RHON1 Mediates a Rho-Like Activity for Transcription Termination in Plastids of Arabidopsis thaliana

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Although transcription termination is essential to generate functional RNAs, its underlying molecular mechanisms are still poorly understood in plastids of vascular plants. Here, we show that the RNA binding protein RHON1 participates in transcriptional termination of rbcL (encoding large subunit of ribulose-1,5-bisphosphate carboxylase/oxygenase) in Arabidopsis thaliana. Inactivation of RHON1 leads to enhanced rbcL read-through transcription and to aberrant accD (encoding β-subunit of the acetyl-CoA carboxylase) transcriptional initiation, which may result from inefficient transcription termination of rbcL. RHON1 can bind to the mRNA as well as to single-stranded DNA of rbcL, displays an RNA-dependent ATPase activity, and terminates transcription of rbcL in vitro. These results suggest that RHON1 terminates rbcL transcription using an ATP-driven mechanism similar to that of Rho of Escherichia coli. This RHON1-dependent transcription termination occurs in Arabidopsis but not in rice (Oryza sativa) and appears to reflect a fundamental difference between plastomes of dicotyledonous and monocotyledonous plants. Our results point to the importance and significance of plastid transcription termination and provide insights into its machinery in an evolutionary context.

INTRODUCTION

Transcription termination, which involves arrest of RNA biosynthesis, release of the transcript, and dissociation of the RNA polymerase from the DNA template, is essential for proper gene expression. It helps avoid interference with transcription of downstream genes. Moreover, it maintains a pool of RNA polymerases available for reinitiation or transcription, prevents formation of antisense RNAs that can interfere with normal pre-RNA production, and thereby avoids aberrant gene expression (Greger and Proudfoot, 1998; Peters et al., 2012). Termination mechanisms vary considerably between different organisms, ranging from relatively simple to exceptionally complex processes. There are at least two different ways to terminate transcription in Escherichia coli and other prokaryotes (Richardson, 2003; Peters et al., 2011). Intrinsic termination requires mainly elements located on the mRNA, whereas Rho-dependent termination relies on both mRNA elements and trans-acting factors (Richardson, 2003; Peters et al., 2011). In eukaryotes, two different pathways for transcriptional termination by RNA polymerase II have been proposed: One uses the Nrd1 complex, whereas the other uses 3’ cleavage and polyadenylation factors together with Rat1 exonuclease (Rondon et al., 2008; Richard and Manley, 2009). By contrast, less is known about the molecular mechanisms that mediate transcription termination in organelles.

Plastids of vascular plants originated from an ancestral cyanobacterial endosymbiont (Timmis et al., 2004). Regulation of plastid gene expression is distinct from that of eubacteria as well as from that in the eukaryotic cytosol and reflects a unique chimeric system assembled from multiple components whose origins can be traced to endocytobiosis (Stern et al., 2010; Barkan, 2011). Plastid transcription is performed by a nucleus-encoded phage-type RNA polymerase (NEP) in addition to the cyanobacterium-derived, plastid-encoded RNA polymerase (PEP) (Barkan, 2011; Liere et al., 2011). However, how transcription termination occurs in plastids remains obscure. Most plastid genes contain inverted repeat sequences at their 3’ ends that are able to form stem-loop structures (Sugita and Sugiura, 1984; Kirsch et al., 1986). Such structures were initially assumed to act as transcription terminators, in a similar way as Rho-independent terminators (Sugita and Sugiura, 1984; Kirsch et al., 1986). However, both in vitro and in vivo assays have shown that in plastids, 3’ inverted repeat sequences function as mRNA processing and stabilizing elements rather than as efficient terminators (Stern and Gruissem, 1987; Monde et al., 2000). Nevertheless, both PEP and NEP polymerases terminate in vitro at strong intrinsic bacterial terminators containing GC-rich RNA hairpins followed by several contiguous uridines (Chen and...
Orozco, 1988; Jeng et al., 1990; Lerbs-Mache, 1993; Kühn et al., 2007). Therefore, it seems likely that RNA polymerases of plastids recognize bacterial terminator sequences with specific features under in vivo conditions. However, no factors involved in plastid transcription termination have been identified and characterized in vascular plants.

Proper expression of plastid genes relies on coordinated interplay between cis-acting elements at the plastid DNA/RNA level and trans-acting factors, most of which are nuclear encoded and specifically bind to their cis-acting targets. Comparative analysis of the plastid genomic evolution of vascular plants has indicated that vascular plants underwent frequent plastome rearrangements, such as large inversions, gene and intron gains and losses, expansion, contraction, and loss of the inverted repeats that vary between plant species (Wicke et al., 2011). The intergenic expansion, contraction, and loss of the inverted repeats that define plastid gene expression. However, how these processes were established during evolution still remains elusive.

Here, we studied the mechanism of transcription termination of plastid genes mediated by the plastid RHON1 protein. RHON1 was originally identified in a screen for interaction partners of the endonuclease RNase E (RNE) and was proposed to be involved in RNE-mediated plastid RNA processing (Stoppel et al., 2012). We show that RHON1 is also involved in the transcription termination of rbcL, encoding the large subunit of ribulose bisphosphate carboxylase oxygenase. Like Rho factors in E. coli, the catalytic activity of RHON1s is based on the use of an ATP-driven mechanism. In addition, we found that RHON1-dependent transcription termination occurs in Arabidopsis thaliana but not in rice (Oryza sativa), which appears to be related to a fundamental difference in the rbcL-accD intergenic region of eudicotyledonous and monocotyledonous plants.

RESULTS

Mutation of RHON1 Affects Transcriptional Termination of rbcL

A mutant with a high chlorophyll fluorescence phenotype was isolated from the Scheible and Somerville T-DNA Arabidopsis lines (Figure 1A). The mutation was mapped to a 100-kb region between markers CER474 and CER448 on BAC F9P14 of chromosome 1 (Figure 1B). Sequencing of the 33 genes in the interval revealed a T-DNA insertion within the second exon of At1g06190 (Figure 1C). In a previous study, the At1g06190 protein was given the name RHON1 because its C-terminal domain is similar to the N-terminal part of the bacterial transcription terminator Rho (Stoppel et al., 2012). Accordingly, we designated our high chlorophyll fluorescence mutant as rhon1-2. RT-PCR and immunoblot analysis showed expression of RHON1 in wild-type plants, but not in rhon1-2 (Figure 1D). Expression of full-length RHON1 cDNA under the control of the constitutive 3SS promoter in rhon1-2 mutants fully restored the wild-type phenotype (Figure 1A). This functional complementation indicates that the RHON1 (At1g06190) gene is indeed responsible for the rhon1-2 phenotype. To explore the possible function of RHON1 in chloroplast gene expression, we examined the plastid gene transcription profile in the rhon1-2 mutant by microarray hybridization. This analysis revealed that the abundance of five plastid transcripts (accD, psal, ycf4, cema, and petA) was increased 20- to 40-fold in rhon1-2 (Supplemental Figure 1). Two NEP promoters (PaccD-172, 252) have been found to drive transcription of the Arabidopsis accD gene (promoters are specified by the gene name and the position of the initiation nucleotide with respect to the start of the coding sequence) (Swiatecka-Hagenbruch et al., 2007). However, a large polycistronic precursor transcript spanning the rbcL, accD, psal, ycf4, cema, and petA genes was also reported (Walter et al., 2010; Stoppel et al., 2012). This large polycistronic precursor is barely detectable in wild-type plants. This raised the question whether the dramatically increased levels of these five transcripts in rhon1-2 were associated with transcriptional and/or post-transcriptional processes related to the rbcL-accD-psal-ycf4-cema-petA operon. To analyze this operon in detail, RNA gel blot analysis using probes specific for the individual mature RNAs was performed. As shown in Figure 2A, the levels of unprocessed precursors (8.5-, 6.7-, 4.3-, and 1.45-kb fragments) were increased, as were those of the mature RNAs except for mature rbcL, which was not significantly changed in rhon1-2.

Levels of both the precursor and mature forms of accD, psal, ycf4, cema, and petA mRNAs were enhanced, suggesting increased transcription of these genes and/or increased stability of the mRNAs in rhon1-2. A run-on assay showed that the transcription rates of accD, psal, ycf4, cema, and petA were increased in rhon1-2 plants, whereas the control (psbA) did not show obvious changes (Figure 2B; Supplemental Figure 2). This increase observed for these five gene transcripts could theoretically be due to augmented transcription of the large polycistronic rbcL transcript in rhon1-2. However, our results showed that the transcription rate of rbcL itself was not increased in rhon1-2, indicating that the transcription of the rbcL-accD-psal-ycf4-cema-petA operon as a whole is not enhanced. We therefore conclude that transcription terminates efficiently after rbcL in wild-type plants but not in rhon1-2, which leads to the increased transcription of downstream genes of rbcL. The unchanged abundance of monocistronic rbcL in rhon1 might be due to endonucleolytic cleavage of the extended polycistronic RNAs downstream of rbcL, followed by 3’ exonuclease trimming to the stabilizing 3’ hairpins. Nevertheless, the possibility of compensation by increased mRNA stability as suggested by Shiina et al. (1998) for the unaltered level of monocistronic rbcL cannot be completely excluded.

Since accD is transcribed by NEP, we reasoned that its increased levels in rhon1-2 might be a response to defects in PEP activity, as plastid transcripts generated by NEP are known to be upregulated in response to impaired PEP activity (Pfalz et al., 2006). However, RNA gel blot analysis showed that the levels of mRNAs transcribed by PEP in rhon1-2 were comparable to those in wild-type plants (Supplemental Figure 3A). In addition, PEP abundance of rhon1-2 was not changed as demonstrated by an immunoblotting assay with RpoB antiserum (Supplemental Figure 3B). This result indicates that PEP activity is not affected in rhon1-2.
Aberrant Transcription Initiation within the rbcL-accD Intergenic Region in rhon1-2

It was reported that the overaccumulation of some plastid polycistronic transcripts is associated with a change of promoter usage (Haley and Bogorad, 1990; Sexton et al., 1990; Schweer et al., 2006). Overaccumulation of polycistronic rbcL transcripts could be due to the activation of an unknown promoter(s) in rhon1-2. To address this possibility, we compared promoter usage in wild-type and rhon1-2 plants by mapping the transcription initiation site(s) using 5′-rapid amplification of cDNA ends (RACE) analysis combined with enzymatic treatment of RNA with tobacco acid pyrophosphatase (TAP). This approach not only can identify primary and secondary 5′-ends of plastid transcripts but also allows discrimination between NEP and PEP promoters by comparing the 5′-ends of transcripts from spectinomycin-treated seedlings with those of untreated seedlings (Swiatecka-Hagenbruch et al., 2007).

Previous work has identified a PEP promoter for rbcL in Arabidopsis, PrbcL-179, which is responsible for the generation of monocistronic rbcL (Hanaoka et al., 2003). Our results revealed two transcription initiation sites for the rbcL gene. Initiation at PrbcL-179 was detected at high levels in green seedlings but at low levels in seedlings treated with spectinomycin (Figure 3A). This suggests that PrbcL-179 is PEP dependent, consistent with the earlier report (Hanaoka et al., 2003). Meanwhile, another transcript initiation site, PrbcL-319, was detected only in spectinomycin-treated seedlings and not in green seedlings (Figure 3A), suggesting that this transcription initiation site is NEP dependent. Recently, a rbcL NEP promoter was identified in barley (Hordeum vulgare) as well (Zhelyazkova et al., 2012). The usage of the PrbcL-179 and PrbcL-319 promoters did not differ between the wild type and rhon1-2, suggesting that the accumulation of rbcL polycistronic transcript in rhon1-2 is not related to rbcL promoter usage. Sequencing of 5′-3′-end ligation products of the 8.5-kb rbcL polycistronic transcript obtained by circularized RNA (CR) RT-PCR indicated that its 5′ end corresponds to the rbcL PEP transcription initiation site (Supplemental Figure 4), which is compatible with the assumption that the rbcL polycistronic mRNA could be the byproduct of inefficient transcriptional termination of rbcL. In addition, no difference in 3′- or 5′-ends of rbcL monocistronic transcript between wild-type and mutant plants was observed (Supplemental Figure 4).

Analysis of accD transcripts in wild-type plants revealed one PCR fragment that was amplified only from TAP-treated RNA isolated from spectinomycin-treated seedlings (Figure 3B). Sequence analysis showed that it corresponds to one of the two reported NEP-dependent transcript initiation sites, PaccD-252 (Swiatecka-Hagenbruch et al., 2007). The other transcript initiation site, PaccD-172, was not detected in wild-type plants under our conditions and was probably not active at this developmental stage, as described by Courtois et al. (2007). However, the amplified products from TAP-treated RNA isolated from rhon1-2 green seedlings displayed a ladder-like pattern, which was markedly different from that of wild-type plants (Figure 3B). This ladder-like pattern of accD transcript initiation sites in rhon1-2 was further confirmed by primer extension assays (Figure 3C). Extensive sequencing of those PCR products indicated that there were multiple transcript initiation sites of accD scattered all over the rbcL-accD intergenic region. In addition, some transcript initiation sites were even mapped within the 5′-end of the accD coding region (Figure 3B). To our...
surprise, these transcript initiation sites disappeared when rhon1-2 seedlings were treated with spectinomycin: As shown in Figure 3B, the rhon1-2 and wild-type plant RNAs showed similar banding patterns in spectinomycin-treated plants. Our results clearly show that absence of RHON1 not only increases transcriptional read-through of the rbcL terminator but also leads to aberrant transcription initiation in the rbcL-accD intergenic region.

In agreement with aberrant transcription initiation, the mature accD transcripts of rhon1-2 plants showed differences in their 5'-ends but had similar 3'-ends when detailed end-mapping of accD transcripts by (CR)RT-PCR was performed (Supplemental Figure 4).

RHON1 Binds to the 3'-Untranslated Region of rbcL

The overaccumulation of rbcL-accD polycistrionic transcript in rhon1-2 suggested that RHON1 could be involved in the regulation of rbcL transcription termination, acting as a trans-factor. If that were the case, it seemed likely that RHON1 would interact with the mRNA of rbcL. The binding activity of the RHON1 protein to rbcL mRNA was investigated by electrophoretic mobility shift assays (EMSA). The C-terminal fragment of 100 amino acids (301 to 401 amino acids) comprising the potential Rho-N domain was expressed for EMSA analysis. Indeed, this fragment was able to bind to the 3'-untranslated region (UTR) of rbcL (Figure 4A). Our results place the RHON1 binding site between 88 and 133 nucleotides downstream of the rbcL termination codon (Figure 4A). This RHON1 binding site covers the region between the two 3'-ends of monocistrionic rbcL assayed by (CR)RT-PCR (−94/95 and −129 with respect to the termination codon of rbcL). RHON1 protein was also found to bind to the corresponding single-stranded DNA of its RNA target (Figure 4A), which is similar to the nucleotide binding activity of Rho factors (Galluppi and Richardson, 1980; Skordalakes and Berger, 2006).

To further test the RHON1-RNA interaction in vivo, RNAs that coimmunoprecipitated with the anti-RHON1 antibody were analyzed by slot-blot hybridization using probes corresponding to the different regions of the rbcL-accD intergenic spaces.

Figure 2. Plastid Gene Expression of the rhon1-2 Mutants.

(A) RNA gel blot analysis of rbcL polycistrionic transcript. A diagram of the rbcL polycistrionic operon and the locations of the probes (a to f) are shown at the top. Blots with reduced exposure time to x-ray film for probes a and b are also included (indicated as a’ and b’). The mature forms of RNA are indicated with asterisks to the left of each blot.

(B) Run-on transcription assay of plastid genes in rhon1-2 and wild-type plants. DNA probe (200 ng) was used for hybridization with labeled transcripts. Another run-on experiment with 100 ng DNA probe was also performed and similar results were observed (Supplemental Figure 2).
The coding region and 3'-UTR of *rbcL* mRNA were detected in substantial amounts in the precipitate, whereas the coding region and 5'-UTR of *accD* were not detected (Figure 4B). This analysis suggests that RHON1 may bind to the monocistronic *rbcL* mRNA but not to the *rbcL-accD* polycistronic precursor or the monocistronic *accD* mRNA. Subsequently, a ribonuclease-protection assay was used to map the 3'-ends of the RNAs that coimmunoprecipitated with RHON1. Our results showed that the RHON1-coimmunoprecipitated RNA was enriched in monocistronic *rbcL* rather than polycistronic *rbcL* precursor (Figure 4C). In addition, the ratio of *rbcL* mRNA ending at position 129 compared with that ending at 94/95 was increased in the coimmunoprecipitated RNAs.

**RHON1 Has Transcription Termination Catalytic Activity**

Transient expression of RHON1-green fluorescent protein (GFP) fusions in protoplasts of *Arabidopsis* gave rise to fluorescence signal exclusively in plastids, suggesting that RHON1 is targeted to the plastids (Figure 5A). Nevertheless, the fluorescence signal of RHON1 inside chloroplasts showed punctuate patterns resembling the distribution of nucleoids visualized by GFP constructs fused with nucleoid proteins on the red background of the chlorophyll autofluorescence (Terasawa and Sato, 2005; Zhong et al., 2013). When RHON1-GFP fusions were transiently coexpressed with red fluorescent protein fusions of pTAC5 protein, a plastid nucleoid-localized protein in protoplasts of *Arabidopsis* (Zhong et al., 2013), we found that green and red fluorescence signals within the chloroplasts were merged (Figure 5B), indicating that RHON1 as well as pTAC5 are localized in chloroplast nucleoids. Immunoblot analysis of chloroplast subfractions showed that RHON1 is mainly localized in the stromal fractions but is absent or present only at low levels in thylakoid and envelope fractions (Figure 5C). This result agrees with previous reports indicating that some stromal proteins accumulate in nucleoids based on GFP assays or mass spectrometry analysis of nucleoids (Majeran et al., 2012; Pfalz and Pfannschmidt, 2013).

Rho is a homohexameric protein with RNA-dependent ATPase and RNA/DNA helicase activities and is distributed throughout the bacterial world with very few exceptions (Allison et al., 1998; Aravind and Koonin, 2000). The first 125 residues of *E. coli* Rho comprise a separate domain that by itself can bind to oligonucleotides, whereas the remaining 290 residues are closely related to the sequences of the corresponding parts of the α- and β-subunits of the F,ATPase (Skordalakes and Berger, 2006). The C terminus of RHON1 displays high similarity to the RNA binding region of Rho, whereas no domain related to F,ATPase was found in this protein. However, a domain of ~100 residues close to the Rho-like domain showed low similarity to the P-type ATPase (Figure 6A).

![Figure 3. The Transcription Initiation Sites of Plastid Genes in Wild-Type and rhon1-2 Plants.](image)

(A) Identification of the transcription initiation site of *rbcL* by 5'-RACE. Total RNA from green (without spectinomycin treatment) and white (with spectinomycin treatment) leaf tissue was ligated to an RNA linker after TAP treatment and, along with untreated RNA as a control, was subjected to RT-PCR with specific primers. The transcription initiation sites identified by sequencing of cloned RT-PCR products are underlined and marked by arrows in the diagram of the 5' region of *rbcL*. The expression of the smallest PCR product in TAP-treated leaves was lower than that in leaves without TAP treatment. Therefore this product most likely corresponds to a processing product rather than to a primary transcript (Hanaoka et al., 2003; also refer to the ~70 band of [C]).

(B) Identification of the transcription initiation sites of *accD*. The procedure was similar to that used for *rbcL*.

(C) Primer extension analysis of *rbcL* and *accD* from wild-type and *rhon1-2* plants. The positions of bands corresponding to transcripts with distinct 5'-ends are indicated at the right.
as Rho factors. We investigate whether RHON1 possesses a similar ATPase activity potential P-type ATPase domain in RHON1 prompted us to investigate whether RHON1 possesses a similar ATPase activity as Rho factors. We first expressed the ATPase domain (AD) of RHON1 alone and examined its ATPase catalytic activity of hydrolysis of ATP to ADP and inorganic phosphate. No obvious ATPase activity of the AD domain was found, whether or not RNA substrates were added. However, when the AD domain of RHON1 was expressed together with the RNA binding domain, a high ATPase activity could be detected in the presence of RNA substrates (Figure 6B). In addition, when the conserved residues for RNA binding were mutated, the ATPase activity of RHON1 was completely eliminated (Supplemental Figure 5). These results clearly indicate that RHON1 indeed has RNA-dependent ATPase activity resembling that of the Rho factor.

We next assayed the transcription termination activity of RHON1 using an in vitro transcription termination assay. In this experiment, full-length RHON1 in which the maltose binding protein (MBP) tag was removed was incubated with DNA templates in an in vitro transcription system with the T7 RNA polymerase (Figures 6C and 6D). When the rbcL-accD intergenic region was used as template, the terminated transcript could be clearly observed among the transcription products in the presence of RHON1, whereas no terminated transcript was detected with the MBP proteins (Figure 6D). When the rm 23-rrn 4.5 intergenic region was used as template, no terminated transcript was observed even when the reaction time was extended to 3 h (Figure 6D). Taken together, these results suggest that RHON1 can specifically terminate rbcL transcription in vitro.

The similarity between the activities of RHON1 and Rho raised the possibility that RHON1 could be a functional analog of the E. coli Rho factor. To test this hypothesis, we introduced RHON1 into a conditional-lethal rho mutant of E. coli termed HD152. The HD152 strain is temperature sensitive; it dies at the restrictive temperature of 42°C and survives at the permissive temperature of 37°C (Das et al., 1976; Inoko et al., 1977). As shown in Figure 6E, the HD152 strain that was transformed with RHON1 as well as with Rho could grow at both 42 and 37°C. As a control, the HD152 strain transformed with GST (glutathione S-transferase) protein could only grow at 37°C. This result clearly demonstrates that RHON1 can functionally replace the E. coli Rho protein in vivo.

The RHON1 Homolog of Rice Cannot Replace RHON1 Function in rbcL Transcription Termination

There is a major difference related to the operon organization associated with the rbcL and accD loci between eudicotyledonous and monocotyledonous plants, as the accD gene has been transferred to the nuclear genome in monocots (Bryant et al., 2011; Supplemental Figure 6A). This raises the question of whether this transcriptional interplay between rbcL and its downstream gene occurs in other plant species besides Arabidopsis. To answer this question, we first compared the 3’-UTRs in a range of plants for which the plastid genome sequence is available. This analysis revealed that two conserved boxes of RHON1 binding nucleotides exist in the examined dicotyledonous but not in monocotyledonous plants (Figure 7A). In addition, the EMSA analysis showed that RHON1 does not bind to the 3’-UTR of rice rbcL (Figure 7B). These results suggest that the cis-elements of rbcL involved in the binding of RHON1 were lost in monocotyledonous plants.

BLAST analysis revealed that RHON1 homologs exist ubiquitously in vascular plants, but no corresponding RHON1 homolog was found in the unicellular alga Chlamydomonas reinhardtii. This suggests that RHON1 is a vascular plant-specific
protein. In correspondence with the evolution of rbcL 3'-UTRs, phylogenetic analysis of the aligned putative homologs of RHON1 revealed two well-supported clades which correspond to eudicotyledons and monocotyledons (Supplemental Figure 6B and Supplemental Data Set 1). This phylogenetic analysis points to a substantial biochemical difference in RHON1 homologs from eudicotyledons and monocotyledons. Indeed, the detailed analysis of the protein sequence of the rice RHON1 homolog Os-BP-73 revealed that the ATPase domain was lost and only the RNA binding domain was retained in Os-BP-73. Accordingly, no ATPase activity of Os-BP-73 was detected under the same conditions used for the At-RHON1 assay (Supplemental Figure 7A). In addition, Os-BP-73 was unable to complement the temperature-sensitive phenotype of the rho mutant strain of E. coli (Supplemental Figure 7B), which suggests that it cannot substitute for the Rho factor of E. coli in vivo.

The distinct biochemical features of Os-BP-73 suggest that it acts in a different manner in vivo compared with RHON1. To address this possibility, we expressed Os-BP-73 in the rho1 background and investigated whether it was able to complement the molecular defects of rho1 plants completely. As shown in Figure 7C, expression of full-length Os-BP-73 cDNA under the control of the constitutive 35S promoter in rho1-2 mutants was able to rescue the high chlorophyll fluorescence phenotype of rho1-2, suggesting that Os-BP-73 was functionally expressed in rho1-2. However, the abundance of transcripts from genes downstream of rbcL in the Os-BP-73/rho1-2 plant was still maintained at a high level, rather than the wild-type level (Figure 7C), suggesting that inefficient transcriptional termination still occurred in Os-BP-73/rho1-2. In contrast to Os-BP-73/rho1-2 plants, the accumulation of those transcripts was restored to wild-type levels in plants complemented with RHON1 (Supplemental Figure 8). Taken together with the in vitro functional analysis of Os-BP-73, we propose that unlike At-RHON1, Os-BP-73 has lost its ability to terminate transcription of rbcL but retained the function related to RNA processing during evolution.

DISCUSSION

RHON1 Is Involved in Transcription Termination of rbcL in Addition to RNA Processing

This study and that of Stoppel et al. (2012) show that a large rbcL polycistronic transcript normally undetectable in wild-type plants is present in two rho1 mutants from different genetic backgrounds (Figure 2A). The accumulation of the rbcL polycistronic precursor might result from processing defects in the rbcL-accD intercistronic region, as suggested by Stoppel et al. (2012). Based on the data presented here, we propose that inefficient transcription termination may account for the over-accumulation of rbcL polycistronic transcripts in rho1-2. At present, neither of these possibilities can be excluded, suggesting
Figure 6. RHON1 Has Transcription Termination Activity.

(A) Schematic diagram of ATPase domain and RNA binding domain of RHON1.

(B) The RNA-dependent ATPase activity of RHON1 protein. The C-terminal region of RHON1 containing ATPase domain (RHON1-AD) and the ATPase domain followed with RNA binding domain (RHON1-AD-RD) were each fused with a MBP tag and expressed in E. coli BL21 (DE3). The purified recombinant proteins with or without poly (rC) were used for the ATPase activity assay by measuring phosphate release. The MBP protein was used as a control. Error bars represent the se of the mean (n = 6).

(C) Preparation of recombinant RHON1 proteins used for the in vitro transcription termination assay. The MBP-RHON1 recombinant protein was expressed in BL21 (DE3) pLysS cells. Following purification, the MBP tag of the recombinant protein was removed with TEV protease.

(D) In vitro transcription termination assay of RHON1. The transcription reaction was performed by the T7 RNA polymerase for 15 min, 30 min, and 3 h. The localization of the probe on the plastid genome is shown on the top lane, and the numbers indicate the positions of the nucleotides with respect to the termination codon of rbcL and the 3’ end of rrn 23. MBP and BSA proteins were used as controls.
that RHON1 might have a function in processing as well as in transcription termination. In fact, RNA polymerase II transcription termination is coupled to 3′-end processing of the pre-mRNA in eukaryotes, and 3′ cleavage and polyadenylation factors are also critical for efficient transcriptional termination of RNA polymerase II (Rondon et al., 2008; Richard and Manley, 2009). Although the coupling of plastid RNA polymerase transcription termination and plastid RNA processing has not been addressed, it is possible that RHON1 acts as a factor coupling these two processes in plastids.

It has been proposed that RHON1 forms a distinct 800-kD complex with RNE and supports RNE function, by conferring sequence specificity to the endonuclease (Stoppel et al., 2012). However, the rhon1 me double mutant displayed a more severe pleiotropic phenotype than that of me, suggesting that RHON1 has additional functions compared with RNE (Stoppel et al., 2012). Indeed, investigation of accD promoter usage in me mutant plants showed that the aberrant transcription initiation observed in rhon1-2 did not occur in me (Supplemental Figure 9). This difference suggests that RHON1 also acts on the rbcL-accD intercistronic region independently of RNE. In addition to the 3′-UTR of rbcL, RHON1 was able to bind multiple RNA species, including rRNAs and petB, in vitro (Stoppel et al., 2012). These multiple functions of RHON1 might be related to its ability to bind multiple protein/RNA partners. RHON1 may form different complexes by interacting with distinct protein/RNA partners, as revealed by the multiple RHON1 complexes fractionated by native acrylamide gradient gel electrophoresis (Stoppel et al., 2012). Those complexes may act independently or cooperate with each other to function in two steps of plastid gene expression.

Possible Mechanisms of Transcription Termination in Plastids

There are conflicting observations concerning transcription termination in plastids of vascular plants. The spinach (Spinacia oleracea) RNA polymerase has been shown to recognize prokaryotic Rho-independent transcription terminators in vitro, suggesting that equivalent sequences might function as terminators of plastid transcription in vivo (Chen and Orozco, 1988). However, transcription termination assays using an in vitro transcription system from spinach showed that plastid 3′ inverted repeats of psbA, rbcL, and rpoB do not function as transcription terminators (Stern and Gruissem, 1987). Instead, two tRNAs, which are not followed by 3′ inverted repeats, do effectively terminate transcription under the same conditions (Stern and Gruissem, 1987). These discordant results raise the possibility that site-specific components are required for proper termination processes in vivo (Blowers et al., 1993). Alternatively, plastid transcription termination could also be mediated in as yet unknown ways that are not associated with 3′ inverted repeats.

The termination of rbcL transcription was originally inferred from the transformation of the plastid genome of tobacco (Nicotiana tabacum) with the rbcL gene (Staub and Maliga, 1994). The tobacco rbcL mRNA is always monocistronic when transcribed in its native context (Sinagawa-García et al., 2009). However, when chimeric genes were introduced into the intergenic region between rbcL and accD of tobacco, rbcL read-through transcripts were readily detectable (Staub and Maliga, 1994; Monde et al., 2000; Sinagawa-García et al., 2009). In addition, when the insertion site of the chimeric genes was moved further downstream, read-through transcription did not occur (Sinagawa-García et al., 2009). These results suggest the presence of a termination or processing site upstream of accD.

Considering the transcriptional independence of rbcL and accD in wild-type Arabidopsis plants, together with the data presented here, we suggest that there is a termination site in the intergenic region between rbcL and accD, although the presence of a processing site cannot be completely excluded. Our results show that efficient transcription termination of rbcL requires the RNA binding protein RHON1, related to bacterial Rho factors. Nucleotide binding assays and ATPase activity analysis suggest that RHON1 acts like Rho factors and is involved in transcription termination of plastid genes. Rho factors usually bind to a wide range of mRNA sequences but have a strong preference for mRNA sequences enriched in rC residues (Graham, 2004). However, the RHON1 binding mRNA region contains few rC residues, which is clearly different from the RNA binding sites of Rho factors. In addition, the ATPase domain of RHON1 does not belong to F-type ATPase and the protein structure is distinct from that of Rho. Therefore, it appears that the mechanism of action of RHON1 is not totally identical to that of Rho factors. Although the Rho-dependent mechanism has been established only in eubacteria, a similar ATP-driven termination machine was also discovered in eukaryotic cells. For instance, a Drosophila melanogaster ATPase, called factor2, and the Sen1p helicase in budding yeast both employ this ATP-driven mechanism to terminate RNA Polymerase II transcription (Xie and Price, 1997; Porrua and Libri, 2013). Therefore it is quite possible that RHON1 uses an unknown ATP-driven device to mediate the dissociation of nascent rbcL transcripts from PEP. A postulated mechanism for RHON1 action is schematically depicted in Figure 8.

In rhon1-2 plants, we observed aberrant transcription initiation in the rbcL-accD intergenic region. It is likely that this is a consequence of inefficient transcription termination of rbcL because it did not occur when rhon1-2 plants were grown on spectino-mycin, which leads to the inhibition of rbcL transcription. This suggests that these processes are coupled. Currently, the
molecular basis for the aberrant accD transcription initiation in rhon1-2 remains unclear. The elongating RNA polymerase complex across the termination site of rbcL might interfere with the initiation transcription complex of the accD transcriptional unit and thus reduce the transcription accuracy especially since plastid transcription initiation sites in vascular plants are not very stringent (Holec et al., 2006; Swiatecka-Hagenbruch et al., 2007). Alternatively, because after termination of rbcL transcription, PEP cannot be released from the template due to the lack of RHON1, it might slide on the template and finally reinitiate transcription nonspecifically at different positions in the rbcL-accD intergenic region and even within the accD gene. Thus, one of the functions of RHON1 may be to avoid aberrant transcription initiation in the rbcL-accD intergenic region due to inefficient rbcL transcription termination.

**Evolution of RHON1-Dependent Transcription Termination**

It is believed that the evolution of land plant organelle genomes is inextricably linked to the evolution of nuclear-encoded trans-acting factors, and vice versa. Plastid RNA editing has been one of the most enigmatic processes in this respect (Takenaka et al., 2013). In this study, both in vitro and in vivo evidence showed that Os-BP-73, the rice ortholog of RHON1, appears to have lost the ability to terminate transcription due to the lack of the ATPase domain. The RNA processing activity of Os-BP-73 is mediated through its conserved RNA binding domain could support RNE functions presumably by conferring sequence specificity to the endonuclease. In agreement with the difference between At-RHON1 and Os-BP-73, the putative RNA binding sites of RHON1 within in rbcL-accD intergenic region of the Arabidopsis plastid genome are absent from the rice plastid genome (Figures 7A and 7B), which suggests coevolution between At-RHON1/ Os-BP-73 and its target elements. Our results indicate that the loss of a cognate cis-element might not result in the complete loss of its corresponding trans-acting factor but lead to the elimination of its related domain, which might represent a more economical strategy for evolution.

One fundamental difference between plastid genomes in dicotyledons and monocotyledons is that three genes, ycf1, ycf2 (functions unknown), and accD, are present in dicotyledons but absent from monocotyledon plastids. The failure to translate one or more of these three genes is likely to be the reason for embryo lethality in Arabidopsis but not in maize (Zea mays), which:

![Figure 7](https://example.com/figure7.png)

**Figure 7.** The RHON1 Homolog of Rice Cannot Replace RHON1 Function in rbcL Transcription Termination.

(A) Comparison of RHON1 binding site sequences in the rbcL 3′-UTR between selected monocots and eudicots. Two conserved regions are boxed.

(B) EMSA assay of the interaction of RHON1 with the rbcL 3′-UTR of rice.

(C) Photographs and chlorophyll fluorescence images of Os-BP-73/rhon1-2 and wild-type plants grown for 3 weeks in a growth chamber. Fluorescence is shown using the pseudocolor index indicated at the bottom.

(D) Accumulation of accD, ycf4, cemA, and Os-BP-73 in Os-BP-73/rhon1-2 and wild-type plants.
suggests that the control of the expression of these genes is critical for plant development in dicotyledons (Asakura and Barkan, 2006; Schmitz-Linneweber and Small, 2008; Bryant et al., 2011). Therefore, it is not surprising that some specific factors have been recruited to serve the expression of these genes in dicotyledons, for instance, the species-specific editing factors for accD (Robbins et al., 2009; Yu et al., 2009). In rhon1 plants, the accumulation of rbcL transcripts was normal, whereas the transcription initiation of accD was affected. Our results thus point to an alternative strategy for regulating accD expression in dicotyledons, which relies on the efficient transcription termination of its upstream gene, rbcL, mediated by RHON1.

METHODS

Plant Materials and Growth Conditions

The rhon1-2 mutant was isolated from a collection of pSKI015 T-DNA-mutagenized Arabidopsis thaliana (ecotype Columbia) lines based on its high chlorophyll fluorescence phenotype. The seeds of wild-type and mutant plants were surface-sterilized, sown on Murashige and Skoog medium containing 2% sucrose, and grown under short-day conditions. Mutant plants were surface-sterilized, sown on Murashige and Skoog medium containing 50 µg mL⁻¹ hygromycin B.

Chlorophyll Fluorescence Measurements

Chlorophyll fluorescence measurements were performed using a CF Imager (Technologica) as described by the manufacturer. Before each measurement, leaves were dark adapted for 10 min. The minimum fluorescence yield (Fₒ) and the maximum fluorescence yield (Fₘₐₓ) were measured. The maximal photochemical efficiency of photosystem II was calculated from the ratio of Fₚ to Fₘₐₓ [Fₚ/Fₘₐₓ = (Fₘₐₓ − Fₒ)/Fₘₐₓ]. Image data acquired in each experiment were normalized to a false color scale, with arbitrarily assigned extreme values of 0.1 (lowest) and 0.8 (highest). This resulted in the highest and lowest Fₚ/Fₘₐₓ values being represented by the red and blue extremes of the color scale, respectively.

RT-PCR, Quantitative PCR, and RNA Gel Blotting

Total leaf RNA was extracted from Arabidopsis seedlings with the Qiagen RNeasy Plant Mini kit (Qiagen) and used to generate first-strand cDNA in a 50-µL reaction with the Superscript III cDNA synthesis system (Invitrogen) following the manufacturer’s instructions. Aliquots of the resulting cDNA samples were used as template for PCR analysis. Quantitative PCR was performed with SYBR PrimeScript Ready Mix (Takara) in an ABI 7900 sequence detection system (Applied Biosystems) according to the manufacturer’s protocol. Total RNA (5 to 10 µg) from wild-type and rhon1-2 seedlings was separated on 1.2 to 2.0% (w/v) agarose-formaldehyde gels, blotted to positively charged nylon membrane by capillary blotting, fixed by cross-linking, and subsequently subjected to hybridization. Hybridization probes were prepared by PCR amplification of fragments of the plastid genes from Arabidopsis. The sequences of all primers used in this study are listed in Supplemental Table 1.

Run-on Transcription Assay

The chloroplast run-on assays were performed using DIG-11-UTP as the labeling nucleotide as described (Chi et al., 2010). Briefly, the leaves were homogenized in buffer containing 50 mM HEPES-KOH, pH 6.8, 0.33 M sorbitol, 2 mM EDTA, 1 mM MgCl₂, 1 mM MnCl₂, and 9 mM DTT. The homogenate was filtered through four layers of cheesecloth and spun at 3000g for 5 min at 0°C. The pellet was fractionated in 20/50/80% Percoll gradients by centrifugation at 6500g for 15 min. Intact chloroplasts were collected at the interface between 50 and 80% Percoll, the integrity of chloroplasts was examined by CP-ISO kits supplied by Sigma-Aldrich, and the number of chloroplasts in the samples was determined by counting the organelles using light microscopy. Transcription was performed in 100-µL reactions containing 5.0 x 10⁻⁶ M radiolabeled nucleotides for 10 min, followed by RNA extraction with phenol/chloroform. Labeled transcripts were hybridized to Hybond N⁺ filters carrying 100 to 200 ng DNA probes of plastid genes. A slot-blotting apparatus (Bio-Dot SF; Bio-Rad) was used for filter preparation. Signals were analyzed using ALPHAVIEW 3.0 (Alpha Innotech).

GFP Assays

The coding sequences of RHON1 were amplified by RT-PCR. The PCR products were then cloned into Xhol/BamHI sites of the transient expression vector pSAT6-EGFP-N1 (Citovsky et al., 2006), which contained the mGFP5 coding sequence under the control of the cauliflower mosaic virus 35S promoter, to generate a RHON1 fusion protein with GFP in the C terminus. Similarly, two controls were then constructed: one in which the floral dip method (Clough and Bent, 1998), respectively. The transformants were selected on Murashige and Skoog medium containing 50 µg mL⁻¹ hygromycin B.
nuclear localization signal of the PTM-N (Sun et al., 2011) and the other in which the transit peptide of chloroplast inner envelop translocation component (TIC21) of Arabidopsis were also fused to the vector. The resulting constructs were transfected into Arabidopsis mesophyll protoplasts according to the method as described by Koytun et al. (2000). Fluorescence analysis was performed on an LSM 510 META confocal laser scanning system (Zeiss).

Expression of Recombinant Proteins and Antibody Production

To produce recombinant MBP-RHON1-Strep proteins, the coding sequence of RHON1 lacking the transit peptide sequence was PCR amplified, digested with SalI and BamHI, and inserted into the SalI/BamHI sites of pMAL-TEV (kindly provided by Alice Barkan), BL21(DE3) pLysS cells (Novagen) harboring the resulting vector were grown at 37°C until OD600 of 0.6 was reached and transferred to 22°C for 30 min. Expression was then induced by 1 mM isopropyl β-D-1-thiogalactopyranoside, and cells were grown for 3 h at 22°C with shaking. All subsequent steps were performed at 4°C. Sedimented cells were lysed in lysis buffer (20 mM Tris-HCl, pH 7.4, 350 mM NaCl, and Complete Protease Inhibitor Cocktail Tablets [Roche]) by sonication and cleared by centrifugation (36,000g, 20 min). Affinity chromatography was performed using amylase resin (NEB). After three washes with lysis buffer, proteins were eluted in Strep-washing buffer (IBA) supplemented with 150 mM maltose and subjected to a second affinity purification using Strept-Tactin-Macroprep (IBA). The column was washed three times with Strep-washing buffer, and proteins were eluted with Strep-elution buffer (IBA). RHON1-Strep was subsequently cleaved from the MBP moiety using AcTEV protease (Life Technologies). The DNA sequences coding for three RHON1 fragments (63 to 200, 201 to 300, and 301 to 401 amino acids) were subcloned into the EcoRI/Xhol sites of the expression vector pMAL-c5x (NEB) to produce MBP fusions. Induction, cell lysis, and purification of the fusion proteins using amylase affinity chromatography were performed according to the manufacturer’s protocol (NEB). A RHON1 fragment (64 to 200 amino acids) with a C-terminal 6× histidine tag was expressed from a pET28a vector (Novagen) and purified with nickel-nitritolitratric acid agarose. The purified protein was used to generate polyclonal antibodies against RHON1 in rabbits.

Protein Isolation and Immunoblot Analysis

Total protein of Arabidopsis leaves was isolated as previously described (Chi et al., 2010). Stroma, thylakoid, and envelope fractions were purified from lysed chloroplasts by sucrose gradient centrifugation as previously described (Williams and Barkan, 2003). The protein levels were quantified using the DC protein assay kit (Bio-Rad). The proteins resolved by SDS-PAGE were blotted onto nitrocellulose membranes and incubated with specific antibodies, and the signals were detected using the enhanced chemiluminescence method.

Mapping of Transcription Initiation Sites

The transcription initiation sites of plastid genes were determined using a 5′-RACE Full-RACE kit (Takara) based on the methods of Swiatecka-Hagenbruch et al. (2007) with minor modifications. Briefly, the total RNA was treated with calf intestine alkaline phosphatase to remove free 5′ phosphates. After extraction, 2 μg purified RNA was then treated with 0.5 units of TAP at 37°C for 1 h in the presence of 40 units of RNase inhibitor. Control reactions were set up without TAP. The RNA was subsequently purified and ligated to a 5′ RNA adapter according to the supplier’s protocol. PCR products were analyzed on agarose gels after two successive PCR amplifications, using two outer primers and two inner primers, respectively. The inner and outer adapter-specific forward primers were included in the Full-RACE kit.

RNA Coimmunoprecipitation Analysis

The stroma was prepared by hypotonic lysis of purified chloroplasts and used for RNA coimmunoprecipitation analysis as described by Barkan (2009). RNA was recovered from the precipitated fractions in the presence of 1% SDS, 0.05 μg μL−1 yeast tRNA, and 5 mM EDTA. The RNA was isolated by phenol–chloroform isooamyl alcohol extraction and precipitated with 3 volumes of ethanol. RNAs from the supernatant fractions were extracted as the controls. Equal portions of each RNA sample were applied to a nylon membrane through a slot-blot manifold (Bio-Dot SF; Bio-Rad) and hybridized with radiolabeled DNA probes specific for the indicated rbCL and accD sequences. Probes were generated by PCR amplification of the relevant chloroplast sequences and labeled by the random priming method. The ribonucleicase-protection assay of the RNA samples was performed as described by Chi et al. (2012).

Primer Extension

Primer extension was performed using the Primer Extension System of Promega. Sequence-specific oligonucleotides were end-labeled with [γ-32P]ATP and then were annealed with 10 μg total RNA for 20 min at the primer annealing temperature. Primer extension was performed for 30 min at 42°C with AMV reverse transcriptase. Extension products were heated at 90°C for 10 min and were separated on a 6 × 5-cm denaturing polyacrylamide gel containing 6% acrylamide (19:1 acrylamide: bis), 7 M urea, and 1× TBE buffer. The signals were visualized by autoradiography.

EMSA

DNA/RNA and protein interactions were detected using the LightShift Chemiluminescent EMSA kit (Pierce). Briefly, the target DNA/RNA was 5′-end-labeled with biotin and terminal deoxynucleotidyl transferase and then incubated with the RHON1 fragments (63 to 200, 201 to 300, and 301 to 401 amino acids) in the DNA/RNA binding reaction mixture supplied by the kit. After incubation for 20 min at 25°C, the samples were resolved on a 6% Tris-borate gel in 0.5× TBE buffer and transferred to a nylon membrane. The biotin end-labeled DNA was detected using a streptavidin-horseradish peroxidase conjugate and chemiluminescent substrate.

ATPase Activity Assay

The ATPase activity was determined by measuring phosphate release in a mixture containing 50 mM Tris-HCl, pH 8.0, 50 mM NaCl, 4 mM ATP, and 2 mM MgCl2, incubated with 0.2 μL reconstitution mixture for 5 min at 37°C. After the reaction was stopped by adding 40 μL trichloroacetic acid, the supernatant was collected and the released Pi was determined by a microcolorimetric method (Taussky and Shorr, 1953).

Complementation of the rho Mutant of Escherichia coli

The rho mutant (HD152) was obtained from the E. coli Genetic Stock Center at Yale University. The coding sequence of RHON1 and Os-Bp-73 lacking the transit peptide sequences were inserted into the EcoRI and Xhol sites of pGEX-4T-2 (GE Healthcare) to produce GST fusions. RHO of E. coli was inserted into the Xmal and Xhol sites of pGEX-4T-2 and used as a control. These constructs then were transformed into HD152 cells by the CaCl2 method. The transformed cells were grown in the presence of 0.4 mM β-1-thiogalactopyranoside. The temperature sensitivity of E. coli cells was assayed as described by Inoko et al. (1977).

In Vitro Transcription Termination Assay

Rbcl-accD (nucleotides 56434 to 56632; GenBank accession number AP000423.1) and Rm23-mm4.5 (nucleotides 107501 to 107598; GenBank...
accession number AP000423.1) intergenic regions were PCR amplified. Amplification products were gel purified. In vitro transcription was performed in 20-µL reactions containing 300 pM each PCR product, 0.5 mM each ATP, CTP, and GTP, 10 µM UTP, 10 µCi [α-32P]UTP, 20 units T7 RNA Polymerase (Fermentas), 4 µL 5× transcription buffer (200 mM Tris-HCl, pH 7.9, 30 mM MgCl2, 50 mM DTT, 50 mM NaCl, and 10 mM spermidine; Fermentas), 20 units Ribloc RNase Inhibitor (Fermentas), and protein concentrations as indicated in the figure. Reaction mixtures were incubated for 15 min, 30 min and 3 h at 30°C, denatured by heating in the presence of 80% formaldehyde, and analyzed on 5% denaturing polyacrylamide gel containing 8 M urea and 1× TBE. Gels were exposed to a phosphor imager screen and analyzed with the Typhoon Variable Mode Imager (GE Healthcare).

**Accession Numbers**

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: RHON1, AT1G06190; accD, ATCG00500; rbcL, ATCG00490; and OsBP-73, Os03g0183100.

**Supplemental Data**

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

- Supplemental Figure 1. Microarray Analysis of Plastid Gene Transcripts in rhon1-2 Compared with the Wild Type.
- Supplemental Figure 2. Run-on Transcription Assay of Plastid Genes in rhon1-2 and Wild-Type Plants.
- Supplemental Figure 3. Analysis of PEP-Dependent Transcription in rhon1-2.
- Supplemental Figure 4. Mapping of 3′- and 5′-Ends of Plastid Transcripts in Wild-Type and rhon1-2 Plants.
- Supplemental Figure 5. The ATPase Activities of Mutated RHON1 Proteins.
- Supplemental Figure 6. Phylogenetic Analysis of RHON1 Homologs in Vascular Plants.
- Supplemental Figure 7. Characterization of a RHON1 Homolog of Rice, Os-BP-73.
- Supplemental Figure 8. The Accumulation of rbcL, accD, psal, ycf4, cemA, and petA Transcripts in RHON1- Complemented Plants Assayed by RT-PCR.
- Supplemental Figure 9. The Transcription Initiation Sites of accD in the me Mutant.
- Supplemental Table 1. List of Primers Used in This Study.
- Supplemental Data Set 1. Text File of the Alignment Used for the Phylogenetic Analysis Shown in Supplemental Figure 6.

**ACKNOWLEDGMENTS**

We thank Alice Barkan for kindly sending us pMAL-TEV. We also thank the reviewers for important insights and for suggesting key experiments that changed and improved our conclusions and insights regarding RHON1 function. This work was supported by the National Natural Science Foundation of China (Grant 31370273), the Major State Basic Research Development Program (Grant 2015CB150106), and the Solar Energy Initiative and Cooperative Team Project of the Chinese Academy of Sciences.

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The authors of the above article request the publication of corrections to Figures 1 to 3, as described below, for clarification of the methods used and the manner in which gel images were arranged. The original results and conclusions are unaffected by these corrections. The authors apologize and accept responsibility for not detecting these errors prior to publication. Both original and corrected figures are shown for ease of comparison.

Figure 1D: In the corrected figure, two white vertical bars have been inserted between the rhon1-2 and rhon1-2 complemented lanes for the RHON1 and ACTIN panels, respectively, to indicate the splicing of separate lanes from the same gel in the figure. The legend of the figure has been changed accordingly for clarity.

Figure 2: In Figure 2A, the RNA gel blots with probes a and f used the same gel and nylon membrane (the first blot was stripped and reprobed with another probe); therefore, their ethidium bromide staining bands used as the loading controls were duplicated. This was also applied to the RNA gel blots of probes b and d. This duplication of loading controls is explained in the corrected legend. Images of Figure 2A have not been changed. Figure 2B has been corrected to indicate the splicing of the cemA and petA lanes of the wild-type panel by a black vertical bar. The legend of Figure 2B has been changed accordingly for clarity.

Figure 3B: The (+) spec and (−) spec samples were run on two separate gels and combined in the published figure. This combination was not necessary and affected the accuracy of data presentation due to the lack of a marker for this combination. Because the original data could not be found, Figure 3B was reconstituted with the data from duplicate experiments. Two separate gels from the duplicated experiments are displayed independently in the new version of Figure 3B shown below. The figure legend has not been changed.
Figure 1. Original: Phenotype and Gene Cloning of rhon1-2.

(D) RT-PCR (left) and immunoblot analysis (right) of RHON1 expression in wild-type, rhon1-2 mutant, and complemented plants.

Figure 1. Corrected: Phenotype and Gene Cloning of rhon1-2.

(D) RT-PCR (left) and immunoblot analysis (right) of RHON1 expression in wild-type, rhon1-2 mutant, and complemented plants. In the RT-PCR analysis, samples were run side-by-side on the same gel. White bars indicate noncontiguous lanes in the gel.
Figure 2. Original: Plastid Gene Expression of the rhon1-2 Mutants.

(A) RNA gel blot analysis of rbcL polycistronic transcript. A diagram of the rbcL polycistronic operon and the locations of the probes (a to f) are shown at the top. Blots with reduced exposure time to x-ray film for probes a and b are also included (indicated as a’ and b’). The mature forms of RNA are indicated with asterisks to the left of each blot.

(B) Run-on transcription assay of plastid genes in rhon1-2 and wild-type plants. DNA probe (200 ng) was used for hybridization with labeled transcripts. Another run-on experiment with 100 ng DNA probe was also performed and similar results were observed (Supplemental Figure 2).
Figure 2. Corrected: Plastid Gene Expression of the *rhon1-2* Mutants.

(A) RNA gel blot analysis of *rbcL* polycistronic transcript. A diagram of the *rbcL* polycistronic operon and the locations of the probes (a to f) are shown at the top. Blots with reduced exposure time to x-ray film for probes a and b are also included (indicated as a’ and b’). The mature forms of RNA are indicated with asterisks at the left of each blot. The blot of probe a was stripped and reprobed with probe f, while that of probe b was stripped and reprobed with probe d. Ethidium bromide (EtBr) staining of the RNA served as loading control.

(B) Run-on transcription assay of plastid genes in *rhon1-2* and wild-type plants. A DNA probe (200 ng) was used for hybridization with labeled transcripts. Another run-on experiment with a 100 ng DNA probe was also performed, and similar results were observed (Supplemental Figure 2). The *petA* signal of the wild-type sample, separated by a black bar, was obtained from a different slot-blot, but all of the blots were developed and processed in parallel.
Figure 3. Original: The Transcription Initiation Sites of Plastid Genes in Wild-Type and rhon1-2 Plants.

(A) Identification of the transcription initiation site of rbcL by 5′-RACE. Total RNA from green (without spectinomycin treatment) and white (with spectinomycin treatment) leaf tissue was ligated to an RNA linker after TAP treatment and, along with untreated RNA as a control, was subjected to RT-PCR with specific primers. The transcription initiation sites identified by sequencing of cloned RT-PCR products are underlined and marked by arrows in the diagram of the 5′ region of rbcL. The expression of the smallest PCR product in TAP-treated leaves was lower than that in leaves without TAP treatment. Therefore, this product most likely corresponds to a processing product rather than to a primary transcript (Hanaoka et al., 2003; also refer to the 70 band of [C]).

(B) Identification of the transcription initiation sites of accD. The procedure was similar to that used for rbcL.

(C) Primer extension analysis of rbcL and accD from wild-type and rhon1-2 plants. The positions of bands corresponding to transcripts with distinct 5′-ends are indicated at the right.
Figure 3. Corrected: The Transcription Initiation Sites of Plastid Genes in Wild-Type and rhon1-2 Plants.

Editor's note: the corrected figures and accompanying text were reviewed by members of The Plant Cell editorial board.
RHON1 Mediates a Rho-Like Activity for Transcription Termination in Plastids of *Arabidopsis thaliana*

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