated amounts of Tim23-2 transcript downstream of the T-DNA insert. This is in agreement with the detection of a smaller protein likely to represent a truncated form of Tim23-2, translated downstream of the T-DNA insert (see Supplemental Figure 3B online). In the case of the tim23-2 OE line, while Tim23-2 transcript abundance was not elevated, and even somewhat reduced (see Supplemental Figure 3B online), protein abundance was increased (Figure 3C). Previous studies in rice (Oryza sativa) and Arabidopsis have reported that changes in the protein abundances of components of the mitochondrial protein import apparatus are generally not accompanied by changes in the respective transcript abundances. This was observed during germination in rice (Howell et al., 2006) and between aerobically and anaerobically grown rice (Howell et al., 2007). Similarly, in Arabidopsis, the mitochondrial protein import components do not show a correlation between changes in protein and transcript abundance during germination (Law et al., 2012). In addition, the inactivation of any Tom20 or OM64 genes, which encode the receptor components of the TOM complex, has been shown to result in an increase in protein abundance of other isoforms, without any change seen in transcript abundance (Lister et al., 2007). Thus, the amount of Tim23-2 in the tim23-2 OE line appears to be regulated posttranscriptionally.

To investigate further possible reasons for the observed growth defect of tim23-2 OE plants, mitochondria from Col-0, tim23-2 OE, and tim23-2 KO lines were analyzed by BN-PAGE. Analysis of the Coomassie blue–stained gel showed that there was a difference in the banding pattern observed for Col-0, tim23-2 OE, and tim23-2 KO mitochondria (Figure 4). Mitochondria from Col-0 plants showed the characteristic pattern of the respiratory chain complexes, with both the monomeric complexes I and III and the supercomplex I+III evident (Figure 4). The monomeric form of complex I appears to be absent in the tim23-2 OE line, while the supercomplex I+III is greatly reduced in abundance (Figure 4). An additional band, between the supercomplex I+III and complex IV was unchanged when detected with antibodies against the complex IV subunit, COXII. Probing with antibodies against the complex III subunit, RISP, showed a reduction in the supercomplex of complex I+III and the presence of this subunit in the MIC complex (Figure 4). Also, while the intensity of complex III appears to have increased, as determined by the intensity of the band probed with the RISP antibody, this may be explained in part by the absence of the supercomplex I+III. An increase in the abundance of the complex V α-subunit of ATP synthase was also observed in the tim23-2 OE line (Figure 4).

The data above suggests that the growth defect observed within the tim23-2 OE plants occurred as a result of a reduction in the amount of complex I. Thus, we characterized several other mutant lines where complex I activity had been dramatically reduced. Three independent complex I mutants, rug, ndufs4, and rpotmp, were used and growth analysis was performed and compared with tim23-2 OE plants. Additionally, as tim23-2 KO appeared to have no deleterious growth defect, it was also included in the growth analysis. Finally, a triple mutant for the Tom20 outer membrane preprotein receptor that displays significantly reduced rates of protein import (Lister et al., 2007) was also used for comparison. The tim23-2 OE line displayed a similar delayed growth phenotype to the complex I mutants when grown on plates and soil (Figures 5A and 5B), and as expected, the tim23-2 KO plants displayed growth characteristics more similar to Col-0 (Figures 5A and 5B), reaching defined development stages earlier than Col-0 (Figure 5B), with plant height, and maximum rosette radius even observed to be significantly greater than Col-0 (P < 0.05) (Figure 5C). Based on these findings, phenotypically the tim23-2 OE is most similar to complex I mutants.

Overexpression of Tim23-2 and Complex I Mutants Shows Increased Rates of Protein Import

As Tim23 is the channel forming subunit in the TIM17:23 translocase in yeast (Truscott et al., 2001), we investigated whether

Figure 5. (continued).

(A) Plate-based growth progression analysis of Col-0, tim23-2 KO, tom20-2 20-3 20-4 (tom20 triple mutant), tim23-2 OE, rug ndufs4, and rpotmp. Arrows define the time (days after sowing) that Col-0 plants have reached the growth stages as defined by Boyes et al. (2001). Boxes define the time between the growth stages, and shading indicates the occurrence of each growth stage. Stage 0.1, imbibition; stage 0.5, radical emergence; stage 0.7, hypocotyl emerge from seed coat; stage 1.0, cotyledons fully opened; stage 1.02, two rosette leaves > 1 mm in length; stage 1.04, four rosette leaves > 1 mm in length. Data are given as averages for 10 plants. Days are relative to the days after sowing after a 3-d stratification at 4°C

(B) Soil-based growth progression as in (A): stage 1.10, 10 rosette leaves > 1 mm; stage 5.10, first flower buds visible; stage 6.00, first flower opens; stage 6.90, flowering complete.

(C) Representative growth parameters of Col-0 and mutants, number of rosette leaves > 1 mm, maximum rosette radius (mm) over time, plant height (cm) over time, and root length (cm) at 14 d. Data are given as averages ± se for 10 plants. P < 0.05 using a Student’s t test.
Figure 6. Overexpression of Tim23-2 or a Decrease in Complex I Results in Increased Import Ability.
protein import was affected in the tim23-2 KO and OE lines in Arabidopsis. Three proteins that contain N-terminal cleavable targeting signals and are imported via the general import pathway were used including the alternative oxidase (AOX) (Whelan et al., 1995), the F$_{0}$d subunit of ATP synthase (F$_{0}$d) (Smith et al., 1994), and the dual-targeted protein monodehydroascorbate reductase (MDHAR) (Chew et al., 2003). Protein import via the carrier import pathway was also tested using the adenine nucleotide carrier (ANT) and the phosphate carrier (Pic) (Murcha et al., 2005b). The tim23-2 OE line exhibited an increase in the import ability of AOX, F$_{0}$d, and MDHAR, while mitochondria isolated from tim23-2 KO plants displayed a reduced import ability of these three precursor proteins (Figure 6A). The effect of altered Tim23-2 levels was seen to be specific to precursor proteins imported via the general import pathway, as no differences were observed with the import of ANT or Pic (Figure 6A), which are both precursors of the carrier import pathway (Murcha et al., 2005b). Thus, while overexpression of Tim23-2 protein resulted in an increased import ability via the general import pathway, it is not clear why a retarded growth phenotype was observed.

Characterization of the import capacity of mitochondria isolated from the complex I mutants also displayed an increase in protein import (Figures 6B and 6C). The rug mutant line revealed increased import ability of all proteins tested via both the general import pathway (AOX, F$_{0}$d, and MDHAR) and the carrier import pathway (ANT and Pic) (Figure 6B). The latter differs from the tim23-2 OE line, in that there was no increase in the import ability of carrier pathway proteins (Figure 6A, ANT and Pic panels versus Figure 6B, ANT and Pic panels). Similarly, the import ability of the ndufs4 mutant line and to a lesser extent the rpotmp line, exhibited increased import ability of AOX, MDHAR, and Pic precursor proteins when compared with Col-0 (Figure 6C).

**A Reduction in Complex I Results in an Increase in Tim23**

To further investigate the similarities between the tim23-2 OE plants and the complex I mutant plants, mitochondria were isolated from Col-0, tim23-2 KO, tim23-2 OE, rug, ndufs4, and rpotmp plants and probed with a series of antibodies raised against mitochondrial PRAT proteins (Murcha et al., 2007), respiratory chain components, and proteins involved in protein import (Figure 7). Antibody dilutions were optimized for specific concentrations to ensure linearity of detection against Col-0 mitochondria (see Supplemental Figure 5 online). Interestingly, probing mitochondria isolated from these lines, with antibodies to Tim23-2, revealed that all complex I mutants displayed an increase in Tim23-2 protein abundance, similar to that of mitochondria isolated from the tim23-2 OE plants (Figure 7A).

In the case of other mitochondrial PRAT proteins, it was evident that an increase or decrease in the amount of Tim23-2 was accompanied by similar variations in Tim17-2 protein abundance (Figure 7A). For B14.7, a subunit of complex I and a PRAT protein, abundance decreased only slightly in the tim23-2 KO line and increased in the tim23-2 OE and ndufs4 lines. B14.7 protein was not detected in the rug mutant plants and was only barely detected in the rpotmp mutant plants (Figure 7A). This indicates that while a T-DNA insertion into the locus At2g42210 (encoding B14.7) is lethal, even the reduced amounts of the protein that were detected in the rug line and rpotmp lines are sufficient to restore viability. As complex I is still largely absent in these lines, it further supports a dual role for B14.7, one in complex I and a second role in the TIM17-23 complex, that is essential for viability. A PRAT protein of unknown function encoded by At3g25120 showed slight increases in abundances in all lines, including in the tim23-2 KO line (Figure 7A). Tim22, a protein encoded by At3g10110 or At1g18320 (two genes which have 100% identity), which has been shown to complement a yeast tim22 deletion mutant (Murcha et al., 2007), also increased in abundance in the tim23-2 OE and complex I mutant lines (Figure 7A).

Analysis of a variety of protein import components revealed an increase in Tim44 protein abundance in all lines (Figure 7B). In addition, all three Tom20 isoforms increased in abundance in all lines. Increases were also observed in the abundance of Sam50, Tim50, Tim21, and Tim9, most prominently in the tim23-2 KO, OE, and rug lines (Figure 7B).

In the case of a variety of other respiratory chain components, the most notable effect observed was a >80% reduction in Ndufs4 protein abundance in the tim23-2 OE, rug, and rpotmp lines (Figure 7C), supporting the previous report of this for mitochondria from rug lines (Kühn et al., 2011). Ndufs4 is a complex I subunit that is proposed to be present in the peripheral portion of complex I (Klodmann et al., 2010; Klodmann and Braun, 2011). Notably, Nad9, a mitochondrial encoded complex I component, present in the matrix arm, Q module (Klodmann et al., 2010; Klodmann and Braun, 2011) was also slightly decreased in abundance, in the complex I mutants, though it was surprisingly higher in abundance in the tim23-2 OE line (Figure 7C). A variety of other respiratory chain components increased...
Figure 7. The Abundance of a Variety of Proteins as Determined by Immunoblot Analysis of Mitochondria Isolated from Col-0 (Wild Type), tim23-2 OE, tim23-2 KO, and rug, ndufs4, and rpotmp Lines.
in abundance to some extent across the tim23-2 KO, tim23-2 OE, and complex I mutant lines, while the α- and β-subunits of the ATP synthase complex increased in abundance most prominently in the tim23-2 OE, rug, and ndufs4 lines (Figure 7C). COXII protein levels similarly increased in these lines, with barely detectable amounts observed in rpotmp (Figure 7C). The Rieske FeS protein of the cytochrome bc\(_2\) complex showed a similar increase in the tim23-2 OE, rug, and ndufs4 lines (with no increase in the rpotmp line) (Figure 7C). Equal loading of total mitochondrial proteins was confirmed with Coomassie blue staining of mitochondrial samples (see Supplemental Figure 5B online), and no changes were observed in the abundance of porin and TOM40 (Figures 7B and 7C). A Link between Mitochondrial Activity and Biogenesis 2687

**Analysis of Global Transcriptional Changes in Tim23-2 Mutant Plants**

tim23-2 OE lines displayed extensive transcriptional reprogramming, with a total of 1645 transcripts significantly altered with respect to Col-0 seedlings (861 transcripts upregulated and 784 transcripts downregulated >1.5-fold, after false discovery rate [FDR] correction [P < 0.05, posterior probability of differential expression (PPDE) (<P > 0.95)] (Figure 8A). tim23-2 KO lines displayed fewer significant changes, with 525 and 466 transcripts significantly down- and upregulated >1.5-fold, respectively, after FDR correction. Comparison of transcript responses between the tim23-2 OE and the complex I mutant rug lines showed significant overlap, not only in the specific genes altered, but also the direction and magnitude of the responses. Significantly more transcripts were coexpressed between rug and tim23-2 OE than would be expected by random chance, (P < 0.001, \(\chi^2\) test) (indicated by #, Figure 8A). This indicates that the transcriptional responses of rug and tim23-2 OE show significant similarity on a genome-wide scale. Transcript abundance changes for components specifically related to mitochondrial import functions were examined in the two mutant lines. For both lines, transcripts encoding mitochondrial intermembrane space chaperones and factors associated with folding in the matrix were consistently upregulated (see Supplemental Table 1 online; Figure 8B).

To investigate further the transcriptomic changes in the tim23-2 OE mutants in relation to mitochondrial function and regulation, significant differential expression was visualized on a custom MapMan image, displaying diverse facets of mitochondrial function and biogenesis (Figure 8B). Overexpression of Tim23-2, and the associated complex I defect, appears to induce a substantial response within the mitochondrion, including the upregulation of numerous nucleus-encoded and organelle-encoded components of all five cytochrome respiratory pathway complexes (Figure 8B). This is accompanied by an upregulation, at the transcript level, of nuclear-encoded mitochondrial transcription/translation machinery, mitochondrial import components, and a specific upregulation of reactive oxygen species/REDOX scavenging systems within the mitochondria, most likely due to an increase in potential for reactive oxygen species production, due to respiratory chain dysfunction (Figure 7C). Furthermore, steady state mitochondrial encoded transcript levels were measured by quantitative RT-PCR, and a fourfold to 10-fold increase was observed for almost all transcripts encoded by the mitochondrial genome (Figures 8B and 8C). This increase in mitochondrial transcription was also accompanied by an increase in the rates of mitochondrial translation, evident for a number of mitochondrial encoded proteins (Figure 8D). Together, these results suggest that overexpression of Tim23-2 protein results in the instability of mitochondrial respiratory chain complex I and signals a mitochondrial biogenesis response to induce transcription/translation of all components necessary to build a functional organelle.

This potential regulation of Tim23-2 with mitochondrial biogenesis is further supported by genome-wide analysis of the coexpression environment of Tim23-2. The top 50 coexpressed *Arabidopsis* genes with Tim23-2 (based on \(r\) values across >300 public microarray data sets) from the BAR expression Angler and ACT (http://www.Arabidopsis.leeds.ac.uk/act/coexpanalyser.php) were combined, leaving a list of 76 unique genes (see Supplemental Table 2 online). Figure 8D shows a functional categorization of this list, compared with the whole genome, and genes are significantly (P < 0.0001) overrepresented in the functional categories of protein synthesis (15.8% versus 2.6%), development (11.8% versus 2.8%), protein targeting (6.6% versus 1.1%), RNA binding (6.6% versus 0.7%), RNA processing (6.6% versus 1.1%), cell division (2.6% versus 0.4%), and protein folding (2.6% versus 0.3%), supporting a potential role for Tim23-2 during mitochondrial biogenesis.

**DISCUSSION**

The results presented indicate that there is an interaction between complex I of the respiratory chain and TIM17:23 in *Arabidopsis*. It
Figure 8. Global Analysis of Transcript Abundance Changes in tim23-2 OE and KO Lines with Other Mutants Affecting Complex I.
appears that Tim23-2 and B14.7 proteins are both imported into
and accumulate in the respiratory complex I and the TIM17:23
complex. Whereas previous studies in yeast have shown inter-
actions between the respiratory chain and protein import com-
plexes, this study reveals an interaction between the protein
import components and complex I. It is likely that this interaction
was not previously detected simply due to the fact that yeast
lack complex I (Melo et al., 2004). In an attempt to investigate
further the biological significance of Tim23-2 and B14.7 being
present in both complexes, T-DNA insertion lines were screened
for B14.7 and Tim23-2. Two homozygous T-DNA insertion lines
for Tim23-2 were obtained, and despite repeated self-pollination
of heterozygous plants, no homozygous lines for B14.7 could be
obtained. For the Tim23-2 T-DNA insertion lines, one line pro-
duced Tim23-2 OE plants, while the other line produced au-
thentic KO plants.

None of the other Tim23 isoforms (i.e., Tim23-1 or Tim23-3)
were detected in the tim23-2 KO plants, suggesting that the
reason this T-DNA KO was not lethal was because B14.7 could
at least partially fulfill the role of Tim23-2. Note that although the
tim23-2 KO plants were viable, a reduced rate of protein import
via the general import pathway was seen, indicating that B14.7
could not completely complement the function of Tim23-2.
However, a reduced rate of import appears to have no dele-
trious effect on the growth phenotype; in fact, tim23-2 KO plants
even grew slightly better than Col-0 (Figure 5). It is notable that
while Tim23-2 and B14.7 proteins do not display high levels of
sequence identity (20 and 33%, respectively), this is not an in-
dication that complementation cannot occur. For example, while
Tim23 and Tim17 from yeast and plants do not display high
levels of sequence identity, the plant genes for these compo-
ents can still complement yeast deletion strains and restore
viability (Murcha et al., 2007), and B14.7 does have a consensus
PRAT domain, whereas Tim23-2 does not (Rassow et al., 1999).
Finally even low levels of B14.7 are sufficient to maintain vi-
ability, as observed in the rug and rpotmp KO lines, where only
low levels of B14.7 are observed (Figure 7A) (Kühn et al., 2009,
2011). The amount of the TIM23 complex is quite low compared
with respiratory chain complexes, even in yeast, where genetic
approaches had to be used to characterize TIM transciptions,
compared with the more abundant TOM components, which
could be characterized by biochemical approaches (Rehling
et al., 2001).

Given the nature of this tim23-2 OE line in Arabidopsis, it was
not possible to complement this mutant, and previous attempts
to overexpress Tim23-2 using the 35S cauliflower mosaic virus
promoter have proved unsuccessful. Therefore, to verify the
findings that an increase in Tim23-2 protein abundance resulted
in a reduction of complex I, we investigated a number of addi-
tional mutants that also exhibit a reduction in complex I activity,
including the complex I rug mutant, a nad2 splicing mutant
(Kühn et al., 2011), the rpotmp mutant (Kühn et al., 2009), and
the ndufs4 mutant (Meyer et al., 2009). In all cases, a reduction
in complex I activity resulted in significantly increased Tim23-2
protein abundance. Collectively, these results show that the
amounts of complex I and Tim23-2 protein abundance appear to
be linked in Arabidopsis. Although the link between the amount
of complex I and Tim17:23 was unexpected, it is likely that this
occurs given that these two complexes share two proteins.
Therefore, a mutation that affects complex I assembly, but not
the amount of B14.7 directly, may result in additional B14.7
protein being available for assembly into TIM17:23, to replace
Tim23-2. In this case, it is proposed that the other subunits of
the TIM17:23 complex, which were observed to increase in
abundance, compensate to assemble all of the available Tim23-2.
However, in tim23-2 OE plants, it appears that this leads to in-
stability of complex I. The mechanism by which this occurs is
unclear but may be explained by the fact that B14.7 is predicted
to be on the periphery of complex I (Klodmann et al., 2010;
Klodmann and Braun, 2011). Thus, an interaction between
complex I and Tim17:23 and complex II may facilitate the formation of a supercomplex of complex I+III, which is
routinely observed in plant mitochondria (Meyer et al., 2008;
Klodmann et al., 2011). However, in the presence of excess
Tim23-2, the dynamics of this interaction or equilibrium are al-
tered, leading to instability of complex I (Figure 4). Note that in
the tim23 OE plants, a new complex (MIC) is observed, and this
complex is not an assembly intermediate or degradation prod-
uct of complex I, as analysis of this complex reveals that it
contains a variety of proteins (see Supplemental Figure 4 online).

Figure 8. (continued).

(A) Venn diagrams showing the overlap of significantly changing genes at a fold change cutoff of 1.5-fold for rug mutants (Kühn et al., 2011) and tim23-2 OE lines. Red indicates induced transcripts and blue indicates downregulated transcripts.

(B) Overview of significant changes (FDR-corrected P < 0.05) in transcript abundance, in response to overexpression of Tim23-2, for transcripts associated with mitochondrial function and organielle biogenesis. Red boxes indicate transcripts that are induced in tim23-2 OE lines, and blue boxes show transcripts that are downregulated. Color scale is shown as log2 fold change.

(C) Quantitative RT-PCR was performed to compare the abundances of mitochondrial transcripts in tim23-2 OE and KO seedlings compared with Col-0 seedlings. Transcript levels are depicted as log2 of the ratio of values determined in mutant versus Col-0 samples for each transcript on the mito-
chondrial genome. Three technical replicates of three independent biological repeats were performed and averaged per genotype; standard errors are indicated. Nuclear 18S (used as a normalization control) along with ACT2 and UBC transcripts were included as controls.

(D) In organellar translations in mitochondria isolated from Col-0, tim23-2 OE, and KO lines. Equal amounts of mitochondria were used in assays and the incorporation of [15S]Met was monitored in proteins over time.

(E) The 76 genes defined as highly coexpressed with tim23-2 were categorized according to function to determine over- or underrepresented groups compared with the whole genome. Categories shown in bold red and blue indicate significantly (P < 0.0001, χ2 test) over- and underrepresented categories, respectively, compared with the whole genome. Percentages shown in parentheses indicate the relevant percentage observed for a whole genome categorization (see Supplemental Tables 1 and 2 online).
We propose that this MIC complex results from the increased expression of a large variety of mitochondrial proteins that are imported but not assembled into functional complexes. To our knowledge, nonassembled proteins have not previously been detected in chemical amounts in mitochondrial complexes that contain several diverse proteins.

The increased abundance of Tim23-2 protein in the tim23-2 OE line was accompanied by additional changes that together can be grouped into the process of mitochondrial biogenesis. It was notable that transcripts encoding many of the components of the mitochondrial respiratory chain were significantly upregulated in abundance in this study. An analysis of a variety of transcriptome data sets also reveals that these components are stable under a variety of stresses (Clifton et al., 2005) or even in KO s of complex I components (Meyer et al., 2009). Furthermore, there are two additional features of the tim23-2 OE plants that are worth noting: First, a comparison of the protein changes observed with quantitative immunoblotting reveals that the increase in Tom20-2, Tom20-3, and Tom20-4 at a protein level is not accompanied by an increase in the transcript abundance of genes encoding these proteins. This suggests that the protein abundance is regulated at a posttranscriptional level. In a previous study, when any one or two of the Tom20 isoforms in Arabidopsis were inactivated, an increase in the other isoforms was observed, again without any change in transcript abundance (Lister et al., 2007), and consistent with posttranscriptional regulation of protein abundance for these components. Similarly, the increase of Tim23-2 protein abundance in the tim23 OE plants also appears to occur at a posttranscriptional level, supporting the findings from several previous studies indicating that the abundance of mitochondrial protein import components occurs posttranscriptionally (Howell et al., 2006, 2007; Law et al., 2012). The insertion of the T-DNA in the 5′ untranslated region may alter translation of Tim23-2 mRNA, leading to the increase in protein abundance observed in this study. Studies in several plants have revealed a role for the 5′ untranslated region in determining translation efficiency (Kawaguchi and Bailey-Serres, 2005; Wever et al., 2010).

Note that examining genes located in close proximity to Tim23-2 from microarray data revealed that transcript abundance for these genes were largely unaffected. Second, while there is a significant overlap in the transcriptome response between the tim23-2 OE line and the complex I mutant rug line, there are also responses that are unique to both (Figure 8A). The response to both mutations displays common features, such as a large reduction in complex I activity, an increase in Tim23-2 abundance and the abundance of various other proteins of the protein import apparatus, and many common changes in the transcriptome. However, differences were also apparent, including the finding that mitochondria from the complex I rug mutant also displays an increase in the rate of protein imported via the carrier import pathway and transcripts encoding components of the mitochondrial respiratory chain are not upregulated in this mutant. Finally, the overexpression of yeast Tim23 in yeast cells using the galactose promoter to drive expression resulted in no detrimental effects (Emtage and Jensen, 1993; Bauer et al., 1996), unlike what we have seen for plants in this study. It is proposed that the difference between plants (and likely other organisms, such as yeast) may be due to the fact that mitochondrial biogenesis and respiratory chain activity in yeast are not linked, given that yeast is a facultative anaerobic organism. Thus, it is not surprising that the expression of the respiratory chain components and mitochondrial biogenesis are not coupled. Rather, in yeast, the expression of mitochondrial respiratory chain components is under the control of heme-activated proteins that are in turn regulated by the availability of oxygen (Kwast et al., 1998). However, mitochondrial biogenesis still occurs in the absence of oxygen; thus, the respiratory chain and mitochondrial biogenesis are not coupled in yeast. By contrast, respiratory chain activity and mitochondrial biogenesis may be linked in multicellular organisms. Thus, the interaction reported in this study, between complex I and TIM17:23 in Arabidopsis, provides a mechanism to link mitochondrial biogenesis and activity.

In conclusion, the results presented here show that in Arabidopsis, the B14.7 and Tim23-2 proteins are imported into and accumulate in two mitochondrial protein complexes: the respiratory chain complex I and the protein import complex TIM17:23. It is also seen that the abundance of complex I is inversely related to the abundance of Tim23. This may provide for an elegant system to regulate mitochondrial biogenesis, as any changes in complex I, due to oxidative damage or increased respiratory demand, could trigger signals that increase the amount of Tim23-2 protein and thus activate mitochondrial biogenesis.

METHODS

Gene Identification and Phylogenetic Analysis of Proteins

The Mitochondrial Protein Import Components database (Lister et al., 2003) and The Arabidopsis Information Resource (www.Arabidopsis.org) was used to identify genes used in this study. Alignments were performed using ClustalW2 (www.ebi.ac.uk).

Plant Material and Culture Conditions

The SALK T-DNA insertion lines were obtained from the ABRC seed stock center (Alonso et al., 2003), and GABI lines were obtained from GABI-Kat (Rosso et al., 2003) for Tim23-2 (SALK_143656 and GABI_689C11). Lines were screened for homozygosity of the T-DNA insert with primers listed in Supplemental Table 3 online, as previously outlined (Carrie et al., 2010b; Kühn et al., 2011). T-DNA mutants and the wild type were germinated on 0.5 × Gamborg’s B5 salt medium containing 3% (w/v) Suc grown at 22°C with a light intensity of 80 μmol quanta m⁻² s⁻¹ in a 16-h photoperiod.

DNA and PCR

Genomic DNA was isolated from 14-d-old plants, and PCR was performed using the REDExtract-N-Amp Plant PCR kit according to the manufacturer’s instructions. Cloning from cDNA was performed as previously described (Murcha et al., 2003) using primers listed in Supplemental Table 3 online.

Isolation of Mitochondria from Arabidopsis thaliana Plants

For the isolation of mitochondria from water-cultured plants, mutant and Col-0 seeds were sterilized and inoculated in half-strength Murashige and
Skoog media, containing 0.5× Gamborg’s B5 salts, 3% (w/v) Suc, 50 µg/mL cefotaxime, and 2 mM MES KOH, pH 5.7. Plants were grown for 2 weeks as described above and mitochondria isolated as described previously (Lister et al., 2007).

BN-PAGE and Immunodetection of Protein Complexes

BN-PAGE was performed according to Jänsch et al. (1996) as described by Meyer et al. (2009). Following separation, gels were stained with Coomassie Brilliant Blue, subjected to an in-gel complex I activity assay, or transferred to polyvinyl difluoride membrane and immunodetected with a selection of antibodies as described below.

In-Gel Complex I Activity Assay

The assay was performed according to Zerbetto et al. (1997).

Immunoblot Analysis

Mitochondrial protein was separated by SDS-PAGE and transferred to Hybond-C extra nitrocellulose membrane (GE Healthcare). Immunodetection of proteins was performed as described previously (Murcia et al., 2005a). The linearity of the response of the antibodies were tested with 10, 20, and 40 µg of mitochondrial proteins (see Supplemental Figure S5A online) and equal protein loading confirmed by staining with Coomassie Brilliant Blue (see Supplemental Figure S5B online). Furthermore, the CV was calculated for fold changes in abundances of porin and Tom40 to ensure that they were of equal abundance in samples. The CV was determined by calculating the ratio of the standard deviation of the mean, providing a statistical indication of the spread/variation in abundance. Antibodies used were raised against Tim17-2 (Murcha et al., 2005a), Tom20-2, 20-3, 20-4, Metaxin (Lister et al., 2007), Tom40, RISP, Sam50 (Carrie et al., 2010b), AOX, α-subunit of ATP synthase, Porin (Elthon et al., 1989), Nad9 (Lamattina et al., 1993), COXII, cytochrome c, β-subunit of ATP synthase (Agrisera), and Ndufs4 (Meyer et al., 2009). For the remaining antibodies, recombinant proteins of Tim9 (At3g46560, amino acids 1 to 93), Tim50 (At1g55900, amino acids 110 to 269), Tim44 (At2g36670, amino acids 100 to 472), Tim21 (At4g00026, amino acids 160 to 376), Tim23-2 (At1g27570, amino acids 3 to 32 and 103 to 143), Tim23-3 (At3g04800, amino acids 6 to 32 and 64 to 132), At3g25120 (amino acids, amino acids 89 to 189), were expressed and purified as previously outlined (Carrie et al., 2010b). For antibodies raised against B14.7 (At2g42210) (amino acids 1 to 100) and Tim22 (At3g01110/At1g18320 amino acids 1 to 50), cDNA was cloned into pDest15 using Gateway technology, and glutathione-S-transferase–tagged recombinant proteins were expressed, bound to GST-Sepharose (Scientific), and electroeluted. Recombinant proteins were confirmed by mass spectrometry prior to inoculation into rabbits as per the standard protocol (Cooper and Paterson, 2008). Immunodetection of newly synthesized antibodies was performed at a 1 in 5000 dilution against purified mitochondria.

In Vitro Import Studies

Mitochondria for in vitro import experiments were prepared from 14-d-old Arabidopsis seedlings grown in liquid culture, as previously described (Lister et al., 2007). [35S]Met-labeled precursor proteins of AOX (X68702) (Whelan et al., 1995), Fd subunit of ATP synthase (X74296) (Smith et al., 1994), ANT (X57536), Pic (AB016063) (Murcia et al., 2005a), and MDHAR (At3g09940) (Chew et al., 2003) synthesized using the rabbit reticulocyte Tnt1 in vitro transcription/translation lysate (Promega). Import of radio-labeled precursor protein into isolated mitochondria was performed as previously described (Lister et al., 2007) at increasing time points. Following import, mitochondria were pelleted at 20,000g for 5 min and subjected to SDS-PAGE analysis. Gels were stained in Coomassie Brilliant Blue, dried, and exposed to a BAS TR2040 phosphor-imaging plate (Fuji) for 24 h. The exposed plate was visualized using the BAS 2500 Bio-Imaging Analyzer (Fuji). Proteinase K–protected, mature, radiolabeled imported proteins were quantified at each time point and normalized to the highest time point for the wild type, which was set to 1 using Quantity One software (Bio-Rad) (Lister et al., 2007). Three biological replicates were performed for each mutant and precursor protein and the average (and st) of import amount relative to Col-0 was determined.

Imports analyzed by BN-PAGE were performed as previously described (Carrle et al., 2010b). [35S]Met-labeled precursor proteins of Tim17-2 (At2g37410), Tim23-2 (At1g72750) (Murcha et al., 2005a), Tim21 (At4g00026), complex I subunit At1g4726, MPPx (At1g51980) (Kühn et al., 2011), B14.7 complex I subunit At2g42210 (Murcha et al., 2007), and cytochrome c subunit of ubiquinone cytochrome c reductase (At4g32470) were generated from Arabidopsis cDNA cloned into pDest14 using Gateway technology (Invitrogen) and used in import assays. Precursor proteins were synthesized and imported as described previously except that 250 µg of mitochondrial protein was used per time point. Following import, mitochondria were pelleted at 20,000g for 5 min and subjected to BN-PAGE analysis as detailed above. BN-PAGE gels were fixed in 40% (v/v) methanol and 10% (v/v) acetic acid and visualized by autoradiography, as described above.

RNA Isolation and Global Transcript Analysis

Analysis of the changes in transcript abundance between Col-0, tim23-2 OE, and tim23-2 KO lines was performed using Affymetrix GeneChip Arabidopsis ATH1 Genome Arrays (Affymetrix). Whole rosettes from several seedlings at the 8 true leaf stage of development, grown under a 16-h photoperiod, were pooled for each biological replicate. Col-0 and mutant tissue samples were collected in biological triplicate. For each replicate, total RNA was isolated and microarrays were performed as described (Kühn et al., 2011).

Quantitative RT-PCR

RNA, isolated as described above, from Col-0, tim23-2 OE, and tim23-2 KO plants was subjected to two consecutive treatments with TURBO DNase (Ambion), confirmed by PCR to be free of detectable amounts of DNA, and reverse transcribed exactly as described (Kühn et al., 2009). Quantitative RT-PCR on the mitochondrial transcriptome was performed using the LightCycler 480 SYBR Green I Master Mix (Roche Applied Science) using the PCR program and primer pairs and analyzed as described (Kühn et al., 2009).

Statistical Analysis of Microarray Data

Data quality was assessed using GCOS 1.4 before CEL files were exported into AVADIS Prophetic (version 4.3; Strand Genomics) and Partek Genomics Suite software, version 6.3, for further analysis. CEL files are available at Array Express (http://www.ebi.ac.uk/arrayexpress/) under the accession number E-MEXP-3309. Public array data for the rug mutant (Kühn et al., 2011) was also independently analyzed and included. MAS5 normalization algorithms were performed only to generate present/absent calls across the arrays. Probe sets that recorded absent calls in two or more of the three biological replicates for all four genotypes conditions were removed from further analysis. Bacterial controls, probe sets for organelle transcripts and ambiguous probe sets for multiple transcripts were also removed, resulting in a final set of 14,831 probe identifiers for downstream analysis. CEL files were also subjected to GC content background robust multiarray average normalization for computing fluorescence intensity values used in differential expression analyses. Correlation plots were examined between all arrays using the scatterplot
function in the Partek Genomics Suite, and in all cases \( r > 0.98 \) (data not shown). Analysis of differential expression was performed using a regularized \( t \) test based on a Bayesian statistical framework using the software program Cyber-T (Baldi and Long, 2001) (http://cybert.microarray.ics.uci.edu/). Cyber-T employs a mixture model-based method described by Allison et al. (2006) for the computation of the global false-positive and false-negative levels inherent in a DNA microarray experiment. To control for FDR accurately and minimize false positives within the differential expression analysis, PPDE(P) values and PPDE(\( > P \)) values were calculated, as a means to measure the true discovery rate (1 – FDR). Changes in transcript abundance were considered significant with a PPDE(\( > P \)) > 0.95 and a fold change >1.5-fold. Overlaps in the transcript abundance responses for the different genotypes were plotted on Venn diagrams to determine statistically significant over- or underrepresentation in the overlap, compared with that which is expected by random chance, using a Pearson’s \( x^2 \) test for independence. Transcript abundance changes for mitochondrial proteins were visualized in a custom MapMan pathway (Thimm et al., 2004). A nonredundant list of 76 transcripts that are the most highly coexpressed transcripts, genome wide, against Tim23-2 was created from the top 50 gene hits from BAR Expression Angler and The Arabidopsis Coexpression Data Mining Tools (coexpression multi-experiment tool, http://www.Arabidopsis.leeds.ac.uk/act/coexpanalyser.php#CO2) shown in Supplemental Table 2 online (Manfield et al., 2006). Cocorrelation scatterplots were generated using the cocorrelation 2-D Pearson correlation coefficients plot, with the 76 coexpressed top hit transcripts highlighted. Functional categorization analysis was performed using simplified annotation categories defined in MapMan software bins to define over/underrepresented functional groups based on percentile distributions of the whole genome.

**In Organello Protein Synthesis Assays**

Mitochondria were isolated as described above and protein synthesis reactions were performed as previously described (Giegé et al., 2005).

**Yeast Two-Hybrid Protein Interactions**

Yeast two-hybrid interaction assays were performed as described (Giraud et al., 2010). AH109 cells transformed with pGAD Tim23-2 were mated with Y187 cells transformed with pGBK Tim17-2, pGBK Tim23-2, pGBK Tim50, pGBK B14.7, pGBK Tim21, pGBK MPPx, and empty pGBK. Serial dilutions of yeast growing on auxothrophic selection media (\( \text{–Leu} \rightarrow \text{Trp} \), \( \text{–His} \rightarrow \text{Ade} \), and \( \text{–Leu} \rightarrow \text{Trp} \) and positive interaction selection (\( \text{–Trp} \rightarrow \text{Leu} \cdot \text{Ade} \cdot \text{His} \) and \( \text{–Leu} \rightarrow \text{Trp} \rightarrow \text{Ade} \cdot \text{His} \) with X-\( \alpha \)-c-galactoside).

**Immunoprecipitations**

Two hundred micrograms of mitochondria was solubilized in digitonin buffer consisting of 30 mM HEPES KOH, pH 7.4, 150 mM potassium acetate, 10% (v/v) glycerol, and 1 mg digitonin and placed on ice for 20 min. One hundred microliters of Protein A Sepharose with or without 10 \( \mu \)L of antibody was added with solubilization buffer without digitonin, including 0.5% (w/v) BSA and Complete Protease inhibitor (Roche), and incubated for 3 h at 4°C. Beads were washed and samples resolved by SDS-PAGE, transferred to nitrocellulose membrane, and immunodetected with antibodies indicated.

**BiFC**

BiFC was performed as outlined previously (Citovsky et al., 2008) in onion (Allium cepa) epidermal cells. Images were captured using Cell imaging software (Olympus) as previously described (Carrie et al., 2009).

**Blue Native Elution and in-Gel Digestion**

The complex of interest was excised from Blue native gels and proteins extracted by electrophoresis. Gel slices were minced by passing back and forth between two syringes and added to the H-shaped eluter vessel of a CBS Scientific electroluter. Proteins were eluted according to the technique detailed previously (Hunkapiller et al., 1983) Protein identification was performed by in-gel digestion and electrospray ionization tandem mass spectrometry according to previous methods (Meyer et al., 2007).

**Accession Numbers**

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: Tim17-1 (AT1g20350), Tim17-2 (AT2g37410), Tim17-3 (AT5g11690), Tim23-1 (AT1g17530), Tim23-2 (AT1g72750), Tim23-3 (AT3g04800), B14.7 (AT2g42210), MPPx (AT3g16840), Tim50 (AT1g55900), Ndufs4 (AT5g67590), RISP (AT5g13430), COXII (ATmG900160), \( \alpha \)-ATP synthase (AT2g07698), rug (AT5g60870), Rpotmp (AT5g15700), Tim22 (AT1g18320/AT3g10110), Tim44 (AT2g36870), Tom20-2 (AT1g27390), Tom20-3 (AT3g27080), Tom20-4 (AT5g40930), Sam50 (AT5g05520), Tim21 (AT4g00026), Tim9 (AT3g45680), Tom40 (AT3g20000), cyc1-1 (AT3g27240), and cyc1-2 (AT5g40810).

**Supplemental Data**

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

**Supplemental Figure 1.** Determining the Specificity of B14.7 Antibody.

**Supplemental Figure 2.** Interaction of Tim21 with Complex III in Arabidopsis.

**Supplemental Figure 3.** Analysis of Tim23 Protein and Transcript Abundance in T-DNA Insertion Lines.

**Supplemental Figure 4.** Identification of the Mixed Inner Membrane Complex in the tim23-2 OE Line.

**Supplemental Figure 5.** Determining the Linearity of Antibodies Used and Protein Abundance in Mitochondrial Isolations.

**Supplemental Table 1.** Summary of Changes in Transcript Abundance for Nuclear Genes Encoding Mitochondrial Import and Assembly Components.

**Supplemental Table 2.** Summary of 76 Transcripts Defined as Being Highly Coexpressed with tim23-2 across Public Arabidopsis Microarray Data Sets.

**Supplemental Table 3.** List of Primers Used in This Study.

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**AUTHOR CONTRIBUTIONS**

J.W. and M.W.M. designed the experiments and wrote the article. Y.W., C.C., D.E., O.D., and M.W.M. performed the experimental procedures and carried out the data analysis. E.G. and R.N. carried out gene expression and transcriptome analysis.
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