Chloroplast RH3 DEAD Box RNA Helicases in Maize and Arabidopsis Function in Splicing of Specific Group II Introns and Affect Chloroplast Ribosome Biogenesis

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Chloroplasts in angiosperms contain at least seven nucleus-encoded members of the DEAD box RNA helicase family. Phylogenetic analysis shows that five of these plastid members (RH22, -39, -47, -50, and -58) form a single clade and that RH3 forms a clade with two mitochondrial RH proteins (PMH1 and -2) functioning in intron splicing. The function of chloroplast RH3 in maize (Zea mays; ZmRH3) and Arabidopsis (Arabidopsis thaliana; AtRH3) was determined. ZmRH3 and AtRH3 are both under strong developmental control, and ZmRH3 abundance sharply peaked in the sink-source transition zone of developing maize leaves, coincident with the plastid biogenesis machinery. ZmRH3 coimmunoprecipitated with a specific set of plastid RNAs, including several group II introns, as well as pre23S and 23S ribosomal RNA (rRNA), but not 16S rRNA. Furthermore, ZmRH3 associated with 50S preribosome particles as well as nucleoids. AtRH3 null mutants are embryo lethal, whereas a weak allele (rh3-4) results in pale-green seedlings with defects in splicing of several group II introns and rRNA maturation as well as reduced levels of assembled ribosomes. These results provide strong evidence that RH3 functions in the splicing of group II introns and possibly also contributes to the assembly of the 50S ribosomal particle. Previously, we observed 5- to 10-fold up-regulation of AtRH3 in plastid Caseinolytic protease mutants. The results shown here indicate that AtRH3 up-regulation was not a direct consequence of reduced proteolysis but constituted a compensatory response at both RH3 transcript and protein levels to impaired chloroplast biogenesis; this response demonstrates that cross talk between the chloroplast and the nucleus is used to regulate RH3 levels.

DEAD box RNA helicases are mainly involved in ATP-dependent rearrangement of intermolecular and intramolecular RNA structures or remodeling of ribonucleoprotein complexes and are found in all eukaryotes and most prokaryotes (Cordin et al., 2006; Banroques et al., 2011; Linder and Jankowsky, 2011). DEAD box RNA helicases in Arabidopsis chloroplasts, namely RH3, RH22, RH26, RH39, RH47, and RH50. These plastid RNA helicases have been shown to be involved in RNA synthesis, modification, cleavage, and degradation as well as in ribosome biogenesis and translation initiation (Banroques et al., 2011; Linder and Jankowsky, 2011). DEAD box RNA helicase proteins possess conserved ATP-binding and -hydrolyzing domains, RNA-binding motifs, and a characteristic Asp-Glu-Ala-Asp (DEAD) sequence (as part of the Walker B motif; Cordin et al., 2006). However, in some cases, the Ala in the DEAD domain is not conserved (resulting into DExD). These conserved domains are clustered in the central core region that spans 350 to 400 amino acid residues. The N- and C-terminal extensions are highly variable and are thought to provide additional interactions with substrates or cofactors, or to confer additional activities such as protease activity (for NS3, see Cordin et al., 2006). DEAD box RNA helicase proteins belong to the helicase superfamily 2, which also includes DEAH and DExH proteins.

It was reported previously that there are 56 predicted DEAD box RNA helicases (and two pseudogenes) in Arabidopsis (Arabidopsis thaliana; Mingam et al., 2004), 11 of which have a TargetP-predicted chloroplast transit peptide. In this study and in the Plant Proteome Database (http://ppdb.tc.cornell.edu/), we adopt the numbering for the RNA helicases as established for Arabidopsis (Mingam et al., 2004). Mass spectrometry analyses identified six DEAD box helicases in Arabidopsis chloroplasts, namely RH3, RH22, RH26, RH39, RH47, and RH50. These plastid helicases are involved in splicing of several group II introns and affect chloroplast ribosome biogenesis.

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proteins were detected in high-molecular-mass (greater than 0.6–3 MD) chloroplast stromal complexes (Peltier et al., 2006; Olinares et al., 2010). Maize (Zea mays) homologs for these six plastid helicases were found by proteome analysis of maize plastids (Friso et al., 2010; Majeran et al., 2012). Most of these maize proteins were enriched in plastid nucleoids (Majeran et al., 2012) and also in Arabidopsis nucleoids (G. Friso, K. Nishimura, X. Qu, and K.J. van Wijk, unpublished data). The tobacco (Nicotiana tabacum) homolog of RH58, named VDL helicase, was identified in tobacco chloroplasts (Wang et al., 2000) but was not (yet) detected in Arabidopsis, likely because of its low abundance. The maize homolog for RH58 was found in a coimmunoprecipitation (co-IP) with anti-WHY1 serum against maize nucleoids (Majeran et al., 2012), but RH58 abundance levels were very low compared with the other plastid RH helicases.

Mutants for several chloroplast RNA helicases have been characterized to various degrees and exhibit delayed or impaired chloroplast and leaf development and defects in chloroplast RNA metabolism. OsBRH1, the rice (Oryza sativa) homolog of Arabidopsis RH50, exhibited RNA helicase activities in vitro and helped to confer plant resistance against various stresses (Li et al., 2008). Arabidopsis mutant nara12-1, with reduced accumulation of RH39, showed strongly reduced levels of the photosynthesis machinery due to inefficient translation elongation, most likely due to the loss of RH39-mediated defects in 23S rRNA maturation (formation of the hidden break) affecting the functional state of the ribosome (Nishimura et al., 2010). Complete loss of Arabidopsis RH22 resulted into embryo lethality, whereas reduced RH22 expression resulted in virescent seedlings in which precursors of 23S and 4.5S rRNA (in the 50S particle) accumulated (Chi et al., 2012). It was concluded that RH22 facilitates the assembly of the 50S ribosomal subunit by interacting with ribosomal protein RPL24 and with the region of 23S rRNA encompassing the RPL24 binding site (Chi et al., 2012). Tobacco mutants with loss of the RH58 homolog (VDL helicase) showed pale-green and variegated leaves and abnormal roots and flowers, but no function was determined (Wang et al., 2000).

In this study, we focus on the molecular function of the chloroplast-localized DEAD box RNA helicase AtRH3 in Arabidopsis and its co-orthologs in maize, ZmRH3A and ZmRH3B. Proteome analysis showed that RH3 is the most abundant plastid DEAD box RNA helicase in maize and Arabidopsis (Zybailov et al., 2008; Friso et al., 2010). Our attention was drawn to RH3 for two reasons. First, we found that AtRH3 was strongly (more than 5-fold) up-regulated in the mutants clpr1-1 and clpr4-1, with defects in the chloroplast stroma-localized Caseinolytic protease (Clp) protease core complex (Rudella et al., 2006; Kim et al., 2009; Zybailov et al., 2009). These clpr2-1 and clpr4-1 mutants are chlorophyll-deficient and developmentally delayed mutants and have partial defects in chloroplast rRNA processing (Rudella et al., 2006; Kim et al., 2009; Zybailov et al., 2009), as observed for the Clp core mutant clpr1-1 (Koussevitzky et al., 2007). Second, we detected ZmRH3 in co-IP with several proteins that promote the splicing of chloroplast group II introns (G. Friso, K.P. Watkins, A. Barkan, and K.J. van Wijk, unpublished data). Here, we address the function of RH3 in both maize and Arabidopsis as well as the basis for its up-regulation in Arabidopsis plastid Clp protease mutants. Our results show that RH3 is directly involved in chloroplast intron splicing and possibly also in 50S ribosome biogenesis. RH3 levels peak during early stages of chloroplast biogenesis, consistent with a role in establishing the chloroplast gene expression machinery. Our results suggest that RH3 overaccumulates in Clp mutants not because it is a direct substrate of Clp protease but rather as a consequence of a disruption in chloroplast biogenesis or protein homeostasis. Finally, we use phylogeny and functional domain analysis of the Arabidopsis DEAD box RNA helicase family to compare RH3 with other plastid DEAD box RNA helicases.

RESULTS

Phylogenetic Analysis of Plant DEAD Box RNA Helicases and Architecture of RH3 Orthologs

To establish the phylogenetic relationships among maize, rice, and Arabidopsis DEAD box proteins, we created unrooted phylogenetic trees based on the 56 Arabidopsis DEAD box helicases listed by Mingam et al. (2004), 46 related maize and 47 rice sequences, as well as the five known DEAD box RNA helicases from Escherichia coli (for accession numbers, annotations, and clade assignments, see Supplemental Table S1; for all amino acid sequences, see Supplemental Text S1). If there was more than one protein model per gene, we selected the longest protein. Phylogenetic trees were built from alignments based on full-length sequences as well as after removal of the variable N- and C-terminal extensions and removing gaps within the central conserved core helicase region. Sixteen clades could be distinguished for the plant proteins (using a minimal bootstrap value of 50 to define clades), with proteins for each of the three plant species represented in each clade (Supplemental Fig. S1; Supplemental Table S1). The relationship between protein accessions inferred from the various trees was not affected by the removal of gaps and extensions, indicating robustness of these relationships.

The cladogram showed that AtRH3 (At5g26742) has two co-orthologs in maize, ZmRH3A (GRMZM2G145491_P01) and ZmRH3B (GRMZM2G163072_P01), and one rice ortholog (Os03g61220; Fig. 1A; Supplemental Fig. S1). Furthermore, these RH3 orthologs formed a distinct clade (clade 7) with the closely related pair RH9/RH33 and the more distant RH7 (Fig. 1A; Supplemental Fig. S1). AtRH9 (At3G22310) and AtRH33 (AT3G22330) are mitochondrial proteins (also named PMH1 and PMH2) and were shown to be part of a large complex in the mitochondrial matrix (Matthes et al., 2007).
Moreover, PMH2 was shown to be involved in group II intron splicing in mitochondria (Köhler et al., 2010). In the same clade, but more distant, was AtRH7, a demonstrated nuclear protein named PRH75 but with unknown function (Lorković et al., 1997). Furthermore, five of the seven known plastid proteins (RH22, RH39, RH47, RH50, and RH58) formed one clade (clade 8), suggesting a common ancestry and likely functions different from RH3 (Supplemental Fig. S1).

To understand the features that are unique to RH3 as compared with other plastid RH proteins, we analyzed the conservation of domains of all 56 Arabidopsis DEAD box RNA helicases and compared that with the conservation in the RH3 clade (Supplemental Table S1). Figure 1B shows the conserved motifs (greater than 60% identity) across all combined maize, rice, and Arabidopsis DEAD box RNA helicase families. The top lines show the conserved motifs (60% minimal conservation threshold) across the 56 Arabidopsis, 46 maize, and 47 rice DEAD box RNA helicases. The bottom lines show the conserved motifs (greater than 90% identity) for the RH3 clade.

**Figure 1.** Phylogenetic and protein domain analyses of the DEAD box RNA helicase family. A, The RH3-containing clade (clade 7) from the phylogenetic tree of all 149 DEAD box helicases in maize, rice, and Arabidopsis and the five *E. coli* DEAD box RNA helicases (see Supplemental Fig. S1). Bootstrap values are indicated. B, Conserved domains of the maize, rice, and Arabidopsis DEAD box RNA helicase families. The top lines show the conserved motifs (60% minimal conservation threshold) across the 56 Arabidopsis, 46 maize, and 47 rice DEAD box RNA helicases. The bottom lines show the conserved motifs (greater than 90% identity) for the RH3 clade.

Plastid RH3 Functions

**RH3 Accumulates in Stroma and Nucleoids of Green Tissues, with Peak Accumulation during Chloroplast Biogenesis**

To examine the temporal and organ expression patterns of AtRH3, we used an antibody raised to a conserved region of ZmRH3 (Fig. 2A and Supplemental Fig. S2). AtRH3 was found to be predominantly expressed in young leaves, peaking in 2-week-old seedlings and decreasing to very low levels in older leaf tissue (Fig. 2B). In addition, the AtRH3 protein was also detected in...
flowers and siliques, but not in stems or roots, of 6-week-old Arabidopsis plants. Mass spectrometry-based analysis of isolated nucleoids and chloroplasts showed a very strong nucleoid enrichment of ZmRH3 (Majeran et al., 2012) as well as AtRH3 (G. Friso, K. Nishimura, X. Qu, and K.J. van Wijk, unpublished data). Both ZmRH3 homologs were detected by mass spectrometry along the development all leaf gradient to expand maize leaves, with expression peaking in the sink-source transition zone (Fig. 2C), together with many other proteins involved in plastid gene expression and protein homeostasis (Majeran et al., 2010), including the nucleoid proteins. In initial accumulation profiles of chloroplast, ribosomal subunits followed ZmRH3 and nucleoids, but accumulation levels of ribosomes remained high and constant in mature tissue, unlike RH3 and (other) nucleoid proteins (Fig. 2C). This is consistent with their respective functions. Western-blot analysis showed that ZmRH3 and DNA/RNA-binding protein ZmWHY1 were present in chloroplast stroma as well as thylakoid membranes, likely reflecting nucleoids associated to the thylakoids (Fig. 2D). Indeed, western blots of nucleoids extracted from the thylakoid membranes identified both ZmRH3 and ZmWHY1 (Fig. 2D), in agreement with previous mass spectrometry analysis of nucleoids (Majeran et al., 2012). We note that the total amount of nucleoid proteins used for the western blot was lower than the other chloroplast fractions, thus underestimating the nucleoid enrichment for ZmRH3 and ZmWHY1. In contrast, the abundant CPN60α chaperone was only found in the stromal fraction (Fig. 2D).

ZmRH3 Associates with Group II Introns and Pre-50S Ribosomal Particles in Vivo

To determine the distribution of the particles with which ZmRH3 associates, maize chloroplast stroma was fractionated by Suc gradient sedimentation under conditions that dissociate 70S ribosomes into 30S and 50S ribosome particles. Immunoblots and RNA staining showed that RH3 primarily accumulated in particles sedimenting between 30S and 50S ribosomal subunits (1–2 MD), consistent with an association with partially assembled 50S ribosomal particles (Fig. 3A).

To identify chloroplast RNAs that associate with RH3 in vivo, RH3 was coimmunoprecipitated with ZmRH3 from maize chloroplasts. The blots stained with Ponceau S is shown in the bottom panel. CPN60α and D1/D2 were used as markers for the stroma and thylakoids, respectively. WHY1 is the control that localizes to stroma, thylakoids, and nucleoids.

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maize chloroplast extracts. RNAs in the co-IP pellets were identified by hybridization to a tilling microarray of the maize chloroplast genome. Figure 3B shows the degree to which each chloroplast RNA sequence was enriched in the RH3 co-IP pellets, plotted according to chromosomal position, using the well-established RNA immunoprecipitation (RIP)-chip procedure as described by Barkan (2009). RNAs from five loci, rps12, rpl2, trnA, trnI, and rmn23, were strongly enriched in the RH3 co-IP pellet in comparison with a negative control assay. These loci encode two kinds of large and highly structured RNAs: group II introns in the rpl2, trnA, trnI, and rps12 loci and the 23S rRNA (rmn23). A minor enrichment of RNA from the ycf3 locus, which includes two group II introns, was also suggested by these data.

The rmn23, trnI, and trnA genes are cotranscribed together with rrm16 in the chloroplast rnm operon. The primary transcript is processed via cleaving between coding regions, removal of group II introns in trnI and trnA, and trimming of the termini once ribosome assembly is complete (Stern et al., 2010). Suc gradient fractionation (Fig. 3A) suggested that ZmRH3 is associated primarily with assembling 50S ribosomal subunits (containing incompletely processed 23S rRNA) rather than with mature 50S subunits. To address that possibility, we used poisoned-primer extension to analyze the processing status of the 5′ end of the 23S rRNA in the RH3 co-IP pellets (Fig. 3C). This showed a strong enrichment for incompletely processed 23S rRNA in the RH3 co-IP pellet compared with the supernatant (Fig. 3C). These results, in conjunction with the Suc gradient sedimentation data, suggest that RH3 associates with assembling 50S ribosomal subunits and, potentially, with mature 50S ribosomal particles as well. The coprecipitation with assembling 50S ribosomal subunits could be direct, or it could be due to tethering to as-yet-unprocessed trnI and trnA, whose introns are associated with RH3. However, the cosedimentation of RH3 with pre-50S subunits favors a direct association, as group II intron particles analyzed to date are considerably smaller (approximately 700 kD) than those observed here (Till et al., 2001; Asakura and Barkan, 2007).

AtRH3 Is Required for Embryogenesis, Chloroplast Development, and Photosynthesis

To further assess how RH3 affects the RNAs with which it associates, three T-DNA insertion lines in the single RH3 gene in Arabidopsis were obtained from

![Figure 3](image-url)

**Figure 3.** RH3 cosediments with pre-50S ribosomal subunits and interacts with specific plastid RNA. A, Maize stroma were sedimented through Suc gradients under the condition that dissociates 30S and 50S subunits. An equal proportion of each fraction was analyzed by probing an immunoblot with anti-RH3 antiserum (top panel). The same blot stained with Ponceau S is shown in the bottom panel and also visualizes the large subunit (RBCL) of the Rubisco holocomplex at 550 kD. RNA extracted from gradients was analyzed by RNA gel blotting and methylene blue staining (middle panel). 16S rRNA marks 30S ribosomal subunits, and 23S rRNA marks 50S ribosomal subunits. B, Identification of RNA ligands of RH3 using the co-IP assay. The RIP-chip assay revealed the enrichment of several RNAs (as indicated) in the RH3 co-IP pellets. C, RH3 associates with the precursor of 23S rRNA determined by poisoned-primer extension assays. Reverse transcriptase reactions were initiated with a radiolabeled primer complementary to the mature and precursor forms of 23S rRNA; a dideoxy nucleotide that terminates the reverse transcription after different distances on mature and precursor forms of 23S was included in the reactions. Pre 23S, Precursor form of 23S rRNA. To, Total; sup, supernatant; ppt, pellet.
the SALK and SAIL T-DNA insertion collections (Sessions et al., 2002; Alonso et al., 2003). We verified the positions of the insertions and found that rh3-2, emb1138, and rh3-4 mutants have T-DNA insertions in intron 2, exon 3, and intron 9, respectively (Fig. 4A).

The different AtRH3 alleles conditioned distinct phenotypes. Self-pollination of heterozygous rh3-2 and emb1138 plants yielded developing siliques with green and white seeds in an approximately 3:1 ratio; these white seeds shriveled after seed maturation (Fig. 4B). We analyzed over 80 progeny of self-pollinated plants for each allele that were germinated on agar plates with Suc at low light intensity (20 μmol photons m−2 s−1) but recovered no homozygous mutants. These results strongly suggest that null mutants for RH3 are embryo lethal (Fig. 4B), consistent with the identification of emb1138 in a large-scale screen for embryo-lethal mutants (Tzafrir et al., 2004). In contrast, an insertion in intron 9 (rh3-4) resulted in seedlings with small chlorotic cotyledons and pale-green leaves with delayed growth (Fig. 4C). Homozygous rh3-4 mutants were able to grow on soil, set viable seeds, and be maintained as a homozygous population. RH3 mRNA was strongly reduced in rh3-4 homozygotes (Fig. 4D), whereas RH3 protein was reduced to approximately 20% of the wild-type level (Fig. 5A).

The heteroallelic progeny of a complementation cross between rh3-4 and emb1138 germinated as albino seedlings, which was an intermediate phenotype with respect to those of the parental lines (Supplemental Fig. S3A). Moreover, the AtRH3 complementary DNA (cDNA) complemented the rh3-4 phenotype (Fig. 4C). Collectively, these results confirm that a reduction in RH3 function causes a pale-green seedling phenotype, whereas complete loss leads to embryo lethality.

Immunoblot analysis of total leaf extracts was used to quantify effects on the thylakoid-localized photosynthetic machinery in rh3-4 homozygotes (Fig. 5B). Accumulation of protein subunits of PSI (psaD/subunit II and psaF/subunit III), PSII (D1, D2, and OEC23), as well as the cytochrome b6f (cytb6f) complex (PetD/subunit IV and PetA/cytb6f) complex was 2- to 3-fold reduced in rh3-4. In contrast, the α- and β-subunits of the ATP synthase increased severalfold. For comparison, we also included an analysis of clpr2-1 mutant thylakoids. The reduction of PSI, PSII, and cytb6f subunits was 2- to 3-fold stronger than in rh3-4 mutants, consistent with the stronger phenotype of clpr2-1 (Rudella et al., 2006). clpr2-1 accumulated the precursor form of the PSI subunit III (PsaF), as observed previously.

**Figure 4.** AtRH3 T-DNA insertion mutants and a double knockdown mutant with rh3-4/clpr2-1. A, T-DNA insertions in AtRH3. Exons and introns are indicated by black rectangles and lines, respectively. 5′ and 3′ untranslated regions are shown as white rectangles. The positions of the T-DNA insertions in emb1138, rh3-2 (SALK_025572), and rh3-4 (SALK_005920) are indicated by triangles. B,Embryo-lethal phenotypes associated with insertions in AtRH3. The green and mature siliques resulting from RH3-2/rh3-2 and Emb1138/emb1138 plants segregated approximately one-third white seeds and brown shriveled seeds, respectively. The shriveled seeds did not germinate. WT, Wild type. C, A rh3-4 homozygote showed a pale-green seedling phenotype, whereas the double knockdown mutant rh3-4/clpr2-1 showed an albino phenotype. Seedlings were grown for 9 d (top row) or 18 d (bottom row) on one-half-strength Murashige and Skoog medium containing 2% Suc. The rh3-4 has white cotyledons at early stages of development and exhibits pale-green true leaves. The double knockdown mutant rh3-4/clpr2-1 showed additive white seedlings. AtRH3 cDNA complemented the rh3-4 phenotype (rh3-4/RH3). Bars = 5 mm. D, RNA gel-blot analysis of RH3, CLPR2, and OEC23 mRNA levels in rh3-4, clpr2-1, and rh3-4/clpr2-1 mutants. Total RNA (6 μg) from leaf samples at leaf stage 1.07 planted on one-half-strength Murashige and Skoog medium with 2% Suc was analyzed by RNA gel blots using RH3, CLPR2, and OEC23 probes (left panels). The same blot stained with methylene blue is shown (right panels).
(Rudella et al., 2006), but rh3-4 did not. Similar to rh4-1, ATP synthase levels increased in clpr2-1. This differential down-regulation was surprising to us, but verification of studies on other chloroplast biogenesis mutants showed that it is frequently observed that the thylakoid ATP synthase is less affected (or even increased), whereas the thylakoid electron transport chain complexes are down-regulated. For instance, the apo1 mutant involved in intron II splicing also shows unchanged levels of ATPase but very strong decreases of PSI, PSII, and the cyt b6f complex (Watkins et al., 2011). Arabidopsis lpa2, with defects in PSII assembly, shows a decrease of PSI and PSII but not of ATP synthase (Ma et al., 2007). This shows that chloroplast and thylakoid biogenesis is a complex process. Chlo-roplast chaperone CPN60 was slightly (approximately 2-fold) up-regulated in both single mutants (Fig. 5B), similar to what we observed previously for clpr2-1 (Rudella et al., 2006).

**AtRH3 Is Required for the Splicing of Group II Introns That Coinmunoprecipitated with ZmRH3**

The splicing of each intron whose ortholog coimmunoprecipitated with ZmRH3 (Fig. 4B) was analyzed in the Arabidopsis rh3-4 mutant. For comparative purposes, we included Arabidopsis cfm2-2 mutants that have reduced levels of the plastid-localized splicing factor CFM2 (AT3G01370). cfm2-2 mutants are pale green and have chloroplast biogenesis defects resulting from defects in the splicing of several group II introns, but not in any of the RH3 intron ligands detected here (Asakura and Barkan, 2007). rh3-4 mutants accumulated markedly reduced levels of mature trnA and trnl but accumulated increased levels of their unspliced precursors (Fig. 6A; compare exon and intron levels), indicating a reduction in trnl and trnA splicing efficiency. Poisoned-primer extension assays revealed a clear decrease in the ratio of spliced to unspliced RNA for the trans-spliced intron rps12-int1 (Fig. 6B). The ratio of spliced to unspliced rpl2 and rps12-int2 was also reduced in rh3-4 mutants, but more modestly (Fig. 6B). The fact that effects on the splicing of trnl, trnA, rps12-int1, rps12-int2, and rpl2 were detected even in the weak rh3 mutant allele, in conjunction with the RNA co-IP data, strongly suggests that RH3 directly promotes the splicing of these introns. The splicing of three chloroplast introns that are found in Arabidopsis but not in maize (clpP-int1, clpP-int2, and rpoC1) and introns found in minor peaks in the maize RIP-chip experiments in Figure 3 (ycf3-int1 and ycf3-int2) was unaffected in rh3-4 (Fig. 7). These results suggest that RH3 does not contribute to the splicing of these introns, but analysis of stronger mutant alleles would be required to firmly address this point.

RH3 coimmunoprecipitates immature and mature 23S rRNA (Fig. 3C). This could indicate that RH3 also promotes the assembly of the 50S ribosomal subunit, a defect that is expected to be reflected by a decrease in...
23S rRNA processing (for references and discussion, see Nishimura et al., 2010; Chi et al., 2012). To further address the role of RH3 in rRNA metabolism, we analyzed the four rRNAs by RNA gel-blot hybridization (Fig. 8A). We observed that the accumulation of each rRNA in the 50S ribosomal subunit (23S, 5S, and 4.5S) was slightly reduced in *rh3-4* mutants, whereas 16S rRNA of the 30S particle was not affected.

RH3 and ClpR2 Interact Genetically, But RH3 Is Unlikely To Be a Substrate for the Clp Protease

RH3 was 5- to 10-fold up-regulated in the two ClpPR protease mutants *clpr2-1* and *clpr4-1* (Kim et al., 2009; Zybailov et al., 2009). This could suggest that RH3 is a substrate of the Clp protease, with the increased accumulation of RH3 due to reduced proteolytic capacity in the chloroplast. Alternatively, the increase in RH3 might be an indirect compensatory or pleiotropic response to
the loss of Clp protease capacity. To explore the relationship between RH3 and ClpPR protease, a double homozygous mutant was generated between rh3-4 and clpr2-1 (Fig. 4C; Supplemental Fig. S4). The double-homozygous seedlings were albino and very slow growing; this phenotype was much stronger than the phenotype of either of its parents (Fig. 4C). However, viable seed could eventually be obtained from the homozygous rh3-4 clpr2-1 double mutants (see “Materials and Methods”).

Immunoblot and RNA gel-blot analyses were carried out to compare levels of RH3 and ClpR2 mRNA (Fig. 4D) and protein (Fig. 5A) in rh3-4 and clpr2-1 single and double mutants. RH3 mRNA was reduced to near undetectable levels in both lines that were homozygous for rh3-4 but was increased severalfold in homozygous clpr2-1 mutants. CLPR2 mRNA levels were unaffected in rh3-4 mutants but strongly reduced in the clpr2-1 background (Fig. 4D). Consistent with our previous mass spectrometry-based analysis (Zybailov et al., 2009), RH3 protein abundance strongly increased (more than 10-fold) in clpr2-1 seedlings (Fig. 5A). Whereas RH3 protein was strongly reduced in the rh3-4 mutant, RH3 protein levels were approximately 2-fold increased in the double mutant as compared with the wild type, indicating that the reduced ClpR2 levels up-regulated RH3 even in the rh3-4 background. The ClpR2 protein level in clpr2-1 was about 20% of the wild type, as reported previously (Rudella et al., 2006), and remained at that level in the double mutant (Fig. 5A).

Consistently, ClpR2 protein levels were not significantly affected in the rh3-4 seedlings. Chloroplast cpHSP70 levels were used as an internal control and were slightly increased in the single and double mutants (Fig. 5A).

The expression of RH3 is under strong developmental control, with highest expression levels in young, developing leaf tissue (Fig. 2, B–D). To better resolve the developmental kinetics and gain insight into why RH3 protein overaccumulated in the clpr2-1 mutant, we determined RH3 mRNA and protein levels in different leaf rosette developmental stages (stages 1.04–1.15). RH3 mRNA levels decreased during the development of the leaf rosette in both the wild type and clpr2-1 (Fig. 9A); in the oldest rosette, transcript levels were identical in the wild type and clpr2-1. However, RH3 mRNA levels were severalfold higher in the clpr2-1 mutant during early leaf development (stages 1.04 and 1.08), indicating delayed transcriptional down-regulation. A similar but more dramatic pattern was observed at the RH3 protein level (Fig. 9B): RH3 levels clearly decreased with progressive leaf rosette development, but RH3 levels were consistently higher in the clpr2-1 mutant (Fig. 9B).

To determine whether RH3 up-regulation is specific for ClpPR core protease mutants or is a general result of defects in chloroplast biogenesis, we determined RH3 protein levels for two other pale-green chloroplast mutants, ffc1-2 and tic40 a1, at leaf stages 1.07 and 1.14 (Fig. 9C). ffc1-2 is deficient for chloroplast signal recognition particle subunit SRP54, involved in the...
targeting of various proteins to the thylakoid membrane (Rutschow et al., 2008). The mutant tic40 is deficient for the inner chloroplast envelope protein TIC40, involved in the import of nucleus-encoded proteins (Kovacheva et al., 2005). At leaf stage 1.07, RH3 levels increased about 2-fold in the ffc1-2 and tic40 mutants, compared with approximately 5-fold in clpr2-1. At least stage 1.14, RH3 levels in ffc1-2 and tic40 mutants increased to nearly the level as in the clpr2-1 mutant. Together with the increased RH3 mRNA levels, this indicates that the increase in RH3 levels is not specific to the ClpPR core mutants, even if up-regulation in the early stages of leaf development is stronger in clpr2-1.

If RH3 were a substrate for degradation by the ClpPR core complex, RH3 or ClpR3 fragments might be found in association with the ClpPR complex. Therefore, we carried out co-IP experiments against stroma from isolated chloroplasts of wild-type and clpr2-1 plants using our anti-ClpR2 and anti-RH3 antisera. Whereas co-IP using anti-ClpR2 and anti-RH3 successfully precipitated ClpR2 and RH3, respectively, anti-ClpR2 did not coimmunoprecipitate with RH3, nor did anti-RH3 coimmunoprecipitate with ClpR2 (Fig. 10). Furthermore, fractionation of stromal extracts from Arabidopsis wild-type and clpr2-1 mutant plants on Suc gradients identified RH3 in high-molecular-mass fractions over 550 kD but never in 350-kD ClpPR complexes (Supplemental Fig. S5). Similarly, tandem mass spectrometry analysis of tagged ClpPR core complexes, such as His6-tagged ClpPR cores (Rudella et al., 2006) or StrepII-tagged ClpPR cores (Olinares et al., 2011), did not find RH3 protein, unlike other putative substrates (data not shown).

Comparison of RNA Metabolism in rh3-4 and clpr2-1 Mutants

So far, we determined that RH3 associates with and promotes the splicing of several group II introns (rps12-int1, rpl2-int, trnA, and trnI) and also coimmunoprecipitates with rrm23 (Fig. 3B). Moreover, we showed that mature forms of rsm4.5 and rsm5 as well as the 0.5-kb fragment of 23S rRNA accumulated at lower levels in rh3-4 (Fig. 8), suggesting a reduced accumulation of the assembled 50S ribosome particles. In contrast to rh3-4, the ratio of spliced to unspliced RNAs from the RH3-dependent introns is nearly normal in clpr2-1 mutants (Figs. 6 and 7). The metabolism of rRNAs also differed between the two mutants: various precursors overaccumulated in the clpr2-1 background, but the fully processed rRNAs accumulated to wild-type levels. Thus, the abundance of assembled 50S ribosomes was unaffected in the clpr2-1 mutant but was reduced in the rh3-4 mutant.

DISCUSSION

RH3 Is a Group II Intron-Splicing Factor in Plastids and May Also Contribute to 50S Ribosomal Subunit Assembly

In this study, we have shown that RH3 is a plastid-localized DEAD box RNA helicase in both maize and
Arabidopsis that functions in the splicing of specific group II introns; RH3 is thus part of the network of plastid splice factors (Supplemental Fig. S6). This was based on a number of complementary measurements, namely (1) the interaction of RH3 with a subset of chloroplast type II introns (Fig. 3), and (2) specific splice defects in the rh3-1 mutant (Figs. 6 and 7). Furthermore, sequence analysis of RH3 clearly shows that RH3 contains the conserved domains for ATP-dependent RNA helicase activity and also has the conservative DEAd domain. Thus, our analysis indicates that RH3 belongs to the class of DEAd box RNA helicase proteins. In contrast, some of the other chloroplast helicases do not have a strictly conserved DEAd domain (but have a DEXD domain), whereas others lack some of the key residues involved in ATP binding and hydrolysis. Strikingly, the role of RH3 in intron slicing is also consistent with the phylogenetic analysis of the RH3 family, in that RH3 is part of a clade with two closely related mitochondrial RH proteins (RH9 and RH39; Matthes et al., 2007), one of which (RH39) has been demonstrated to function in intron II splicing in mitochondria (Köhler et al., 2010).

Other results suggest that RH3 may have an additional function in the assembly of 50S ribosomal subunits. First, the abundance of mature 50S ribosomal subunits is reduced in rh3 mutants, as determined by the significant decrease in the accumulation of 23S, 4S, and 5S rRNA but not the 16S RNA of the 30S particle (Fig. 8). In addition, RH3 is found in particles of a suitable size (approximately 45S; Fig. 3A), and it coimmunoprecipitates with pre-23S rRNA (Fig. 3C). Such a role would be in keeping with the fact that other DEAd box proteins promote ribosome assembly in plastids and in other systems (Nishimura et al., 2010; Linder and Jankowsky, 2011; Chi et al., 2012). However, we cannot rule out the possibility that the decrease in ribosome abundance in rh3 mutants is a secondary effect of the splicing defects that impacts several components of the translation machinery (trnI, trnA, and rps12). The only viable rh3 allele was quite weak, and it is possible that a ribosome assembly defect would be revealed more clearly with a stronger (but not total) loss of function. The fact that RH3 is essential for embryogenesis and viability, based on the phenotypes of the two null RH3 alleles, is consistent with the need for functional plastid ribosomes and translation during embryogenesis in Arabidopsis (Stern et al., 2010).

**RH3 Increase in Clp Protease Mutants Constitutes a Compensatory Effect and Must Involve Retrograde Signaling from the Plastid to the Nucleus**

Both RH3 mRNA transcripts and protein levels of RH3 were severalfold increased in the clpr2-1 background. The strong increase in RH3 is not specific for clpr2-1 but applies also to pale-green mutants in plastid ClpPR genes, such as clpr4-1 (Kim et al., 2009) and clpp3 (J. Kim, P.D. Olinares, and K.J. van Wijk, unpublished data). Because these ClpPRs are subunits of the same Clp core protease complex, this shows that RH3 levels increase in response to the loss of ClpPR protease capacity. However, we did not find evidence that RH3 increases are directly due to the loss of degradation capacity; rather, our data suggest that the simultaneous increase in RH3 mRNA and protein constitutes a compensatory effect to the Clp mutant phenotype. The increase in RH3 mRNA level suggests a transcriptional response due to the loss of Clp core activity and must involve retrograde signaling from plastid to nucleus; the nature of this signal is still unclear (Sun et al., 2011), but many studies have indicated that inhibition of chloroplast homeostasis does contribute to the signal (Chan et al., 2010; Šimková et al., 2012). Indeed, multiple proteins involved in regulating chloroplast biogenesis and homeostasis, such as nucleoid-associated proteins (e.g. pTAC1/Whyl), translation factors (e.g. EF-Tu-1, Ef-Tu-G/Sco1, and Ef-Tu-TypeA/BipA), as well as the major chaperones (CPN60, HSP70, HSP90, and ClpB3), were up-regulated in the Clp mutants (Kim et al., 2009; Zybaliov et al., 2009), indicative of significant protein stress.

**The Difference between RH3 and Other Plastid-Localized DEAd Box Helicases**

Out of the seven identified plastid-localized DEAd box RNA helicases (RH3, -22, -26, -39, -47, -50, and -58), only the functions of RH39 and RH22 were previously studied in detail in Arabidopsis (Nishimura et al., 2010; Chi et al., 2012). Our phylogenetic analysis showed that five of these plastid helicases (RH22, -39, -47, -50, and -58) form a separate clade, distinct from the clade with RH3 and its mitochondrial and nuclear homologs, and that RH26 is in a clade with extraplastidic proteins with unknown functions. The five plastid RH proteins in clade 8 lack several conserved motifs likely reducing RNA helicase activity, whereas plastid RH3 and RH26 show full conservation of the
DEAD box RNA helicase motifs. Recently, we provided indirect evidence that RNA metabolism and ribosome assembly occur in association with nucleoids (Majeran et al., 2012). The identified plastid RH helicases were found in high-mass complexes (greater than 1 MD) in Arabidopsis stroma (Oliñares et al., 2010) or present in maize nucleoids (Majeran et al., 2012), consistent with a role in RNA metabolism and ribosome assembly. Future experiments should focus on the identification of functions of plastid-localized RH26, -47, -50, and -58.

MATERIALS AND METHODS

Phylogenetic and Functional Domain Analysis

Selected proteins were aligned using MUSCLE (http://toolkit.tuebingen.mpg.de/muscle) using 15 to 25 iterations, and the aligned selected sequences in Clustal format were viewed and edited (removal of gaps and variable extensions) in Jalview. Sequences were then converted into PHYLF format using http://searchlauncher.bcm.tmc.edu/seq-util/readseq.html. Phylogenetic trees were generated (1,000 iterations) using the CLIPRES Web-portal (http://www.phylo.org/sub_sections/portal/) using the RAAML HPC Blackbox, with the general time reversal model selected as the protein substitution matrix. The resulting phylogenetic trees were annotated in FigTree (http://tree.bio.ed.ac.uk/software/figtree/).

Plant Materials and Growth Conditions

Maize (Zea mays inbred line B73) seedlings for stroma and cDNA preparations were grown in a growth chamber under a 12-h-light (31°C, 400 μmol photons m⁻² s⁻¹) and 14-h-dark (22°C) cycle unless otherwise indicated. Arabidopsis seedlings for Rip-chip and Suc gradient were isolated as described previously (Kovacheva et al., 2000; Tzafrir et al., 2004; Asakura and Barkan, 2007; Rutschow et al., 2006) with LR Clonase II enzyme mix (Invitrogen) according to the manufacturer.

RNA Extraction and Determination of RNA Concentration

Maize stroma for RIP-chip and Suc gradient were isolated as described previously (Majeran et al., 2012). The isolated stroma and cDNA preparations were grown in a growth chamber under a 12-h-light (31°C, 400 μmol photons m⁻² s⁻¹) and 12-h-dark (22°C) cycle. They were harvested 7.5 d after planting for the nucleoid isolation or 8.5 d for chloroplast isolation.

Arabidopsis (Arabidopsis thaliana) T-DNA insertion lines SALK_025572 (rh3-2), SALK_005920 (rh3-4), and CS16011 (emb1138) were obtained from the Salk Institute Genomic Analysis Laboratory. emb1138, SALK_046378 (clpr2-1), SAIL_6E07_05 (Atcfm2-2), and clpr2-1 mutants were described previously (Amin et al., 1999; Tzafrir et al., 2004; Asakura and Barkan, 2007; Rutschow et al., 2008). The tci40 mutant (SAIL_92_C10) was described previously (Kovacheva et al., 2005). Arabidopsis plants used for protein, DNA, and RNA extraction were grown on one-half-strength Murashige and Skoog plates supplemented with 2% Suc under short-day and low-light (60 μmol m⁻² s⁻¹) conditions followed by transfer to soil and growth for 6 months under long days at low light intensity (20 μmol photons m⁻² s⁻¹). Tagged fusion proteins from pET28a(+) or pET21a(+) vectors (Novagen), respectively. A ZmRH3A recombinant protein was expressed in BL21(DE3) Star (Novagen) and induced by the addition of 1 μg isopropyl-1-thio-β-D-galactopyranoside for 2 h. AtClpR2 recombinant protein was expressed in Rosetta (DE3) cells (Novagen) and induced by the addition of 0.5 μg isopropyl-1-thio-β-D-galactopyranoside for 4 h. Both proteins were recovered in a soluble fraction, purified with nickel-nitrilotriacetic acid agarose beads (Qia- gen) according to the manufacturer’s protocol, and equilibrated with phosphate-buffered saline. Polyclonal antisera from 10 mg of antigens were generated in rabbits at the University of Oregon antibody facility for anti-ZmRH3A or at Alfa Diagnostic International for anti-AtClpR2. Antisera were affinity purified against the same antigen that had been used for the immunizations coupled to a HiTrap N-hydroxysuccinimide (NHS) ester-activated column (GE Healthcare Life Science) or cyagen bromide-activated Sepharose (Sigma).

Protein Extraction and Immunoblot Analysis

Arabidopsis seedlings from six to 10 plants at leaf stage 1.07, unless otherwise mentioned, were ground with a mortar and pestle in liquid nitrogen by adding an extraction buffer (2% SDS, 50 mM Tris-HCl, pH 8.8, and 5 mM EDTA) with protease inhibitors as described above and vortexed for 30 s to solubilize proteins. Insoluble materials were removed by centrifugation (0.8-Ml column, 30 μm; Pierce) for 1 min at 10,000 rpm. Protein concentrations were determined by bicinchoninic acid assay (Smith et al., 1985). Samples were denatured in 3% or 2% Laemmli buffer (Laemmli, 1970) and heated at 75°C for 10 min. Proteins were resolved by SDS-PAGE followed by electroblotting to nitrocellulose membranes and then the membranes were stained with Ponceau S (0.3% Ponceau S, 3% TCA). Immunoblotting and enhanced chemiluminescence were performed as described previously (Barkan, 1998). The antisera against PsaA (Pisum sativum) CPN60 and spinach (Spinacia oleracea) cpHS70 were kindly provided by Masato Nakai (Osaka University). The antisera against barley (Hordeum vulgare) PsaA was provided by Hendrik Scheller (Royal Veterinary and Agricultural University, Denmark). The antisera against sorghum (Sorghum bicolor) OEC23 and OEC16, barley PsaD, and maize PetD and WHY1 were described previously (Voeleker and Barkan, 1995; Priksyl et al., 2008).

RIP-chip and rrn23 Poisoned-Primer Extension Assays

RIP-chip assay of RNAs that communoprecipitate with RH3 and OEC16 was performed as described previously (Schmitz-Linneweber et al., 2005). Each ccr-IP used affinity-purified antibodies and 100 μL of stromal extract (approx. 1 μg of stromal protein). A poisoned-primer assay to distinguish mature and precursor forms of 23S rRNA was performed as described previously (Asakura and Barkan, 2006).

Arabidopsis DNA Extraction and PCR Amplification

Arabidopsis DNA isolation was as described previously (Asakura and Barkan, 2006). T-DNA insertions were confirmed by PCR amplification. Primers are provided in Supplemental Table S2.

Arabidopsis RH3 Complementation

A cDNA clone of AtRH3 (stock no. U2189) was obtained from the Arabidopsis Biological Resource Center. A cDNA fragment of AtRH3 was amplified using platinum Pfx DNA polymerase (Invitrogen) with primers YAAAHR3_3'-F (start) (5'-CACCATCGCGCGGACAGTT-3') and YAAAHR3_244-R (stop) (5'-CTAAAATCCTTCACTTACAGGAC-3') and subcloned into a pENTR/D-TOPO vector. The resulting pENTR-RH3 was cut with Mfil gel purified, introduced into a binary pEAREGATE100 vector (Earley et al., 2006) with LR Clonase II enzyme mix (Invitrogen) according to the manufacturer’s instruction, and electrotransformed into E. coli strain Top10. The resulting pEAREGATE-AtRH3 clone was electrotransformed into Agrobacterium tumefaciens (strain GV3101). Transformation of the homologous rh3-4 was carried out using the floral dip method (Clough and Bent, 1998). Transformants were selected on soil with spraying three times with BASTA herbicide with a 3-d interval. T1 and T2 generations of transformants were assayed by PCR genotyping.
Arabidopsis RNA Analysis

RNA was extracted from the Arabidopsis leaf tissues by using Trizol reagent (Invitrogen) according to the manufacturer’s instruction. RNA gel-blot hybridizations and probes for chloroplast ribosomes were as described previously (Zybaliov et al., 2009). Other probes were amplified by PCR and reverse transcription-P-CR from Arabidopsis genomic DNA or RNA with the primers listed in Supplemental Table S2. Poisoned-primer extension assays were performed using 10 μg of leaf RNA as described previously (Asakura and Barkan, 2006).

Maize Stroma Suc Gradient to Dissociate 50S and 30S Ribosomes

A maize stroma Suc gradient was carried out as described with minor modifications (Barkan et al., 2007). A total of 100 μL of maize stroma (approximately 1 mg of protein) was diluted with 400 μL of dilution buffer (20 mM Tris-4CL, pH 7.8, 100 mM ammonium chloride, and 5 mM 2-mercaptoethanol) to reduce the magnesium concentration to 2 mM in maize stroma. A total of 500 μL of diluted stroma was layered onto a 10% to 40% Suc gradient containing KEX buffer (20 mM Tris-HCl, pH 7.8, 1 mM MgCl2, and 100 mM ammonium chloride) and was ultracentrifuged in a Beckman SW41 rotor at 35,000 rpm for 7 h at 4°C. RNA was extracted from 300 μL of each fraction by adding 50 μL of 5% SDS and 0.2 mM EDTA, followed by phenol/chloroform extraction and ethanol precipitation. An equal proportion of each fraction was used for immobil blot and RNA gel-blot analyses.

Arabidopsis Stroma Suc Gradient

A total of 500 μL of Arabidopsis stroma (1 mg of protein from the wild type and clp2r-1) with 120 units of RNAsin (Promega) was fractionated by a 10% to 40% Suc gradient containing KEX buffer [30 mM HEPES-KOH, pH 8, 100 mM KOAc, 10 mM Mg(OAc)2, and 5 mM dithiothreitol] in a Beckman SW41 Ti rotor at 35,000 rpm for 3 h at 4°C. An equal proportion of each fraction was used for immunoblot analysis.

Co-IP

For Co-IP for Clp2r and RH3, 240 μL of Dynabeads protein A (Invitrogen) was washed three times with phosphate-buffered saline, pH 8.0, with 0.1% (v/v) Igepal CA-630. About 400 μL of stroma (390 μg mL−1) for one reaction from wild-type and clp2r-1 Arabidopsis was clarified by centrifugation at top spin for 10 min at 4°C. Twenty microliters of washed Dynabeads was added in the stroma and incubated for 30 min at 4°C, and beads were removed by a Dyna Magnet (Invitrogen). Preclarred stroma were split into three tubes to 400 μL and 40 μL was saved as a total sample. Beads were divided into two 60-μL aliquots for wild-type and clp2r-1 stroma and washed once with co-IP buffer (20 mM Tris-HCl, pH 7.0, 150 mM NaCl, 1 mM EDTA, 0.2% Igepal CA-630, and 5 μg mL−1 aprotinin). Preclarred stroma aliquots (approximately 400 μL) were incubated with anti-ZnRH3A-Dynabeads or anti-AKL22-Dynabeads in 0.4 mL of co-IP buffer. Samples were incubated for more than 90 min at 4°C and then washed six times with co-IP buffer. Bound proteins were eluted with 40 μL of 1.5× Laemml buffer without 2-mercaptoethanol at 5 min at 70°C or 95°C, and beads were removed by Dyna Magnet. Before running SDS-PAGE gels, 4 μL of 2-mercaptoethanol was added to the samples, heated for 10 min at 75°C, and microdiluted for 5 min to remove insoluble materials.

Supplemental Data

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

Supplemental Figure S1. Phylogenetic tree of the RH3 DEAD box helicase family in Arabidopsis, maize, rice, and E. coli based on the core domains with major gaps removed.

Supplemental Figure S2. Multiple sequence alignment of ZmRH3A, ZmRH3B, and AtRH3.

Supplemental Figure S3. An albino seedling phenotype of heteroallelic rh3-4/clp2r-1.

Supplemental Figure S4. PCR genotype of the double knockdown mutant rh3-4/clp2r-1.

Supplemental Figure S5. RH3 in the wild type and the clp2r-1 mutant is in the same fraction in Suc gradients.

Supplemental Figure S6. RH3 functions and chloroplast splicing factors and their intron target in land plants (updated from Barkan, 2011).

Supplemental Table S1. Protein accession numbers, nomenclature, annotation, and clade assignments for the RH3 DEAD box helicase family in Arabidopsis, maize, and rice used for phylogenetic analysis.

Supplemental Table S2. Primer sequences used for the verification of Arabidopsis T-DNA insertions and RNA analysis.

Supplemental Table S3. All 154 DEAD box RNA helicase sequences (all with DEAD or DExD motifs) used for the phylogenetic analysis.

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