A RanBP2-type zinc finger protein functions in intron splicing in Arabidopsis mitochondria and is involved in the biogenesis of respiratory complex I

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ABSTRACT

The RanBP2 zinc finger (Znf) domain is a prevalent domain that mediates protein interaction and RNA binding. In Arabidopsis, a clade of four RanBP2 Znf-containing proteins, named the Organelle Zinc (OZ) finger family, are known or predicted to be targeted to either the mitochondria or the plastids. Previously we reported that OZ1 is absolutely required for the editing of 14 sites in chloroplasts. We now have investigated the function of OZ2, whose null mutation is embryo lethal. We rescued the null mutant by expressing wild-type OZ2 under the control of the seed-specific ABSCISIC ACID-INSENSITIVE3 (ABI3) promoter. Rescued mutant plants exhibit severely delayed development and a distinctive morphological phenotype. Genetic and biochemical analyses demonstrated that OZ2 promotes the splicing of transcripts of several mitochondrial nad genes and rps3. The splicing defect of nad transcripts results in the destabilization of complex I, which in turn affects the respiratory ability of oz2 mutants, turning on the alternative respiratory pathway, and impacting the plant development. Protein-protein interaction assays demonstrated binding of OZ2 to several known mitochondrial splicing factors targeting the same splicing events. These findings extend the known functional repertoire of the RanBP2 zinc finger domain in nuclear splicing to include plant organelle splicing.

INTRODUCTION

Plant mitochondria were once free-living organisms, related to contemporary alpha-protobacteria, that were engulfed about 2 billion years ago by the ancestor of eukaryotic cells. It is now accepted that endosymbiosis gave rise to mitochondria and, in plant cells, to the chloroplasts, whose ancestors were cyanobacteria (1). Both organelles have retained their own genomes; however, a gradual loss of genetic content has occurred with a massive transfer of genes from the mitochondria and the chloroplasts towards the nucleus (2).

Because of their prokaryotic origins, gene expression in both organelles has retained some bacterial-like traits, but gene transfer and co-evolution with the nuclear genome has resulted in overlay of eukaryotic features (3,4). The collection of genes still present in the mitochondrial genome is comparable, though not identical, across plant, fungal and animal lineages, usually incorporating some components of the OXPHOS system, tRNAs and ribosomal RNAs and some proteins. The proper function of mitochondrial respiratory complexes and translation apparatus, due to the dual origin of some of their components, either mitochondrial- or nuclear-encoded, require a coordination of gene expression from these two physically different genomes. In addition to the organellar genes that have been transferred to the nuclear genome, many other nuclear-encoded proteins targeted to the organelles function in the control of organelle gene expression. The very complex organelle RNA metabolism, which encompasses several common post-transcriptional processes such as processing, editing and splicing, is a striking illustration of the increasing control of the nucleus over organelle gene expression. Many proteins involved in these processes are not de-
rived from the endosymbiont but were acquired by the host-cell during the nuclear-organellar co-evolution.

Most of the introns found in the angiosperm mitochondrial genomes belong to the group II class, with the presence of a group I intron limited to the coxl gene, only in certain lineages (5). Mitochondrial introns are found in the coding sequences of protein coding genes and must be spliced out in order to produce functional products. Some of the introns are fragmented and need to be reassembled in trans before being removed. These trans-splicing events occur in the mitochondrially-encoded subunits of the NADH:ubiquinone oxidoreductase complex (complex I), nadl, nad2 and nad5, and also in the cox2 intron in Allium cepa (6–9). Since endosymbiosis, these group II introns have degenerated compared to their prokaryotic ancestors. In particular, most of the mitochondrial introns have lost their intron-encoded proteins or maturases; only one matR, has been retained in the fourth intron of nad1 (10). As a result, splicing of these introns does not occur in vivo without the assistance of nuclear-encoded splicing factors (11).

Another type of RNA processing in chloroplasts and mitochondria that is facilitated by nuclear proteins is plant organelle RNA editing, which is a post-transcriptional event in which cytidines are converted to uridines at many sites in chloroplast and mitochondrial transcripts. We have identified OZI (Organelle Zinc finger 1), a protein containing a RanBP2-type zinc finger (Znf), as a chloroplast editing factor, raising the possibility that other OZ family members could be editing factors. Here, we investigate the function of OZ2 (Atlg55040), which, when mutated, results in an embryo defective (EMB) phenotype, arresting development at the cotyledon stage (12,13). We rescued the homozygous oz2 mutant embryo with a wild-type transgene driven by an embryo-specific promoter, a strategy developed to study post-embryonic functions of EMB genes (14).

The oz2 rescued mutant plants exhibit a marked delay in development with dark green and curled rosette leaves. These partially complemented mutants have a dramatic decrease in the splicing efficiency of six mitochondrial introns, five of which are located in nad transcripts, which encode subunits of Complex I. Our data show that the assembly of complex I is seriously compromised as a result of the defective splicing. Therefore, unexpectedly, OZ2 is not an editing factor, but a mitochondrial splicing factor whose knockout leads to an embryo-lethal phenotype. Thus, we have determined that a RanBP2 zinc finger-containing protein has a novel function as an essential splicing factor in higher plant mitochondria.

**MATERIALS AND METHODS**

**Plant material**

The Arabidopsis T-DNA insertion lines SALK_043472C and WiscDsLox233237,13N (CS849339) in the OZ2 gene were ordered from the Arabidopsis Biological Resource Center (https://abrc.osu.edu/). After 3 days of stratification, seeds were planted in soil growing in a growth room (14 h of light/10 h of dark) at 26°C. Genotyping was done by polymerase chain reaction (PCR) with OneTaq Quick-Load 2X Master Mix (New England Biolabs) using primer pairs listed in Supplementary Table S1. The wild-type alleles were amplified using the primer pairs SALK_043472C-LP and SALK_043472C-RP, and CS84933-LP and CS84933-RP. The mutant alleles were amplified using the primer pairs SALK_043472C-RP and LBB1.3, and CS84933-RP and p745. The PCR products were sequenced at Cornell University Life Sciences Core Laboratories Center.

**Constructs used in this study**

A 2411-bp fragment of the ABI3 (AT3G24650) promoter was amplified using a HindIII restriction site forward primer 5’-AAAGCTT ATGGATCCATGTAACGCTACTGATC and a HindIII restriction site reverse primer 5’-GTACCGTGAAGTGAAATGAAACAATAAACTAG, and the Platinum SuperFi Green PCR Master Mix (Invitrogen) and cloned into the pCR8/GW/TOPO vector (Invitrogen). The sequence of the cloned ABI3 promoter was verified by using the primers listed in Supplementary Table S1 and digested with HindIII and NheI. The insert was separated from the vector by electrophoresis, gel purified (PureLink Quick Gel Extraction Kit, Invitrogen) and ligated to a Gateway-converted pBII121 binary vector previously digested with HindIII and XbaI. The ligated product resulted in swapping the CaMV35S promoter for the ABI3 promoter in the Gateway pBII121 vector. The Gateway pBII121 vector had been engineered previously by changing the resistance marker from kanamycin to Basta resistance using a strategy similar to exchanging the promoter. The Basta resistance cassette was amplified from the pEarleygate 205 vector (15) using a Pmel restriction site forward primer 5’- GCTATTTACACTGTGATATTCGGTGAAGTGGAAATGAAACAATAAACTAG, and a HindIII restriction site reverse primer 5’- GTTAAACTTTCTGAGATTTTTCAAAATCTAGTC and a HindIII restriction site reverse primer 5’- AAGCCTTCCGGATCTGATAATTTTATTTTGAAGATT. The amplified Basta cassette was cloned into the pCR8/GW/TOPO vector, digested with HindIII and Pmel, gel purified and ligated to a Gateway pBII121 vector previously digested with HindIII and Pmel to release the Kanamycin resistance cassette. The final binary vector pBII121-ABI3-GW-Basta is a Gateway-compatible binary vector whose transgene is under the control of the ABI3 promoter, and the selection marker is resistance to Basta.

The cDNA clone of OZ2 used in this study was reverse-transcribed by SuperScript® III Reverse Transcriptase (Invitrogen) with the OZ2-R1 primer (Supplementary Table S1) from RNA extracted from wild-type Arabidopsis Columbia using PureLink® RNA Mini Kit (Invitrogen). The full length OZ2 CDS was amplified with OZ2-F1 and OZ2-R1 primers and cloned into the pCR8/GW/TOPO vector; after verifying the sequence with primers listed in Supplementary Table S1, the OZ2 CDS was transferred to the pBII121-ABI3-GW-Basta by LR Clonase II (Invitrogen).

A recombinant protein that included the two RanBP2 Zn finger domains (residues 343P to 454R of the OZ2 protein) was expressed in pET28a (Novagen). An ampiclon was produced from OZ2 cDNA with oligonucleotide primers (AtOZ2_ZnFing_For; AtOZ2_ZnFing_Rev. Supplementary Table S1) and cloned into the EcoRI and HindIII sites of the pET28a plasmid.
Generation of transgenic plants

AB13-OZ2 (hereafter OZ*) in the pBI121 vector was transformed into Agrobacterium tumefaciens GV3101, and floral dip transformation of heterozygous o2/OZ2 plants (SALK_043472C) was performed as described in (16). Progeny were selected on soil by spraying Basta on young seedlings; resistant plants were genotyped to determine whether they were homozygous wild-type OZ2/OZ2/OZ*, heterozygous o2/OZ2/OZ*, or homozygous mutant o2/o2/OZ*. The wild-type plants were discarded, the homozygous mutant plant was investigated, and the heterozygous transgenic plants were grown to the next generation. T1 progeny from the heterozygous T0 parental plants were also subjected to Basta selection, and several homozygous o2/o2/OZ* plants were recovered in the T1 generation.

OZ2 localization

OZ2 was amplified from OZ2 cDNA with Phusion polymerase (Thermo Scientific) and the primers OZ2-F1 and OZ2-nostop-R1 using standard protocols. 3′-A overhangs were added with Taq (QIAGEN) by incubating at 37°C for 10 min. After purification, the amplicons were cloned into pCR8-TOPO/TOPO (Invitrogen) to use in a Gateway cloning reaction with a modified pEXSG vector (17) containing an EYFP C-terminal tag using LR Clonase II (Invitrogen) to produce pEXSG-OZ2-YFP.

Arabidopsis Col-0 plants and Nicotiana benthamiana were grown on soil in a long day (16 h) conditions for 3–5 weeks for Arabidopsis and 5–6 weeks for N. benthamiana. pEXSG-OZ2-YFP was transfected into Arabidopsis and N. benthamiana protoplasts using the method outlined in (18), using ~3.0 × 10^5 cells per transformation.

Protoplast mitochondria were stained with Mitotracker® Orange CM-H2TMRos (500 nM; Thermo Scientific), using DMSO as the solvent and W5 buffer. Protoplasts were incubated in the dark for 45 min and then were pelleted (1000 × g, 5 min) and resuspended in W5 buffer (500 µl). Protoplasts were imaged using a Zeiss Axio Observer LSM 710 microscope and C-Apochromat 40×/1.20 W Korr M27 objective.

RNA analysis

Total RNA was extracted from Arabidopsis rosette leaves (4–5-week-old plants) by pulverizing green tissue with liquid nitrogen and using Trizol (Ambion) according to the manufacturer’s instructions. RNA was further purified with the PureLink® RNA Mini Kit (Invitrogen). The purified RNA was treated with Turbo DNAse (Invitrogen). Mitochondrial RT-PCR products were amplified with the same set of primers used in (19). The real-time quantitative RT-PCR conditions and analysis are similar to the ones outlined in (20), except for the use of an updated CFX Maestro Software (Bio-Rad) that allows performance of an ANOVA. The primers used for the quantitative RT-PCR to measure the expression of mitochondrial transcripts are described in (21), except for nad2-ex2ex3 and nad2-ex2in2, which are given in Supplementary Table S1. The qRT-PCR primers to measure the expression levels of alternative respiratory pathway genes are described in (22) for AOX1a, NDA1, NDA2, NDB1, NDB2, NDB3, NDB4 and NDC1. The qRT-PCR primers for AOX1c, and AOX1d are described in (23) while the primers for AOX1b, and AOX2 were retrieved from qPrimerDB, a qPCR primer database (24). The poisoned primer extension assays were performed as described previously (25) with primers listed in Supplementary Table S1.

Blue native gel and respiratory complex in-gel activity assay

Some of the rescued homozygous o2 mutants (o2/o2/OZ*) were able to produce a reasonable amount of viable seeds. Mitochondria were extracted from liquid-grown Arabidopsis seedlings following the protocol of Murcha and Whelan (26). The yield of mitochondrial extraction was evaluated by Bradford assay (Bio-Rad). The sample preparation for the Blue native (BN) gel was done according to the protocol of Schertl and Braun (27). The gel used for the BN-PAGE was a pre-cast Native PAGE Novex 3–12% Bis–Tris gel (Invitrogen). The conditions for the run were: 4°C, 150 V for 60 min, 250 V for 180 min. Approximately 50 µg of mitochondrial protein was loaded per lane. After completion of the run, slices of the gel were either stained with Coomassie R-250 according to the manufacturer’s instructions, or incubated overnight with freshly prepared solutions for in-gel-activity at room temperature following the protocol of Schertl and Braun (27). The reactions were then stopped by transferring the gels into fixing solution.

Zinc binding analysis

Recombinant OZ2 protein containing the two RanBP2 Zn finger domains was expressed in E. coli cells (Rosetta 2 (BL21) DE3 pLys) in 1 l cultures at 18°C using 1 mM of IPTG and purified on Ni-NTA resin. Recombinant protein was digested overnight with thrombin (1 U/mg protein) at 4°C, and dialyzed twice in 20 mM ammonium acetate. The mass spectrometry analysis of the recombinant OZ2 containing the two RanBP2 Zn finger domains was performed as in (28) with a Waters QTOF2 instrument with an electrospray ion voltage of 3.6 kV, a cone voltage of 40 V, and a desolvation temperature of 120°C. In addition to native conditions performed in 20% ammonium acetate proteins were denatured using 50% acetonitrile 0.1% formic acid to release bound zinc ligands.

Yeast two-hybrid assay

The mature coding sequences (without predicted N-terminal mitochondrial transit peptide) of OZ2 and mitochondrial splicing factors ABO5, BIR6, MAT2, mCSF1, MISF26, MISF74, MTL1, OTP439, PMH2, RUG3, SLO3 and WTF9 were amplified using Y2H primer pairs to measure the expression levels of alternative respiratory pathway genes described in (22) for AOX1a, NDA1, NDA2, NDB1, NDB2, NDB3, NDB4 and NDC1. The qRT-PCR primers for AOX1c, and AOX1d are described in (23) while the primers for AOX1b, and AOX2 were retrieved from qPrimerDB, a qPCR primer database (24). The poisoned primer extension assays were performed as described previously (25) with primers listed in Supplementary Table S1. The qRT-PCR vectors were used as negative controls in yeast two-hybrid

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assays. Yeast mating strains PJ69-4a and PJ69-4c were individually transformed with pGADT7/F and pGBK77/F plasmids, respectively. Single transformants were mated to produce diploid double-transformant yeast on YPAD agar plates. Yeast harboring testing pairs were grown in leucine- and tryptophan-deficient media overnight; then, 10 \( \mu l \) of each culture was spotted onto leucine-, tryptophan-, histidine-, adenine-deficient plates and then diluted with water to OD600 0.5, 0.05 and 0.005. Survival/growth plates were imaged after three days of incubation at 30°C.

Bimolecular fluorescence complementation (BiFC)

The full-length coding sequences (including N-terminal mitochondrial transit peptides but without the stop codon) of OZ2 and mitochondrial splicing factors ABO5, BIR6, MAT2, mCSF1, MISF26, MISF74, MTL1, OTP439, PMH2, RUG3, SLO3 and WFT9 were amplified using BiFC primer pairs listed in Supplementary Table S1. PCR products were first cloned into pCR8/GW/TOPO and then pEXSG-nYFP and pEXSG-cYFP BiFC destination vectors via Gateway cloning. Strain A. tumefaciens GV3101::pMP90RK were transformed via electroporation with 1 \( \mu g \) plasmid. Electroporation was conducted with the following parameters: capacitance 25 \( \mu F \), voltage 2.0 kV, resistance 200 \( \Omega \), pulse length \( \sim 5 \) ms. Agrobacteria were selected on LB agar plates containing kanamycin (50 \( \mu g/ml \)), gentamicin (25 \( \mu g/ml \)) and carbenicillin (50 \( \mu g/ml \)).

5-ml cultures of individual transformed Agrobacteria were incubated for 2 days at 28°C and resuspended in a solution of 2-(N-morpholine)-ethanesulphonic acid (pH 5.6), 20 mM Na3PO4, and 150 mM acetosyringone. Infiltration samples were made by mixing bacterial cultures carrying pEXSG-nYFP, pEXSG-cYFP and P19 at equal OD600 to a final OD of 0.9. Leaves of 4- to 6-week-old N. benthamiana plants grown in long-days condition were agroinfiltrated as described in (29). 2–3 days post-infiltration, 2-mm squares of plants grown in long-days condition were agroinfiltrated as described in (29). 2–3 days post-infiltration, 2-mm squares were cut from the infiltrated leaf area and imaged using fluorescence microscopy as described above.

RESULTS

Arabidopsis \( oz2 \) mutant plants are embryo-lethal but can be rescued by the use of the ABI3 promoter

Both OZ1 and OZ2 contain two Znf domains while OZ3 and OZ4 have three and four Znf domains, respectively (Figure 1A). The RanBP2 Znf domain is 30 aa long and is a C4-type member of the broader zinc finger family, known to have unique functions and an unusually diverse distribution in plants (30).

In order to investigate the function of OZ2, we obtained two independent insertional mutant lines from the Arabidopsis Biological Resource Center (https://abrc.osu.edu/) (Supplementary Figure S1A). Sequencing of the mutant allele confirmed the location of the T-DNA insertion in the third exon for SALK_043472C, while the insertion in Wis-cDsLox233237,13N was found to be in the fourth intron instead of the fourth exon as annotated in the database. No homozygous mutant plants were found in the segregating progeny of a heterozygous parent plant of either insertional line, indicating that the \( oz2 \) mutation is embryo lethal (Supplementary Figure S1B).

To overcome the embryo lethality of the \( oz2 \) mutation and study the function of OZ2, we attempted to rescue the mutant embryo by complementing heterozygous lines (\( oz2/oz2 \)) with a cDNA carrying the wild-type coding sequence under the control of the seed-specific ABSCISIC ACID-INSensitive3 (ABI3) promoter: ABI3- 

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The Arabidopsis subcellular database SUBA (http://suba.live/) strongly supports a plastid location for OZ2; however, the most recent version of TargetP (TargetP-2.0), a predictive software for the subcellular localization of proteins, predicts the localization of OZ2 to be mitochondrial with high confidence (31). Predotar, another predictive software, favours a plastidial over a mitochondrial localization of OZ2, but only by a slight margin (32). To resolve these ambiguities and find out the subcellular localization of OZ2, we transiently expressed in Arabidopsis or N. benthamiana protoplasts a YFP translational fusion comprising the complete coding sequence of OZ2 (Figure 2). In protoplasts of both species, confocal microscopy demonstrated that the YFP fluorescence co-localizes with the fluorescent signal from Mitotracker Orange that specifically labels mitochondria. Thus, this analysis demonstrated that OZ2 was targeted to mitochondria and not plastids, supporting a previous localization study in Arabidopsis (12).

Splicing of most transcripts of the complex 1 and \( rps3 \) is severely impaired in \( oz2 \) mutants

Since OZ1 is an essential plastid editing factor, and OZ2 is a mitochondrial-targeted protein closely related to OZ1, we tested whether a defect in mitochondrial editing could be the source of the morphological phenotype caused by the \( oz2 \) mutation. Survey of the mitochondrial editome by bulk-sequencing of the RT-PCR products corresponding to the whole set of mitochondrial genes did not reveal any defects in editing in the \( oz2 \) mutant T0 plant \((\text{oz2/oz2/ABIS})\) (Supplementary Figure S2A). However, during the amplification of the RT-PCR products, four mitochondrial genes, \( rps3 \), \( nad2 \), \( nad5 \), \( nad7 \) and \( rps3 \), exhibited a ladder pattern with a marked reduction of the mature product in the \( oz2 \) mutant T0 plant (Supplementary Figure S2B). A feature common to these four genes is that splicing is involved in their transcript maturation. In addition, even though the RT-PCR...
Figure 1. OZ2 belongs to a four-member family, and its partial complementation in a mutant background leads to a stunted growth phenotype. (A) Schematic representation of each OZ protein with the location of their RanBP2-type zinc finger motifs shown in blue. (B) The rescued oz2/oz2/OZ2* mutant plant exhibits slow growth and dark curled leaves. (C) Identification of several independent T1 oz2/oz2/OZ2* plants confirm the morphological defect caused by the oz2 mutation. Six T1 oz2/oz2/OZ2* plants show a stunted phenotype similar to one of the T0 plants previously identified. In each panel, the three T1 plants are progeny from a T0 Oz2*/oz2/OZ2 plant. The three panels represent three different transgenic events.

Figure 2. OZ2 is targeted to mitochondria. (A) Transfection of Arabidopsis protoplasts with OZ2 fused to YFP. From left panel to right panel: OZ2 location appears as green dots, mitochondria are stained in red with MitoTracker, chloroplasts autofluorescence is artificially colored blue, overlay of these three panels demonstrate the location of OZ2 in mitochondria as yellow (green + red) dots. (B) Transfection of Nicotiana benthamiana protoplasts with OZ2 fused to YFP. Some of the green spots (OZ2-YFP) can be seen to co-localize with the red spots (mitochondria).

reaction was not meant to be quantitative, there was also a noticeable reduction of the amplified cDNA for two other nad genes, nad1 and nad4 in the oz2 mutant (Supplementary Figure S2C).

We performed quantitative RT-PCR (qRT-PCR) assays for all the mitochondrial transcripts using a set of primers that allowed survey of not only the abundance of the transcripts, but also the efficiency of splicing by discriminating between the spliced and unspliced transcripts (21). This analysis was conducted on three wild-type T1 plants and four oz2 mutant (oz2/oz2/OZ*) plants: the T0 oz2/oz2/OZ* plant and three of the T1 plants derived from three different heterozygous T0 oz2/OZ2/OZ* plants (Figure 1B and C). Therefore, the four oz2 mutant plants anal-
yzed were produced via four independent transgenic events, presumably with random insertion of the ABI3:OZ2 construct in the genome.

The qRT-PCR analysis sheds light on the molecular defect caused by the oz2 mutation that resulted in the reduction of mature products for some of the nad genes as observed after RT-PCR (Supplementary Figure S2B and C). The maturation of some mitochondrial nad genes is quite complex and requires both cis- and trans-splicing events \((8,9,33)\). For instance, the nad1 gene is split into five coding segments localized on three precursor transcripts that are transcribed from three genomic areas scattered over 136 kb in the Arabidopsis mitochondrial genome (Figure 3A). Two trans-splicing events and two cis-splicing events are necessary for the complete maturation of the nad1 transcript. The qRT-PCR results revealed a major and highly significant \((P < 0.0001)\) reduction of the intron 1 spliced transcript, denoted nad1-ex1ex2 in Figure 3B. Its transcription abundance is reduced around 60 times in the oz2 mutant compared to the wild-type plant. The RNA comprising intron IB, one of the precursors of the intron 1 spliced product, exhibited a significant \((P < 0.05)\) increase in the oz2 mutant (Figure 3B). This increase is more moderate than the decrease of the intron 1 spliced transcript, 5 times versus 60 times, respectively, supporting a true defect in splicing instead of a destabilization of the precursor transcript. None of the other nad1 spliced species, pertaining to the splicing of introns 2, 3 or 4, exhibited a reduced amount in the oz2 mutant; the abundance of these spliced products was significantly \((P < 0.05)\) but moderately \((5–7\) times) increased in the oz2 mutant (Figure 3B). The specificity of the effect of oz2 mutation, with a marked reduction of only the intron 1 spliced transcript, strongly suggests a defect in splicing rather than a global destabilization across the entire nad1 transcript.

Defective splicing events caused by the oz2 mutation were observed in other nad transcripts. Like nad1, the nad2 gene showed a very significant \((P<0.01)\) reduction in abundance of only one of its spliced species (Figure 4A). The intron 3 spliced transcript is reduced 830 times in abundance on average in the oz2 mutant plant relative to wild-type. None of the other transcript species assayed in nad2 showed a significant variation in the mutant plant. The increase in the precursor to this spliced species, nad2-in3 in Figure 4A, although not significant, further supports the involvement of OZ2 specifically in splicing, since the reduced amount of the spliced product can thus not be caused by a decreased availability of the precursor. Unlike the nad1 first intron which is trans-spliced, the third intron of nad2 is cis-spliced. Two of the nad5 spliced products are very significantly \((P < 0.01)\) reduced in abundance in the oz2 mutant (Figure 4B). The intron 1 cis-spliced transcript reduction is about 25 times in the oz2 mutant plant while the intron 3 or intron 4 trans-spliced transcript reduction amounts to 60 times. Due to the short length of nad5 exon 3 (22 nt), we did not try to discriminate between a defective splicing of intron 3 or intron 4 by using a qRT-PCR assay. The splicing of intron 2 in the nad7 transcript is the last example of a defective splicing event in the nad transcripts found in the oz2 mutant plants (Figure 4C).

The only non-complex I gene whose transcripts exhibit defective splicing is rps3, encoding the ribosomal protein S3 (Figure 4D). The significant reduction of spliced rps3 in oz2 is moderate, reaching about a tenth of the amount of the wild-type plant, while the amount of unspliced transcript is about 10 times the amount found in the wild-type.

In summary, among 68 transcript species tested by qRT-PCR assays, six transcripts, all of them spliced products, showed a very significant reduction of their abundance in the oz2 mutant plant versus the wild-type plant (Supplementary Figure S3). The decrease ranged from 0.11 \((rps3)\) to 0.0011 \((nad7-ex2ex3)\). Eleven transcripts exhibited a significantly increased abundance in the oz2 mutant relative to the wild-type, ranging from 3 to 10 times. These transcripts could be spliced products, e.g. ccmFe-ex1ex2, unspliced products, e.g. nad1-in1ex2, or ones not requiring splicing for their maturation, e.g. atp1 (Supplementary Figure S3).

In order to confirm the splicing defect caused by the oz2 mutation, we performed a semi-quantitative RT-PCR assay comparing the amount of amplified products, spliced and unspliced, in the wild-type versus oz2 \((oz2/oz2/OZ*)\) at different number of cycles (Supplementary Figure S4). At 30 cycles, a rps3 spliced product is detectable in the wild-type lanes, but not in the oz2 lanes (Supplementary Figure S4A). Conversely, an unspliced rps3 product is detectable in the oz2 lanes but not in the wild-type lanes. At 35 cycles, the rps3 spliced product is now detectable in the oz2 lanes; however, its abundance is reduced when compared to the wild-type lanes. The opposite is true for the unspliced product; its abundance is higher in the oz2 lanes than in the wild-type lanes. The results of this assay recapitulate the data from the qRT-PCR assay with a roughly 10-time reduction of spliced (unspliced) transcript in the oz2 (wild-type) plant. We also tested the abundance of the nad2-intron 3 spliced and unspliced product (Supplementary Figure S4B). At 35 cycles, the spliced product is readily detectable in the wild-type lanes but not in the oz2 mutant lanes. The unspliced product is more abundant in the oz2 lanes than in the wild-type lanes. At 40 cycles, there is still no detectable trace of spliced nad2 in the oz2 lanes, while the amount of amplified product is easy to detect in the wild-type lanes. These results validate the qRT-PCR data that estimated the spliced product to be on average 830 times less abundant in the oz2 mutants than in the wild-type plants.

A poisoned primer extension (PPE) assay was also conducted to confirm the splicing defect for two of the affected transcripts in the oz2 mutant plants (Figure 5). In this assay, a fluorescent primer is extended in the presence of a template, either RNA or cDNA, and a mix of deoxyribonucleotide triphosphates and one deoxyribonucleotide triphosphate. The primer is located on the exon just upstream of the junction intron/exon to be tested. The extension reaction will stop at the first nucleotide in the template that is complementary to the deoxyribonucleotide in the reaction mix. The deoxyribonucleotide is chosen so that the length of the extension product differs between the spliced versus unspliced template. Assaying the splicing of the third exon in the nad2 transcript by a PPE reaction shows both spliced and unspliced extension products for the wild-type lanes but
the amount of spliced intron 1 transcript (four splicing events, two cis-splicing events (II and IV), and two trans-splicing events (I and III)). Arrows indicate the primers used in the qRT-PCR analysis; red arrows indicate the primers used to quantify the amount of unspliced transcripts. (B) qRT-PCR analysis of nad1 transcript species demonstrated that the amount of spliced intron 1 transcript (nad1-ex1ex2) is severely reduced in oz2/oz2/OZ* (around 60 times), while the amount of unspliced precursor (nad1-in1) is slightly higher in oz2/oz2/OZ* than in the wild-type (wt) plants (around 5 times). Splicing of intron 1 is the only splicing defect for nad1 as the other spliced transcripts, nad1-ex2ex3, nad1-ex3ex4, nad1-ex4ex5, exhibit a slight but significant increased amount in oz2 compared to the wild-type plants. The assay was performed on the RNAs from plants shown in Figure 1: three wild-type T1 plants, three oz2/oz2/OZ* T1 plants, and one oz2/oz2/OZ* T0 plant. Three technical replicates were measured for each data point (*P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001).

no detectable spliced product in the oz2 lanes (Figure 5A). We performed this PPE assay with two different dideoxynucleotides, either ddCTP or dTTP, and in both experiments there was no detectable spliced product in the oz2 lanes. We also assayed the splicing of nad7-intron 2 by PPE with two different dideoxynucleotides, either ddATP or ddCTP (Figure 5B). Like for nad2, both spliced and unspliced products are present in wild-type lanes, while the spliced product is absent in three of the oz2 mutant lanes. A faint band representing the spliced product is detectable in one of the oz2 lanes, but with much less intensity than in the wild-type lanes.

In conclusion, our data strongly support a role of OZ2 in the splicing of most mitochondrial complex I and rps3 transcripts. The splicing defect affects one to two splicing events per transcript and can alter both transcripts. The splicing defect affects one to two splicing events (I and III). Arrows indicate the primers used in the qRT-PCR analysis.

oz2 mutants are severely impaired in respiratory complex I

Given the molecular defects caused by the oz2 mutation, we considered that a respiratory dysfunction, more specifically a deficiency in complex I, could be the source of the developmental alterations observed in the oz2 mutant plants. Most of the NAD proteins are highly hydrophobic and therefore lack effective antibodies. We examined the steady-state levels of the different respiratory chain complexes after separation by blue-native PAGE (BN-PAGE) (Figure 6). An approximately equivalent amount of mitochondrial protein (ca. 50 μg/lane) from both the wild-type and the oz2 mutant plant (oz2/oz2/OZ*) was loaded onto the gel. Coomassie staining of the gel after BN-PAGE shows that most of the bands representing the different complexes are present in the same amount in both the wild-type and the oz2 lanes, except for complex I (Figure 6). A marked reduction of complex I bands, representing both complex I and supercomplex I + III, is apparent in the oz2 lane compared to the wild-type lane. An in-gel NADH dehydrogenase activity assay confirmed the significantly decreased amount of complex I observed in the Coomassie-stained gel (Supplementary Figure S5). Succinate dehydrogenase (complex II), cytochrome c oxidase (complex IV), and ATPase (complex V) were also tested and showed similar activities in oz2 and wild-type plants, confirming the observations from the Coomassie stained gel (Supplementary Figure S5).

Previously described Arabidopsis mutants lacking complex I have elevated levels of genes involved in the alternative respiratory pathway (22,34–36). We assessed the level of expression of alternative oxidase (AOX) genes, isoforms AOX1a-d and AOX2, and the alternative external (NDBI–4) and internal (NDA1–2 and NDA1C) NAD(P)H dehydrogenase genes in the oz2 mutant by qRT-PCR analysis (Figure 7). This assay revealed a significant (P < 0.01) and very high induction of the expression of AOX1d in the oz2 mutant plants, whose steady state level was ~270-fold higher than in the wild-type plants. The NDA2 gene also exhibited a significant (P < 0.05) but moderate induction of its expression in the oz2 mutant plants, with a steady state level ~3-fold higher than in the wild-type plants. Three alternative NAD(P)H dehydrogenase genes, NDA1, NDB1, and
Figure 4. OZ2 is a mitochondrial splicing factor that controls the splicing events of complex I transcripts and rps3. (A) Survey of the splicing in nad2 by qRT-PCR analysis. The spliced species without intron 3 shows a significant reduction in oz2/oz2/OZ*. (B) Survey of the splicing in nad5 by qRT-PCR analysis. Two of the spliced species of this transcript show a significant reduction in the oz2/oz2/OZ* mutant. (C) Survey of the splicing in nad7 by qRT-PCR analysis. The spliced species without intron 2 shows a significant reduction in oz2/oz2/OZ*. (D) Survey of the splicing in rps3 by qRT-PCR analysis. The spliced rps3 shows a significant reduction in the oz2/oz2/OZ* mutant while the unspliced precursor amount is significantly increased. The order of the plants assayed is similar to Figure 3: three wild-type T1 plants, three oz2/oz2/OZ2*T1 plants, and one oz2/oz2/OZ2*T0 plant. Three technical replicates were measured for each data point (*P < 0.05, **P < 0.01)
Figure 5. Poisoned primer extension (PPE) assay confirms that \( \text{o}2 \) mutation results in a splicing defect. Above the PPE gels is a schematic representation of the two cDNAs, spliced or unspliced, that were used as templates for the PPE reaction. The position of the primer represented by an arrow is shown upstream of the junction of exon (square) with intron (line). The length of the PPE products is indicated for each chain terminator used and each extension product is color coded (red for ddCTP, green for ddTTP, and blue for ddATP). The first nucleotide complementary to the dideoxynucleotide used in the PPE reaction that will stop the extension is indicated on either the exon (spliced) or the intron (unspliced). (A) \( \text{nad}2 \) PPE assay shows a detectable amount of spliced product in the wild-type lanes, while there is no detectable spliced product in the \( \text{o}2\text{/o}2\text{/OZ}^* \) lanes. Conversely, some unspliced product is detectable in both the wild-type and the \( \text{o}2\text{/o}2\text{/OZ}^* \) lanes. Left panel: the PPE reaction included the chain terminator ddCTP such that the extension terminated after the addition of 8 or 10 nucleotides on unspliced or spliced templates, respectively. Right panel: the PPE reaction included the chain terminator ddTTP such that the extension terminated after the addition of 2 or 11 nucleotides on unspliced or spliced templates, respectively. (B) \( \text{nad}7 \) PPE assay shows a detectable amount of spliced product in all the wild-type lanes, while there is detectable spliced product in only one of the \( \text{o}2\text{/o}2\text{/OZ}^* \) lanes. Conversely, some unspliced product is detectable in both the wild-type and the \( \text{o}2\text{/o}2\text{/OZ}^* \) lanes. Left panel: the PPE reaction included the chain terminator ddATP such that the extension terminated after the addition of one or five nucleotides on unspliced or spliced templates, respectively. Right panel: the PPE reaction included the chain terminator ddCTP such that the extension terminated after the addition of two or eight nucleotides on spliced or unspliced templates, respectively. P: primer, U: unspliced, S: spliced, wt: wild-type plant, \( \text{o}2\text{/o}2 \): \( \text{o}2 \) mutant plant, NT: no template.
Figure 6. Blue native gels demonstrate that only complex I is seriously impaired in the oz2/oz2/OZ* mutant. Arabidopsis mitochondrial proteins were solubilized by digitonin and subsequently separated by 1D BN-PAGE. After electrophoresis, slices of the gel were stained with Coomassie.

NDC1, show a significant reduction of their expression in the oz2 mutants; however, the magnitude of this decrease (3–11-fold) is much less pronounced than the induction of AOX1d (Figure 7).

Altogether these results lead us to conclude that oz2 mutants are complex I-deficient plants. The splicing defect of nad transcripts results in the destabilization of complex I, which in turn affects the respiratory ability of oz2 mutants, turning on the alternative respiratory pathway, and impacting the plant development with stunted growth and curly leaves. These developmental abnormalities are typical of complex I mutants, e.g. (22,35,37).

OZ2 binds two zinc ions

The structure of the RanBP2 Zn finger domain was solved with the first domain of the ZNF265 (also called ZRANB2 and Zis), a human splicing factor (38). NMR spectra in both the absence and presence of Zn showed a substantial increase in chemical shift dispersion in the presence of Zn, demonstrating this domain to be a genuine zinc-binding domain. We expressed 112 aa of OZ2 encompassing the two RanBP2 Znf domains in E. coli and performed mass spectrometry (MS) analysis of this polypeptide under native and denaturing conditions (Figure 8). Native MS can determine the molecular mass of polypeptides under conditions that retain prosthetic groups so that covalently bound ligands can be identified based on the difference between native and denaturing MS spectra. Recombinant OZ2 protein showed two peaks under native conditions with masses of 15 234 and 13 717 (Figure 8B). The presence of two polypeptides after digestion with thrombin is explained by the presence of two cleavage sites, one canonical (LVPRGS) and one cryptic (QMGRGS) (Figure 8A). Under denaturing conditions, the masses of these two polypeptides are 15 108 and 13 590, respectively. The difference in mass between these two polypeptides, 126–127 Da, corresponds to two zinc atoms. Although zinc has an atomic mass of 65.4, cysteines coordinate zinc directly to the sulfur atom with loss of a proton. Therefore, the observed mass difference is consistent with two bound zinc ions, likely one zinc atom per RanBP2 Zn finger domain.

OZ2 binds specifically to some known splicing factors

The requirement of OZ2 for certain mitochondrial splicing events may be mediated by its binding with other splicing factors. We tested the possibility of OZ2/mitochondrial splicing factor interactions by in vitro yeast two-hybrid (Y2H) assay and in vivo bimolecular fluorescence complementation (BiFC). Mitochondrial splicing factors were chosen for testing based on their sharing of target introns with OZ2; we also included factors that do not share target sites with OZ2 as possible negative controls (BIR6, MISF74 and WTF9). Some of the mitochondrial splicing factors sharing common targets with OZ2 are general factors like MAT2, mCSF1 and PMH2, which affect the splicing of several sites, while other are specific and control one to two splicing events, e.g. ABO5 and RUG3, respectively. The Y2H assays showed fewer OZ2/splice factor interactions than the BiFC assay, 4 versus 7, respectively (Figure 9, Table 1). All the interactions detected by the Y2H assay were confirmed by the BiFC assay (Table 1, Figure 9, Supplementary Figure S6). Among the four positive interactions detected by both assays, three involved splicing factors sharing targets with OZ2, one general factor (PMH2), and two specific factors (ABO5 and MISF26). One consistent interaction identified by both Y2H and BiFC assays involves BIR6, a protein that does not share a target splicing site with OZ2. The other two negative controls, MISF74 and WTF9 did not show any interaction with OZ2. The BiFC assay revealed three additional interactions not detectable by the Y2H assay, two with the general splicing factors MAT2 and mCSF1 and one with the specific factor MTL1, all of which share common splicing targets with OZ2 (Table 1, Figure 9). Overall, our interaction data are consistent with the specificity of the OZ2 splicing function being mediated in some instances by the interaction of OZ2 with other mitochondrial splicing factors.

DISCUSSION

We have established the function of OZ2 in mitochondrial splicing by rescuing oz2 homozygous mutants. These plants, in addition to a delayed growth phenotype, show a significant and pronounced reduction in the splicing of six mitochondrial introns.

The only referenced domain found in OZ2 is a RanBP2-type zinc finger (PFAM00641, IPR001876), 30 residues long, which contains four cysteines required for coordinating a zinc ion that is necessary for the structural fold of the
Figure 7. Quantitative RT-PCR analysis of alternative transport pathway transcripts show a strong induction of AOX1d in the oz2/oz2/OZ2* mutants. The assay was performed on the same RNAs from the same plants tested for the whole mitochondrial transcriptome: three wild-type T1 plants, three oz2/oz2/OZ2* T1 plants, and one oz2/oz2/OZ2* T0 plant. Three technical replicates were measured for each data point (*P < 0.05, **P < 0.01).

Figure 8. OZ2 binds two zinc atoms per polypeptide. (A) Amino acid sequence of the recombinant protein subjected to mass spectrometry analysis. Bolded sequences represent part of the recombinant OZ2 protein (aa 343-454), underlined are the two RanBP2Znfinger domains. Thrombin cleavages confirmed by MS are indicated by arrows. (B) Left, MS spectra indicate the projected mass for OZ2 under native conditions. At right, spectra are for denaturing conditions. The mass difference between the native and denatured conditions (126–127 Da) corresponds to the mass of two zinc atoms.

Our data support the existence of a zinc ion bound to each of the zinc fingers (Znf) present in OZ2. This domain is ancient, prevalent and found in all domains of life, archaea, bacteria, and eukaryotes. Originally described in the RanBP2 nucleoporin, the domain mediates binding of RanBP2 with high specificity to exportin-1, a protein which is a key component of the nuclear export pathway (39). The RanBP2-Znf domain is found in Nup153, another nucleoporin, where it also mediates protein-protein interactions (40). More relevant to the present work is the fact that variants of the domain are found in ZRANB2, a human spliceosomal protein able to induce alternative splicing (41). The two RanBP2-type Znf domains found in ZRANB2 are able to bind ssRNA with high affinity and specificity (42). Similar Znf domains in human proteins implicated in mRNA processing are also able to bind ssRNA with a preference for a GGU motif (43). In addition to its two Znf domains located at its N-terminus, ZRANB2 contains a C-terminal arginine-serine rich (RS) domain typical of splicing factors, where it facilitates both protein-protein and protein-RNA interactions. Interestingly, OZ2 shares a similar architecture with two Znf domains, albeit in the middle of the protein, followed by a glycine-rich region in its C-terminus (E-value: 4.2e-07). The C-terminus of OZ2 is also enriched in arginine and serine, although the significance is below the threshold level (E-value: 0.16 and 0.35, respectively). An X-ray crystallography of the second Znf of ZRANB2 bound to a 6-nt RNA with the sequence AGGUAA was resolved at 1.4 Å and identified six residues that directly contact the RNA (42). Three of these six residues are identical or chemically similar to the ones found in the Znf domains of OZ2, suggesting that OZ2 likely binds ssRNA (Figure 10). A notable difference concerns the two asparagines that specify uridine in the GGU motif; these have been substituted with pheny-
Figure 9. OZ2 interacts with several site-specific and general mitochondrial splicing factors in vitro and in vivo. (A) Yeast two-hybrid assay for OZ2/mitochondrial splicing factors demonstrate interactions with certain factors that share splicing targets with OZ2, as well as OZ2 dimerization. Haploid yeast were transformed with constructs expressing OZ2 or a mitochondrial splicing factor with N-terminal fusions of parts of the yeast GAL4 transcription factor. Transformed yeast were mated to produce diploids expressing two fusion proteins and selected on synthetic dropout media lacking leucine and tryptophan (-L-W). Interactions were tested on plates lacking leucine, tryptophan, histidine, and adenine (-L-W-H-Ade). Yeast dilution spots contained 10⁶, 10⁵ and 10⁴ cells per mL. Em = empty vector. (B-I) Bimolecular fluorescence complementation (BiFC) was performed in N. benthamiana using transient nYFP/cYFP fusion protein expression constructs. Green = YFP fluorescence; magenta = chlorophyll autofluorescence; scale bars = 10 μm. (B) ABO5 + OZ2. (C) BIR6 + OZ2. (D) MAT2 + OZ2. (E) mCSF1 + OZ2. (F) MISF26 + OZ2. (G) MTL1 + OZ2. (H) PMH2 + OZ2. (I) OZ2 + OZ2. The interaction of OZ2 with itself serves as a positive control for this assay.

Table 1. Summary of protein–protein interactions between OZ2 and mitochondrial splice factors. Y2H = yeast two-hybrid; BiFC = bimolecular fluorescence complementation

| Splicing factor | Target introns                | OZ2 interaction (Y2H) | OZ2 interaction (BiFC) |
|----------------|-------------------------------|-----------------------|------------------------|
| ABO5           | nad2-t3                      | Yes                   | Yes                    |
| BIR6           | nad7-t1                      | Yes                   | Yes                    |
| MAT2           | cox2, nad1-t2, nad7-t2        | No                    | Yes                    |
| mCSF1          | rps3*, nad2-t3*, nad5-t1*, -t2*, -t3*, nad7-t2* and others | No | Yes |
| MISF26         | nad2-t3                      | Yes                   | Yes                    |
| MISF74         | nad1-t4, nad2-t4             | No                    | No                     |
| mTERF15        | nad2-t3                      | No                    | No                     |
| MTL1           | nad7-t2                      | No                    | Yes                    |
| OTP439         | nad5-t3                      | No                    | No                     |
| PMH2           | nad5-t1*, -t2*, -t3*, rps3* and others | Yes | Yes |
| RUG3           | nad2-t2, nad2-t3*            | No                    | No                     |
| SLO3           | nad7-t2*                     | No                    | No                     |
| WTF9           | rpl2, ccmFc                  | No                    | No                     |

*Splice sites targeted by OZ2.

It is noteworthy that the same domain has now been implicated in both nuclear and organelle splicing. The origin of nuclear introns is still under debate; however, one theory proposes that the spliceosome and nuclear introns evolved from bacterial group II introns that invaded the eukaryotic genomes (44). Convergent evolution alone can hardly explain the numerous similarities found not only in mechanisms, two identical transesterifications resulting in lariat formation and exon junction, but also in the structures of group II and nuclear introns. In addition to the striking parallel between the function and structure of the small nuclear RNAs (snRNAs) and the group II intron RNA, some homologies are clear between Prp8, the protein core of the spliceosome, and the group II intron encoded protein (45). The discovery of OZ2 as a mitochondrial splicing factor containing a domain involved in nuclear splicing establishes another link between these two possible evolutionary related processes.
The specificity of OZ2 function in the splicing of certain introns might be mediated by interaction with other splicing factors. Eighteen splicing factors have been identified that share some common targets with OZ2; some are required for the processing of many introns like CRM-containing proteins mCSF1 and CFM9. (46,47). RNA DEAD-box helicases PHM2 and ABO6 (48,49), and matures nMAT1, nMAT2, nMAT4 and MatR (10,50–52). Other more specific splicing factors are involved in the removal of one to two introns; most of these proteins belong to the PPR family (OTP43, MSP1, ABO5, MISF26, OTP439, TANG2, SLO3, MTL1) (22,34,37,53–56). The splicing of the third intron in the nad2 transcript. Many chloroplast splicing factors have been found in high molecular weight ribonucleoprotein complexes by sucrose gradient sedimentation experiments (59–63). Similar experiments have shown that PHM2 and nMAT2 are also associated with their intron-RNA targets in large ribonucleoprotein particles in vivo (64). Our results suggest that OZ2 might also be found in these high molecular weight ribonucleoprotein complexes, particularly the one controlling the splicing of the third intron in the nad2 transcript. The localization of OZ2 as punctate foci in Arabidopsis mitochondria as seen in Figure 2A might be caused by its interaction with other splicing factors.

An inconsistency in our interaction data concerns BIR6, a protein shown to interact with OZ2 by both Y2H and BiFC assays even though it does not share a common splicing target with OZ2. BIR6 is involved in the splicing of intron 1 of the mitochondrial nad7 transcript (21). However, two other splicing events are affected in the bir6 mutants though to a much lesser extent than the splicing of nad7-intron 1, the removal of introns 1 and 2 from the nad2 transcript (Figure 4A). The interaction between BIR6 and OZ2 might thus be explained by their minor involvement in the splicing of the first intron of the nad2 transcript.

The BiFC assay revealed three additional interactions when compared to the Y2H assay. The heterologous system represented by the Y2H assay might not allow the proper folding of some of these factors preventing their interactions. In addition, we removed the transit peptide from the factors assayed in the Y2H assay based on software predictions; a possible error in the predicted cleavage sites might have led to the removal of too much or too little of the protein, impairing its proper folding.

Splicing efficiency is a metric often used to measure the effect of a splicing factor and is calculated as the ratio of spliced to unspliced forms of each transcript in the mutant normalized to the same ratio in the wild-type plant; it is generally expressed as a base 2 logarithm ratio. The reduction in splicing efficiency in mutants affected either in general factors such as CFM9 or nMAT1 or in factors having another function such as ODB1 or MTL1 is moderate and ranges from 2\(^{-6}\) to 2\(^{-10}\) (34,47,50,65). On the other hand, mutants in specific factors like PPR proteins tend to exhibit a more pronounced decrease in splicing efficiency, ranging from 2\(^{-10}\) to 2\(^{-35}\), as observed for SLO3 and OTP43, respectively (22,56). OZ2 mutant plants exhibit a marked decrease in splicing efficiency in mutants affected either in general factors such as CFM9 or nMAT1 or in factors having another function such as ODB1 or MTL1 is moderate and ranges from 2\(^{-6}\) to 2\(^{-10}\) (34,47,50,65). On the other hand, mutants in specific factors like PPR proteins tend to exhibit a more pronounced decrease in splicing efficiency, ranging from 2\(^{-10}\) to 2\(^{-35}\), as observed for SLO3 and OTP43, respectively (22,56). OZ2 mutant plants exhibit a marked decrease in splicing efficiency in mutants affected either in general factors such as CFM9 or nMAT1 or in factors having another function such as ODB1 or MTL1 is moderate and ranges from 2\(^{-6}\) to 2\(^{-10}\) (34,47,50,65). On the other hand, mutants in specific factors like PPR proteins tend to exhibit a more pronounced decrease in splicing efficiency, ranging from 2\(^{-10}\) to 2\(^{-35}\), as observed for SLO3 and OTP43, respectively (22,56). OZ2 mutant plants exhibit a marked decrease in splicing efficiency in mutants affected either in general factors such as CFM9 or nMAT1 or in factors having another function such as ODB1 or MTL1 is moderate and ranges from 2\(^{-6}\).
script. For instance, the splicing efficiency of nad1-intron 1 is reduced to 2−3 in oz2, while the other introns in nad1 transcript are not affected (Figure 3B).

Whatever the molecular mechanism by which OZ2 is involved in the splicing of mitochondrial introns, its impairment leads to a reduction in complex I assembly and activity and a subsequent morphological phenotype characteristic of complex I mutant plants (Figure 6). All the other mitochondrial complexes were similar in abundance and activity between the oz2 mutant and the wild-type plants. Complex I mutant plants have been studied in different species and originate from: 1) deletion in mitochondrial genomes, e.g. the CMS1 (cytoplasmic male sterile) and CMSII plants in Nicotiana sylvestris (66), the NCS2 (non-chromosomal stripe) mutant in maize, 2) point or T-DNA insertional mutation in nuclear genes encoding mitochondrial subunits like the fro1 (frostbite1), ndufs4 or ca2 mutants in Arabidopsis (67–69), and 3) nuclear mutants in genes encoding factors required for the maturation of mitochondrial transcripts, like NMS1 plants in Nicotiana sylvestris (70) and the otp43 and mtsf2 mutant plants in Arabidopsis (22,35). The range of phenotypes observed in these mutants varies quite dramatically both between and within species regarding embryo development and vegetative growth. In NCS2 maize mutant plants, deletion of the 3’ end of the nad4 gene causes lethality during kernel development; only heteroplasmic plants that contain normal mitochondria in addition to mutant ones can be propagated. The segregation of mutant mitochondria during development results in mutant sectors that appear as pale stripes, hence the name of the mutants (71). The Nicotiana sylvestris CMS plants exhibit delayed development and conditional male sterility depending on light intensity (66). In Arabidopsis, the ca2 mutants affected in a gene encoding a plant specific subunit of complex I with similarity to carboxylic anhydrases do not show any morphological phenotype even though Complex I activity was decreased by 80% (69). On the other hand, Arabidopsis otp43 mutant plants defective in the trans-splicing of nad1-intron 1 exhibit a total lack of complex I activity; in addition to delayed development and severe morphological phenotype, the mutant seeds require germination on half-strength MS medium supplemented with 1% sucrose (22). This mutation belongs to the comprehensive dataset of 510 cloned EMBRYO-DEFECTIVE (EMB) genes established by Meinke (13). Thus, it appears that the effect of a complex I dysfunction on embryo development in Arabidopsis depends on the severity of the reduction in the activity of this complex. Measurement by in-gel activity assay lacks the sensitivity to detect trace amounts of complex I activity. This limitation might explain the ambiguity surrounding the essential function of complex I in embryo development (76). For instance, ndufs4 mutants lacking the 18-kDa subunit of complex I that do not need sucrose to germinate were first reported to lack complex I activity (68), but in a later report were found to exhibit trace amounts of activity (72). A recent report on msp1 mutant plants similarly affected in trans-splicing of nad1-intron 1 as otp43 mutant plants further supports the essential role of complex I in embryo development (53). Like otp43, the msp1 mutation is annotated as an EMB gene (EMB1025) (13). A modified method for embryo rescue of msp1 plants had to be developed by sowing mutant seeds on MS-agar plates supplemented with 1–3% sucrose and various vitamins (53).

The oz2 mutation was first reported as the slow embryo development (sed1) mutation, which results in retarded embryogenesis and aborted seeds (12). Light microscopy data indicated that embryonic arrest occurs by approximately early globular stage in sed1 (oz2) seeds. Attempts to rescue the mutant embryos by growing them on half-strength MS medium failed as they died after 3 weeks (12). What can be the origin of the embryo defect caused by the oz2 mutation? Given the splicing deficiency in rps3 (the gene encoding the mitochondrial ribosomal protein S3) and nad transcripts observed in oz2 rescued mutants, both a defective mitochondrial translation apparatus and/or complex I could explain the embryo phenotype. NCS3 and NCS4 maize mutants present deletions of the mitochondrial rps3 and are associated with very reduced levels of mitochondrial protein synthesis (73,74). Surprisingly, the abundance of Complexes III, IV and V was not reduced in oz2 rescued mutant plants (Figure 6 and Supplementary Figure S5) even though they contain mitochondrial-encoded subunits, suggesting that the reduced rps3 mature transcript does not impair mitochondrial protein synthesis. However, as noted earlier, expression under the control of the ABI3 promoter is not totally turned off during seedling development, the stage at which the RNA was extracted from oz2 rescued plants. It is possible that the amount of rps3 spliced product is more drastically reduced during the embryo development of oz2 mutant, thus contributing to its arrest. The cumulative effect of splicing deficiencies in several nad transcripts probably explains the observation of a reduced complex I abundance and activity in oz2 rescued mutant plants. Like for rps3, the decrease in nad1, 2, 5 and 7 mature transcripts, and hence in complex I, is likely to be more severe in an oz2 mutant that has not been rescued by the ABI3 promoter.

Like many mutants affected in complex I biogenesis, oz2 rescued mutants show an induction of alternative respiratory pathway genes, especially AOX1d whose transcript steady state level was ~270-fold higher than in the wild-type plants (Figure 7). This observation is reminiscent of the strong induction of AOX1d at the transcript and protein level in a double mutant impaired in both the cytochrome c oxidase pathway and the AOX pathway (aox1a:rhoTmp) (75). The induction of AOX1d could not compensate for the absence of AOX1a, as the double mutant exhibited a more severe growth retardation than the single rhoTmp mutant suggesting differential activity of AOX isoforms. A differential activity of AOX isoforms was later substantiated by a differential response to tricarboxylic acid cycle intermediates (76). Another feature shared by oz2 rescued mutant plants and complex I mutants is the upregulation of steady state levels of mitochondrial transcripts, which may correspond to compensatory effects due to the altered organelar functions (Supplementary Figure S3), e.g. (50,51,54,58).

In this work, we have investigated the function of OZ2 and discovered its involvement in mitochondrial splicing. Another OZ family member, OZ1, was previously identified to be a chloroplast editing factor (77). The remaining members of this family, OZ3 and OZ4, both predicted to be localized either in the mitochondrion or in the plastid (30)
have not yet been assigned any function, though it seems likely that they will be found to have a role in organelle RNA metabolism.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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REFERENCES
1. Archibald,J.M. (2015) Endosymbiosis and eukaryotic cell evolution. Curr. Biol., 25, R911–R921.
2. Timmis,J.N., Ayliffe,M.A., Huang,C.Y. and Martin,W. (2004) Endosymbiotic gene transfer: organelle genomes forge eukaryotic chromosomes. Nat. Rev. Genet., 5, 123–135.
3. Hammami,K. and Giege,P. (2014) RNA metabolism in plant mitochondria. Trends Plant Sci., 19, 380–389.
4. Barkan,A. (2011) Expression of plastid genes: organelle-specific elaborations on a prokaryotic scaffold. Plant Physiol., 155, 1520–1532.
5. Bonen,L. (2008) Cis- and trans-splicing of group II introns in plant mitochondria. Mitochondrion, 8, 26–34.
6. de Longevialle,A.F., Small,I.D. and Lurin,C. (2010) Nuclearly encoded splicing factors implicated in RNA splicing in higher plant organelles. Mol Plant, 3, 691–701.
7. Kim,S. and Yoon,M.K. (2010) Comparison of mitochondrial and chloroplast genome segments from three onion (Allium cepa L.) cytoplasm types and identification of a trans-splicing intron of cox2. Curr. Genet., 56, 177–188.
8. Chapdelaine,Y. and Bonen,L. (1991) The wheat mitochondrial gene for subunit I of the NADH dehydrogenase complex: a trans-splicing model for this gene-in-pieces. Cell, 65, 465–472.
9. Conklin,P.L., Wilson,R.K. and Hanson,M. (1991) Multiple trans-splicing events are required to produce a mature nad1 transcript in a plant mitochondrion. Genes Dev., 5, 1407–1415.
10. Sultmann,H., Mileshina,D., Grewe,F., Rolle,K., Abudraham,S., Giege,P., Niazi,A.K., Keren,I., Shevtsov,S., Klipcan,L. et al. (2016) The reverse transcriptase/RNA maturase protein MatR is required for the splicing of various group ii introns in brassicaceae mitochondria. Plant Cell, 28, 2805–2829.
11. Brown,G.G., Colas des Frans-Small,C. and Osterterz-Biran,O. (2014) Group II intron splicing factors in plant mitochondria. Front Plant Sci., 5, 35.
12. Ju,Y., Liu,C., Lu,W., Zhang,Q. and Sodmergen (2016) Arabidopsis mitochondrial protein slow embryo development1 is essential for embryo development. Biochem. Biophys. Res. Commun., 474, 371–376.
13. Meinke,D.W. (2020) Genome-wide identification of EMBRYO-DEFECTIVE (EMB) genes required for growth and development in Arabidopsis. New Phytol., 226, 306–325.
14. Després,B., Delseny,M. and Devic,M. (2001) Partial complementation of embryo defective mutations: a general strategy to elucidate gene function. Plant J., 27, 149–159.
15. Earley,K.W., Haag,J.R., Pontes,O., Opper,K., Juehne,T., Song,K. and Pikaard,C.S. (2006) Gateway-compatible vectors for plant functional genomics. Plant J., 45, 616.
16. Zhang,X., Henriques,R., Lin,S.S., Niu,Q.W. and Chua,N.H. (2006) Agrobacterium-mediated transformation of Arabidopsis thaliana using the floral dip method. Nat. Protoc., 1, 641.
17. Jakoby,M.J., Weinl,C., Pusch,S., Kujit,S.J., Merkle,T., Dissmeyer,N. and Schnittger,A. (2006) Analysis of the subcellular localization, function, and proteolytic control of the Arabidopsis cyclin-dependent kinase inhibitor IKK1/1RPI. Plant Physiol., 141, 1293–1305.
18. Yoo,S.D., Cho,Y.H. and Sheen,J. (2007) Arabidopsis mesophyll proplasts: a versatile cell system for transient gene expression analysis. Nat. Protoc., 2, 1565.
19. Bentolila,S., Elliott,L.E. and Hanson,M.R. (2008) Genetic architecture of mitochondrial editing in Arabidopsis thaliana. Genetics, 178, 1693.
20. Shi,X., Hanson,M.R. and Bentolila,S. (2015) Two RNA recognition motif-containing proteins are plant mitochondrial editing factors. Nucleic Acids Res., 43, 3874–3875.
21. Koprivova,A., des Francs-Small,C.C., Calder,G., Mugford,S.T., Tanz,S., Lee,B.R., Zechmann,B., Small,l. and Kopriva,S. (2010) Identification of a pentatricopeptide repeat protein implicated in splicing of intron 1 of mitochondrial nad1 transcripts. J. Biol. Chem., 285, 32192–32199.
22. de Longevialle,A.F., Meyer,E.H., Andres,C., Taylor,N.L., Linur,C., Millar,A.H. and Small,I.D. (2007) The pentatricopeptide repeat gene OTP43 is required for trans-splicing of the mitochondrial nad1 Intron 1 in Arabidopsis thaliana. Plant Cell, 19, 3256.
23. Umbach,A.L., Fiorani,F. and Siedow,J.N. (2005) Characterization of transformed Arabidopsis with altered alternative oxidase levels and analysis of effects on reactive oxygen species in tissue. Plant Physiol., 139, 1806–1820.
24. Lu,K., Li,T., He,J., Chang,W., Zhang,R., Liu,M., Yu,M., Fan,Y., Ma,J., Sun,W. et al. (2016) q PrimerDB: a thermodynamics-based gene-specific qPCR primer database for 147 organisms. Nucleic Acids Res., 45, D1239–D1236.
25. Bentolila,S., Knight,W. and Hanson,M. (2010) Natural variation in Arabidopsis leads to the identification of REME1, a pentatricopeptide repeat-DYW protein controlling the editing of mitochondrial transcripts. Plant Physiol., 154, 1966–1982.
26. Murcha,M.W. and Whelan,J. (2015) Isoforms of the model plant species Arabidopsis thaliana and Orzya sativa. Methods Mol. Biol., 1305, 1–12.
27. Schertl,P. and Braun,H.P. (2015) Activity measurements of mitochondrial enzymes in native gels. Methods Mol. Biol., 1305, 131–138.
28. Hayes,M.L., Giang,K., Berhane,B. and Mulligan,R.M. (2013) Identification of two pentatricopeptide repeat genes required for RNA editing and zinc binding by C-terminal cystidine deaminase-like domains. J. Biol. Chem., 288, 36519–36529.
29. Lin,M.T., Occhialini,A., Andralojc,P., J.Devonshire,J., Hines,K.M., Parry,M.A. and Hanson,M.R. (2014) β-Carboxysomal proteins assemble into highly organized structures in Nicotiana chloroplasts. Plant J., 79, 1–12.
30. Gipson,A.B., Giletto,A., L., Hanson,M.R. and Bentolila,S. (2020) Arabidopsis RanBP2-Type zin finger proteins related to chloroplast RNA editing factor OZ1. Plants (Basel), 9, 307.
31. Hooper,C.M., Castleden,I.R., Tanz,S.K., Aryamanesh,N. and Millar,A.H. (2017) SUBA4: the interactive data analysis centre for Arabidopsis subcellular protein locations. Nucleic Acids Res., 45, D1064–D1074.
32. Small,I., Peeters,N., Legeai,F. and Linur,C. (2004) Predator: a tool for rapidly screening proteomes for N-terminal targeting sequences. Proteomics, 4, 1581.
33. Malek,O. and Knoop,V. (1998) Trans-splicing group II introns in plant mitochondria: the complete set of cis-arranged homologs in ferns, fern allies, and a hornwort. RNA, 4, 1599–1609.
34. Halli,N., Planchard,N., Arnal,N., Quadrado,M., Vrielynck,N., Dahan,J., des Francs-Small,C. and Mireau,H. (2016) The MT1l pentatricopeptide repeat protein is required for both translation and splicing of the mitochondrial NADH DEHYDROGENASE SUBUNIT7 mRNA in arabidopsis. Plant Physiol., 170, 354–366.
35. Wang,C., Aube,F., Planchard,N., Quadrado,M., Dargel-Graffin,C., Nogue,F. and Mireau,H. (2017) The pentatricopeptide repeat protein MT5F2 stabilizes a nad1 precursor transcript and defines the 3 end of its 5′-half intron. Nucleic Acids Res., 45, 6119–6134.
36. Halli,N., Arnal,N., Quadrado,M., Amiar,S., Tcherkez,G., Dahan,J., Briozzo,P., Colas des Francs-Small,C., Vrielynck,N. and Mireau,H. (2013) The pentatricopeptide repeat MT5F1 protein stabilizes the
40. Higa, M.M., Alam, S.L., Sundquist, W.I., and Ullman, K.S. (2007) The molecular characterization of the Ran-binding zinc finger domain of Nup153. J. Biol. Chem., 282, 17090–17100.

41. Adams, D.J., van der Weyden, L., Mayeda, A., Stamm, S., Morris, B.J., Singh, B.B., Patel, H.H., Roepman, R., Schick, D., and Ferreira, P.A. (1999) The zinc finger cluster domain of RanBP2 is a specific docking site for the nuclear export factor, exportin-1. J. Biol. Chem., 274, 37370–37378.

42. Higa, M.M., Alam, S.L., Sundquist, W.I., and Ullman, K.S. (2007) Molecular characterization of the Ran-binding zinc finger domain of Nup153. J. Biol. Chem., 282, 17090–17100.

43. Adams, D.J., van der Weyden, L., Mayeda, A., Stamm, S., Morris, B.J., Singh, B.B., Patel, H.H., Roepman, R., Schick, D., and Ferreira, P.A. (1999) The zinc finger cluster domain of RanBP2 is a specific docking site for the nuclear export factor, exportin-1. J. Biol. Chem., 274, 37370–37378.

44. Adams, D.J., van der Weyden, L., Mayeda, A., Stamm, S., Morris, B.J., Singh, B.B., Patel, H.H., Roepman, R., Schick, D., and Ferreira, P.A. (1999) The zinc finger cluster domain of RanBP2 is a specific docking site for the nuclear export factor, exportin-1. J. Biol. Chem., 274, 37370–37378.

45. Adams, D.J., van der Weyden, L., Mayeda, A., Stamm, S., Morris, B.J., Singh, B.B., Patel, H.H., Roepman, R., Schick, D., and Ferreira, P.A. (1999) The zinc finger cluster domain of RanBP2 is a specific docking site for the nuclear export factor, exportin-1. J. Biol. Chem., 274, 37370–37378.

46. Adams, D.J., van der Weyden, L., Mayeda, A., Stamm, S., Morris, B.J., Singh, B.B., Patel, H.H., Roepman, R., Schick, D., and Ferreira, P.A. (1999) The zinc finger cluster domain of RanBP2 is a specific docking site for the nuclear export factor, exportin-1. J. Biol. Chem., 274, 37370–37378.

47. Adams, D.J., van der Weyden, L., Mayeda, A., Stamm, S., Morris, B.J., Singh, B.B., Patel, H.H., Roepman, R., Schick, D., and Ferreira, P.A. (1999) The zinc finger cluster domain of RanBP2 is a specific docking site for the nuclear export factor, exportin-1. J. Biol. Chem., 274, 37370–37378.

48. Kohler, D., Schmidt-Gattung, S. and Binder, S. (2010) The DEAD-box protein PMH2 is required for efficient group II intron splicing in Arabidopsis. Plant J., 67, 1067–1080.

49. Keren, I., Tal, L., des Francs-Small, C.C., Zhang, B., Murcha, M.W. and Whelan, J. (2011) The RCC1 family protein RUG3 is required for the splicing of nad2 and complex I biogenesis in Arabidopsis thaliana. Plant J., 67, 1067–1080.

50. Keren, I., Bezawork-Geleta, A., Kolton, M., Maayan, I., Belausov, E., Small, I. and Ostersetzer-Biran, O. (2013) mCSF1, a new RNA-binding protein, facilitates the trans-splicing of the maize chloroplast rps12 pre-mRNA. Plant Cell, 18, 2650.

51. Keren, I., Tal, L., des Francs-Small, C.C., Zhang, B., Murcha, M.W. and Whelan, J. (2011) The RCC1 family protein RUG3 is required for the splicing of nad2 and complex I biogenesis in Arabidopsis thaliana. Plant J., 67, 1067–1080.

52. Keren, I., Bezawork-Geleta, A., Kolton, M., Maayan, I., Belausov, E., Small, I. and Ostersetzer-Biran, O. (2013) mCSF1, a new RNA-binding protein, facilitates the trans-splicing of the maize chloroplast rps12 pre-mRNA. Plant Cell, 18, 2650.

53. Keren, I., Tal, L., des Francs-Small, C.C., Zhang, B., Murcha, M.W. and Whelan, J. (2011) The RCC1 family protein RUG3 is required for the splicing of nad2 and complex I biogenesis in Arabidopsis thaliana. Plant J., 67, 1067–1080.

54. Keren, I., Tal, L., des Francs-Small, C.C., Zhang, B., Murcha, M.W. and Whelan, J. (2011) The RCC1 family protein RUG3 is required for the splicing of nad2 and complex I biogenesis in Arabidopsis thaliana. Plant J., 67, 1067–1080.

55. Liu, Y., He, J., Chen, Z., Ren, X., Hong, X. and Gong, Z. (2010) ABA over-sensitive 5 (ABOS), encoding a pentatricopeptide repeat protein required for cis-splicing of mitochondrial nad2 intron 3, is involved in the abscisic acid response in Arabidopsis. Plant J., 63, 749–765.

56. Wang, C., Aube, F., Quadroado, M., Dargel-Graffin, C. and Mireau, H. (2018) Three new pentatricopeptide repeat proteins facilitate the splicing of mitochondrial transcripts and complex I biogenesis in Arabidopsis. J. Exp. Bot., 69, 3131–3140.

57. Huawei, W., Liao, J.C., Chang, C.Y., Harrison, T., Boucher, C. and Hu, H.M. (2015) The SLOW GROWTh1 pentatricopeptide repeat protein is required for the splicing of mitochondrial NADH dehydrogenase Subunit2 intron 2 in Arabidopsis. Plant Physiol., 168, 490–501.

58. Hsu, Y.W., Wang, H.J., Hsieh, M.H., Hsieh, H.L. and Jauh, G.Y. (2014) Arabidopsis mTERF15 is required for mitochondrial nad2 intron 3 splicing and functional complex I activity. PLoS One, 9, e112360.

59. Kuhn, K., Carr, C., Giraud, E., Wang, Y., Meyer, E.H., Narsai, R., des Francs-Small, C.C., Zhang, B., Murcha, M.W. and Whelan, J. (2011) The RCC1 family protein RUG3 is required for the splicing of nad2 and complex I biogenesis in Arabidopsis thaliana. Plant J., 67, 1067–1080.

60. Kohler, D., Schmidt-Gattung, S. and Binder, S. (2010) The DEAD-box protein PMH2 is required for efficient group II intron splicing in Arabidopsis. New Phytol., 189, 755–767.

61. Kohler, D., Schmidt-Gattung, S. and Binder, S. (2010) The DEAD-box protein PMH2 is required for efficient group II intron splicing in Arabidopsis. New Phytol., 189, 755–767.

62. Kohler, D., Schmidt-Gattung, S. and Binder, S. (2010) The DEAD-box protein PMH2 is required for efficient group II intron splicing in Arabidopsis. New Phytol., 189, 755–767.

63. Kohler, D., Schmidt-Gattung, S. and Binder, S. (2010) The DEAD-box protein PMH2 is required for efficient group II intron splicing in Arabidopsis. New Phytol., 189, 755–767.

64. Kohler, D., Schmidt-Gattung, S. and Binder, S. (2010) The DEAD-box protein PMH2 is required for efficient group II intron splicing in Arabidopsis. New Phytol., 189, 755–767.

65. Kohler, D., Schmidt-Gattung, S. and Binder, S. (2010) The DEAD-box protein PMH2 is required for efficient group II intron splicing in Arabidopsis. New Phytol., 189, 755–767.

66. Kohler, D., Schmidt-Gattung, S. and Binder, S. (2010) The DEAD-box protein PMH2 is required for efficient group II intron splicing in Arabidopsis. New Phytol., 189, 755–767.

67. Kohler, D., Schmidt-Gattung, S. and Binder, S. (2010) The DEAD-box protein PMH2 is required for efficient group II intron splicing in Arabidopsis. New Phytol., 189, 755–767.
71. Karpova, O.V. and Newton, K.J. (1999) A partially assembled complex I in NAD4-deficient mitochondria of maize. *Plant J.*, 17, 511–521.

72. Kuhn, K., Obata, T., Feher, K., Bock, R., Fernie, A.R. and Meyer, E.H. (2015) Complete mitochondrial complex I deficiency induces an up-regulation of respiratory fluxes that is abolished by traces of functional complex I. *Plant Physiol.*, 168, 1537–1549.

73. Hunt, M.D. and Newton, K.J. (1991) The NCS3 mutation: genetic evidence for the expression of ribosomal protein genes in *Zea mays* mitochondria. *EMBO J.*, 10, 1045.

74. Newton, K.J., Mariano, J.M., Gibson, C.M., Kuzmin, E. and Gabay-Laughnan, S. (1996) Involvement of S2 episomal sequences in the generation of NCS4 deletion mutation in maize mitochondria. *Dev. Genet.*, 19, 277–286.

75. Kuhn, K., Yin, G., Duncan, O., Law, S.R., Kubiszewski-Iakubiak, S., Kaur, P., Meyer, E., Wang, Y., Small, C.C., Giraud, E. *et al.* (2015) Decreasing electron flux through the cytochrome and/or alternative respiratory pathways triggers common and distinct cellular responses dependent on growth conditions. *Plant Physiol.*, 167, 228–250.

76. Selinski, J., Hartmann, A., Deckers-Hebestreit, G., Day, D.A., Whelan, J. and Scheibe, R. (2018) Alternative oxidase isoforms are differentially activated by tricarboxylic acid cycle intermediates. *Plant Physiol.*, 176, 1423–1432.

77. Sun, T., Shi, X., Friso, G., Van Wijk, K., Bentolila, S. and Hanson, M.R. (2015) A zinc finger motif-containing protein is essential for chloroplast RNA editing. *PLoS Genet.*, 11, e1005028.