Unique Translation Initiation at the Second AUG Codon Determines Mitochondrial Localization of the Phage-Type RNA Polymerases in the Moss Physcomitrella patens

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The nuclear genome of the moss Physcomitrella patens contains two genes encoding phage-type RNA polymerases (PpRPOT1 and PpRPOT2). Each of the PpRPOT1 and PpRPOT2 transcripts possesses two in-frame AUG codons at the 5′ terminus that could act as a translational initiation site. Observation of transient and stable Physcomitrella transformants expressing the 5′ terminus of each PpRPOT cDNA fused with the green fluorescent protein gene suggested that both PpRPOT1 and PpRPOT2 are not translated from the first (upstream) AUG codon in the natural context but translated from the second (downstream) one, and that these enzymes are targeted only to mitochondria, although they are potentially targeted to plastids when translation is forced to start from the first AUG codon. The influence of the 5′-upstream sequence on the translation efficiency of the two AUG codons in PpRPOT1 and PpRPOT2 was quantitatively assessed using a β-glucuronidase reporter. The results further supported that the second AUG codon is the sole translation initiation site in Physcomitrella cells. An Arabidopsis (Arabidopsis thaliana) RPOT homolog AtRpoT;2 that possesses two initiation AUG codons in its transcripts, as do the RPOTs of P. patens, has been regarded as a dually targeted protein. When the localization of AtRpoT;2 was tested using green fluorescent protein in a similar way, AtRpoT;2 was also observed only in mitochondria in many Arabidopsis tissues. These results suggest that, despite the presence of two in-frame AUGs at the 5′ termini of RPOTs in Physcomitrella and Arabidopsis, the second AUG is specifically recognized as the initiation site in these organisms, resulting in expression of a protein that is targeted to mitochondria. This finding may change the current framework of thinking about the transcription machinery of plastids in land plants.

Plant cells contain two organelles having semiautonomous genetic systems, namely, mitochondria and plastids. They arose from eubacteria-like endosymbionts, closely related to extant α-proteobacteria and cyanobacteria, respectively (Gray, 1992, 1993; Howe et al., 1992). During the course of evolution, they lost many of their genes, while a number of genes were transferred to the cell nucleus. Therefore, the majority of the proteins involved in the biogenesis of mitochondria and plastids are encoded in the nucleus, and must be translated in the cytoplasm and then imported into the respective organelles (Schatz and Dobberstein, 1996; Neupert, 1997; Soll and Tien, 1998; Keegstra and Cline, 1999). Most of these proteins have a transit peptide at the N terminus that is necessary for their import into the target organelles (von Heijne, 1986; van Loon et al., 1988; Hand et al., 1989; Ko and Cashmore, 1989; Sidorov et al., 1999). The localization of these organelar proteins has been analyzed by in vitro import experiments using isolated organelles and in vivo experiments using a reporter, such as green fluorescent protein (GFP).

Nuclear-encoded RNA polymerases (RPOTs) consisting of a single polypeptide, which are similar to the RNA polymerase of bacteriophages T3 and T7, are widely distributed among eukaryotes and act as mitochondrial RNA polymerases (Cermakian et al., 1996, 1997). In higher plants, there is an additional plastid-targeted RPOT (Hedtke et al., 1997; Chang et al., 1999; Kobayashi et al., 2002), which is also called NEP (for nuclear-encoded polymerase). Plant RPOTs constitute a small gene family with different targeting properties. For example, Arabidopsis (Arabidopsis thaliana) RpoT;1 (AtRpoT;1) is targeted to mitochondria (Hedtke et al., 1997, 1999), while AtRpoT;3 is targeted to plastids (Hedtke et al., 1997, 1999). In addition, AtRpoT;2 was postulated to be targeted to both organelles (Hedtke et al., 2000). AtRpoT;2 possesses the property of dual targeting by the use of two probable initiation codons. Hedtke et al. (2000) reported that the polypeptide translated from the first AUG codon was targeted to the plastids and mitochondria, whereas the polypeptide translated from the second AUG was targeted only to mitochondria. Dually targeted RPOT was also identified in Nicotiana sylvestris (NsRpoT-B; Kobayashi et al., 2001a), which also

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contains two putative translation initiation codons. The dual targeting is thought to be effected by alternative translation initiation within a single transcript (Kobayashi et al., 2001a). In the moss Physcomitrella patens, two RPO7 genes (named PpRPO1 and PpRPO2, respectively) have been identified (Kabeya et al., 2002; Richter et al., 2002). The translated sequence for each of these contains two putative translation initiation codons at the N terminus, just as do AtRpoT2 and NsRpoT-B (Fig. 1A). In our previous study, the two RPO7 proteins were targeted only to mitochondria (Kabeya et al., 2002). However, Richter et al. (2002) reported that the two RPO7 proteins were targeted to both mitochondria and chloroplasts.

In general, mRNA structure can influence translation initiation, e.g. the m7G cap, the length of the 5′-untranslated region (UTR), upstream open reading frame (uORF), the secondary structure of RNA, and the sequence context surrounding the initiation codon (Kozak, 1991), as well as interaction of the 5′- and the 3′-UTR (Bailey-Serres, 1999). Subcellular localization of several RPO7s was often examined with GFP-fusion protein, but the native 5′-UTR was not used in most of these targeting experiments, including the experiments by Richter et al. (2002). In our previous experiments, the native 5′-UTR was used (Kabeya et al., 2002). Effect of 5′-UTR on the translational start sites is a hypothesis that explains experimental discrepancy, and the resolution of this discrepancy will lead to physiologically important consequences on the framework of thinking about the plastid transcription machinery.

In this study, we examined the localization of the two PpRPO7s by immunoblot and enzymatic analyses, as well as targeting experiments using several GFP-fusion proteins. The results strongly indicated that the localization of these proteins is determined by the use of a unique initiation site, namely, both PpRPO1 and PpRPO2 proteins are translated from the second AUG codon and localized to mitochondria in Physcomitrella tissues. In addition, AtRpoT2, so far regarded as dual-targeting RPO7 in Arabidopsis, was also suggested to be localized only to mitochondria in wide plant tissues.

RESULTS

Presence of Two In-Frame AUG Codons in the 5′ Region of RPO7 Sequences

The cDNA sequence for the two RPO7s in P. patens that we published previously (Kabeya et al., 2002) showed a long 5′ sequence that preceded the conserved RPO7-coding sequence (Fig. 1A). Some cDNA sequences were also available in the expressed sequence tag database (Nishiyama et al., 2003), but all of them were mapped within the cDNA sequences shown in Figure 1. Genomic sequences were published by Richter et al. (2002), and the comparison of cDNA and genomic sequences indicated that the long 5′ sequence of PpRPO1 was shared by the two except two terminal residues (this could be due to cloning artifact or vector sequence), but that a short sequence fragment at the 5′ end of the genomic sequence of PpRPO2 might be an intron sequence, which ends by the consensus AG. In other words, the transcription starts from an exon further upstream whose genomic sequence is still not available. Both of the cDNAs are likely full-length ones, but the 5′ end was not mapped due to unavailability of upstream genomic sequences and the low level of transcripts in the total mRNA pool.

In both RPO7 sequences, the long 5′ sequence contains two in-frame ATG triplets. No further in-frame ATG is present upstream of the conserved polymerase sequence. Available sequence data indicate the transcript contains both AUG codons. No splicing variant containing only one of the AUGs is known. Therefore, the mRNA contains a long 5′-UTR (291 nucleotides) upstream of the first ATG in PpRPO1. The length of the 5′-UTR of PpRPO2 mRNA is 132 nucleotides. In addition, the interval between the first ATG and the second ATG was 141 and 105 nucleotides, respectively, for PpRPO1 and PpRPO2. In the AtRpoT2 sequence, the 5′-UTR is 231 nucleotides long, and the interval between the two ATG codons is 117 nucleotides.

Immunoblot Analysis and Effect of Tagetitoxin on the Organellar Transcription in P. patens

We first tested if the two PpRPO7s are present in the plastids by immunoblot analysis and by measuring the sensitivity to tagetitoxin of the transcription in isolated mitochondria and plastids of P. patens protonemata (Fig. 2, B and C, respectively). Tagetitoxin is known to inhibit transcription by plastid-encoded RNA polymerase (PEP) and bacterial RNA polymerase (Matthews and Durbin, 1990), but not the activity of RPO7. In P. patens, the core of PEP consists of plastid-encoded subunits (β, β′, and β″) and a nuclear-encoded α-subunit (Sugiiura et al., 2003), but is expected to be sensitive to tagetitoxin because its structure is analogous to that of bacterial RNA polymerases. Protonemata were used in these experiments because we already showed that the two PpRPO7 genes are transcribed in the protonemata (Kabeya et al., 2002). First, polyclonal antisera were raised against the glutathione S-transferase (GST)-PpRPO1 and GST-PpRPO2 recombinant proteins. (Primers for fusion constructs are summarized in Table I.) In these proteins, the N-terminal half of the protein sequence was used to avoid cross-reaction due to the conserved C-terminal active center domain. The cross-reaction of these antibodies with the full-length polymerase enzymes (expressed as His-tagged proteins that were used in the experiment in Fig. 2C) was indeed undetectable and estimated to be less than 1% (Fig. 2A). The two organelles were purified over Percoll gradients and then tested by immunoblot analysis.
Figure 1. Sequence context for the putative initiation codons of *PpRPOT1*, *PpRPOT2*, and *AtRpoT1*.  
A. The 5’ sequences upstream of the first and the second ATGs of *PpRPOT1*, *PpRPOT2*, and *AtRpoT1*. Nucleotide sequences of *PpRPOT1* (top), *PpRPOT2* (middle), and *AtRpoT1* (bottom). The first ATG codon is surrounded by a rectangle in each sequence. The second ATG codon is marked with underline. The 5’ end of the genomic sequences of *PpRPOT1* and *PpRPOT2* that diverge from the cDNA sequences are shown in uppercase letters. The extensive difference in the 5’ sequence in *PpRPOT2* could be due to an intron in the genomic sequence. 

B. Upstream context for the putative initiation codons. Top half shows actual sequences upstream of the respective ATG. The numbers above indicate nucleotide position with respect to the A of the putative initiation codon. Nucleotides corresponding to moss consensus sequence are indicated by uppercase letters. Bottom half shows moss and plant consensus sequences as well as scoring matrix for moss based on information content analysis. The score at the right side of the top half was calculated by adding respective values corresponding to the nucleotide sequence. The plant consensus was taken from Joshi et al. (1997). GenBank accession numbers are as follows: *PpRPOT1* cDNA, AB055214; *PpRPOT1* genomic, AJ416854; *PpRPOT2* cDNA, AB055215; *PpRPOT2* genomic, AJ416855; and *AtRpoT1* genomic, AJ001037.
with anti-PpRPOT1 or anti-PpRPOT2 (Fig. 2B). An immunoreactive protein with a molecular mass identical to that expected for PpRPOT1 (approximately 110 kD) or PpRPOT2 (approximately 108 kD) was detected in the mitochondrial fraction but not in the chloroplast fraction (Fig. 1B). Control antibodies directed against known plastid (SIR or sulfite reductase; Sato et al., 2001) and mitochondrial (PpGR3, an RNA-binding protein; Nomata et al., 2004) proteins were also tested to confirm the purity of the two fractions. No significant cross-contamination was found (Fig. 2B). These results suggest that both PpPROT1 and PpRPOT2 proteins are localized in the mitochondria but not in the chloroplasts in the protonema.

The two PpRPOTs were demonstrated to be functional RNA polymerases in a previous study (Kabeya et al., 2002). The transcription activity of the two recombinant PpRPOT enzymes was not inhibited by tagetitoxin (Fig. 2C). In mitochondria, the transcription was not inhibited by tagetitoxin or even stimulated to some extent (Fig. 2C). This result was just as expected because the two PpRPOTs are present in the mitochondria as described above, and no other type of RNA polymerase is known to function in mitochondria of higher eukaryotes. In the plastids, the transcription was almost completely (to less than 1%) inhibited by the addition of tagetitoxin. In the in vitro transcription system using the plastidial nucleioids of tobacco BY-2 cells, which contain an appreciable level of NEP, the residual transcription activity in the presence of tagetitoxin (about 50% of total activity) was ascribed to the plasmid of N. patens, however, there was no measurable level of tagetitoxin-insensitive transcription activity. This is consistent with the results of immunoblot analysis that indicated the absence of RPO1 enzymes in the chloroplasts. If the RPO1s are massively targeted to chloroplasts as reported by Richter et al. (2002), all these results are hard to explain. Rather, this is evidence that RPO1 enzymes or NEP is absent in the plastids of P. patens.

Targeting of PpRPOT1 and PpRPOT2

Next, we reexamined targeting of GFP-fusion proteins. The discrepancy in the targeting experiments, as described in the introduction, might be due to the constructs used in the two reports. In the experiments of Richter et al. (2002), translation was forced to start from the AUG codon just downstream the translation leader sequence, which was taken from tobacco etch virus (using the vector pOL S65C; Peeters et al., 2000). The native 5'-upstream sequence was not used in their targeting experiments. By contrast, we used GFP-fused constructs containing the native 5'-upstream sequences to investigate the situation in the natural translation context. Thus, the sequence context of the translational initiation site is a likely candidate for the discrepancy. Since the nucleotide sequence context of the AUG plays a role in the efficiency of translation initiation, the 5'-upstream sequences of the first and second AUGs were compared (Fig. 1B). Although the consensus sequence of the plant context is AAAAACAA(A/C)AUG (Joshi et al., 1997), the sequence context of both the first and second AUG were scarcely analogous to the consensus sequence, and it was difficult to estimate functional initiation codon just by such sequence comparison.
We therefore tried to obtain as much information as possible from the sequence database of *P. patens*. First, all available protein-encoding sequences (266 in total) were retrieved from the GenBank database. Then, the 20 nucleotides upstream of the initiation site were extracted from each entry. The information content of each site or SequenceLogo was calculated for these 214 UTR sequences and is presented in Figure 1B. Based on this statistic, we deduced moss consensus, as shown in the figure. The information content values were used as a weight matrix to calculate a score for each possible initiation site, which we hoped would represent the probability of the initiation site. The result (Fig. 1B, right column) indicated that the score was higher for the second AUG than for the first AUG in both PpRPOT1 and PpRPOT2. This is the first positive computational data suggesting that the second AUG may be used preferentially.

To reexamine the targeting of PpPROT1 and PpRPOT2 by GFP, we prepared several new GFP-fusion constructs using both the natural 5′-upstream sequence and the translation leader sequence (TL) taken from the *RBCS3A* gene of pea (*Pisum sativum*; Fig. 3, A and B). In addition, in TP1rM48I-GFP and TP2rM36I-GFP, the second AUG codon was mutated, while the translation was forced to start from the first one. These constructs were introduced into moss protoplasts by polyethylene glycol-mediated transformation.

In the experiments with TP1-GFP, the construct having the entire 5′-UTR plus both Met codons (wild-type construct having the original upstream sequence), and constructs having only the second Met codon, TP1M2-GFP and TP1rM2-GFP, the fluorescence of GFP was localized to mitochondria (Fig. 4A, a, b, and d), whereas the fluorescence of GFP was localized to plastids with the construct having a mutated second Met codon, TP1rM48I-GFP (Fig. 4A, e). In analogous constructs with the PpRPOT2, namely, the construct having the entire 5′-UTR, TP2-GFP, and the constructs having the second Met codon, TP2M2-GFP and TP2rM2-GFP, the GFP fluorescence was localized to mitochondria (Fig. 4A, f, g, and i), while the fluorescence of GFP was localized to plastids with TP2rM36I-GFP (Fig. 4A, j). The fluorescence of GFP with the constructs having a translation leader that replaced the 5′-UTR, TP1rM1-GFP and TP2rM1-GFP, was localized to both mitochondria and plastids (Fig. 4A, c and h). These results indicate that forced

Figure 3. Schematic representation of DNA constructs. A, PpRPOT1-GFP fusion plasmids used in *P. patens* protoplast transformation. B, PpRPOT2-GFP fusion plasmids. C, Plasmids for the measurement of translation efficiency. D, Plasmids for Arabidopsis transformation. Open rectangles represent the 5′ part of *PpRPOT1*, *PpRPOT2*, or *AtRpoT2* cDNAs. Vertically hatched rectangles represent reporter part, encoding either GFP (in A, B, and D) or GUS (in C). Black rectangles represent TL from the pea *RBCS3A* gene. Maps are not drawn to scale for readability. In C, u1 and u2 are upstream sequences of the first and the second AUG codons, respectively. TP1 and TP2 in the construct names are used here to designate putative transit sequence (with or without upstream sequence) of PpRPOT1 and PpRPOT2, respectively. "r" indicates *RBCS* TL. M1 and M2 indicate the first and the second Met, respectively. Crosses over ATG indicate that the ATG was mutated to ATC (M48I and M36I mutations, respectively). "+1" indicates that the insert begins from the A of the first ATG, while "−10" indicates that the insert begins from the −10 position.
translation from the first AUG codon using the TL of pea RBCS 3A resulted in the localization to plastids, just as in the reported results with pOL S65C vector (Richter et al., 2002). However, TP1-GFP and TP2-GFP with the natural 5′-upstream sequence were localized only to mitochondria. These results are explained if the first AUG codon is not used as the translation initiation site in vivo, although the amino acid sequence beginning from it has a property of plastid targeting sequence.

No secondary structure such as stem-loop is predicted in the 5′-upstream sequence of the two PpRPOTs with the software RNAstructure version 3.71 (Mathews et al., 1999). However, some uORFs were detected in the 5′-UTR. We examined the influence of the 5′-upstream sequence on the translation efficiency with GFP-fusion constructs (Fig. 4B, TP1M1210-GFP; containing the 10 bp sequence upstream of the first AUG codon but no uORF, TP1M11-GFP; the 35S promoter directly joined to the first AUG). The results suggest that the GFP fluorescence of TP1M110-GFP was localized to mitochondria (Fig. 4B, a). The fluorescence of TP1M11-GFP was localized to both mitochondria and plastids (Fig. 4B, b). In the experiment with TP1M110-GFP, GFP fluorescence was observed in mitochondria as in the case of TP1-GFP containing the full-length 5′-upstream sequence of the first AUG codon. These results suggest that the uORF has no effect on the translation efficiency of the first AUG codon, and that only the 10-nucleotide sequence upstream of the first AUG is necessary to suppress translation from this site. However, this may not be suppression because not all AUGs within the mRNA act as initiation codons. What can be concluded from this experiment is that the proximal 10-nucleotide sequence but not the long 5′-UTR sequence upstream is important to determine whether translation is started from this site of RROT mRNA.

### Quantitative Estimation of the Effect of the 5′-Upstream Sequence on the Translation of PpRPOT1 and PpRPOT2

To quantitate the effects of the 5′-upstream sequence on the translation from the first and the second AUGs of PpRPOT1 and PpRPOT2, five additional plasmids with β-glucuronidase (GUS) reporter (Fig. 3C) were constructed and tested in transient expression. These constructs contain either the 5′-upstream sequence of the first (PpRPOT1u1-GUS and PpRPOT2u1-GUS) or the second AUG codon (PpRPOT1u2-GUS and PpRPOT2u2-GUS), or the TL from the pea RBCS 3A gene (TL-GUS), which were fused to the niaD gene and driven by the cauliflower mosaic virus (CaMV) 35S promoter. They were introduced into moss protoplasts by polyethylene glycol-mediated transformation. One
day after transformation, GUS activity was measured. Two constructs, PpRPOT1u2-GUS and TL-GUS, gave high levels of GUS activity, whereas the GUS activity with PpRPOT1u1-GUS was at a level of pUC18 control (Table II). In contrast with PpRPOT1 constructs, PpRPOT2u2-GUS gave lower GUS activity, but this activity was significantly higher than the activity with PpRPOT2u1-GUS and pUC18 control. Therefore, it seems that translation is initiated only at the second AUG codon in both PpRPOT1 and PpRPOT2. In other words, the first AUG codon is unlikely to be recognized as a translation initiation site.

### Subcellular Localization of PpRPOT1 and PpRPOT2 in Stably Transformed P. patens

In our transient expression experiments, both PpRPOT1 and PpRPOT2 were translated from the second AUG codon and were only localized to mitochondria in the protoplasts. However, the first AUG codon could be used as an initiation site in some particular types of cells or tissues. To address this question, we examined in detail the localization of PpRPOT1 and PpRPOT2 in various cell types and tissues in stably transformed moss (stable is used here to mean not transient or stably integrated in the chromosome). The pPpMADS-TP1GFP and pPpMADS-TP2GFP plasmids containing the natural 5′-upstream sequence of the first AUG codon and N-terminal sequence were used for the transformation. In all cell types and tissues of stably transformed P. patens, including protonemata and gametophores, GFP fluorescence was only observed in mitochondria but never in chloroplasts (Fig. 5). When immunoblot analysis was performed with plastids and mitochondria isolated from stably transformed P. patens protonemata, the GFP-fusion protein was detected in the mitochondrial fraction but not in the plastid fraction.

### Table I. Oligonucleotides used in this study

Underlines indicate restriction sites.

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Thus, subcellular localization of the GFP-fusion protein in stably transformed protonemata and other tissues was the same as that in transiently transformed protoplasts, confirming that no detectable level of translation occurs from the first AUG codon of PpRPOT1 and PpRPOT2. It was concluded that the N-terminal extension in each of the two PpRPOTs (Fig. 6A) beginning from the first AUG codon is not translated in vivo.

**Phylogenetic Analysis of the Subcellular Localization of AtRpoT;2**

Phylogenetic analysis of the RPOT proteins (Fig. 6A, left) suggested three major clusters, with a relationship (moss, ((Cluster I, Cluster II), Cluster III)). This result is essentially consistent with the structure of N-terminal sequences: Cluster I is characterized by the conserved sequence block (MWR) and mitochondrial localization. The conserved sequence block is not found in Cluster III, which contained plastid proteins. N-terminal extension as well as the conserved sequence block are found in the sequences in Cluster II (Fig. 6A). However, ZmRpoT1 belongs to Cluster II but lacks N-terminal extension. AtRpoT;2 and NsRpoT-B have been shown to be dually targeted to both plastids and mitochondria in experiments with GFP-fusion proteins (Hedtke et al., 2000; Kobayashi et al., 2001a). Immunological evidence suggested that Wheat-G is a mitochondrial RNA polymerase (Ikeda and Gray, 1999), but the plastid localization was not tested with GFP. The localization of Wheat-C was not reported (Fig. 6B). We chose AtRpoT;2 to test its targeting since the 5′-upstream sequence was not included in the upstream sequences in Cluster II (Fig. 6A). However, ZmRpoT1 belongs to Cluster II but lacks N-terminal extension. AtRpoT;2 and NsRpoT-B have been shown to be dually targeted to both plastids and mitochondria in experiments with GFP-fusion proteins (Hedtke et al., 2000; Kobayashi et al., 2001a). Immunological evidence suggested that Wheat-G is a mitochondrial RNA polymerase (Ikeda and Gray, 1999), but the plastid localization was not tested with GFP. The localization of Wheat-C was not reported (Fig. 6B). We chose AtRpoT;2 to test its targeting since the 5′-upstream sequence was not included in the upstream sequences in the previous report. The sequence context of two AUGs was compared to the consensus sequence of plants AAAACAAA(A/C)A-AUG. However, both of the upstream sequences were not analogous to the consensus sequence (Fig. 1B). However, a computer prediction of translation initiation site with the NetStart program (Pedersen and Nielsen, 1997), which is said to be specialized for Arabidopsis, suggested that AtRpoT;2 is not translated from the first AUG codon but is translated from the second one (Fig. 6B). Analogous prediction data are presented for other plants, but they might not be correctly predicted because of limitation of neural network prediction using the Arabidopsis training data set. Two plasmids were constructed (Fig. 3D). pBI-AtRpoT;2GFP contained the 5′-upstream sequence of the first AUG codon and coding sequence (126 amino acids), and pBI-AtRpoT;2+1GFP contained the coding region and no native 5′-upstream sequence. These constructs were used to transform Arabidopsis (Fig. 7), and then subcellular localization of the GFP-fusion proteins was observed in cotyledon, leaf, and root. In the experiment with pBI-AtRpoT;2GFP, the fluorescence of GFP was localized to mitochondria (Fig. 7A, a, d, and g). By contrast, the fluorescence of AtRpoT;2+1GFP was observed in both mitochondria and plastids (Fig. 7B, j, m, and p). Additionally, we investigated into the localization of AtRpoT;2 during the early stage of seedling development and during the deetiolation process because Baba et al. (2004) reported that the mutation of (data not shown). Thus, subcellular localization of the GFP-fusion protein in stably transformed protonemata and other tissues was the same as that in transiently transformed protoplasts, confirming that no detectable level of translation occurs from the first AUG codon of PpRPOT1 and PpRPOT2. It was concluded that the N-terminal extension in each of the two PpRPOTs (Fig. 6A) beginning from the first AUG codon is not translated in vivo.

**Table II. Effects of the 5′-upstream sequences on translation efficiency**

GUS activity in P. patens protoplasts that were transformed with respective plasmid is normalized with the coexpressed GFP. The GUS values are expressed in arbitrary unit, with setting the average values of expression driven by the TL-GUS as 100. Each value (±SE) represents the average of three independent assays. N/A: Not applicable.

<table>
<thead>
<tr>
<th>Construct</th>
<th>Relative GUS Activity</th>
<th>P Value</th>
</tr>
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<tbody>
<tr>
<td><strong>Experiment 1</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PpRPOT1u1-GUS</td>
<td>0.85 ± 0.34</td>
<td>0.51</td>
</tr>
<tr>
<td>PpRPOT1u2-GUS</td>
<td>93.03 ± 21.85</td>
<td>0.02</td>
</tr>
<tr>
<td>TL-GUS</td>
<td>100 ± 24.77</td>
<td>0.02</td>
</tr>
<tr>
<td>pUC18</td>
<td>0.67 ± 0.21</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Experiment 2</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PpRPOT2u1-GUS</td>
<td>1.10 ± 0.04</td>
<td>0.76</td>
</tr>
<tr>
<td>PpRPOT2u2-GUS</td>
<td>12.40 ± 0.44</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>TL-GUS</td>
<td>100 ± 9.75</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>pUC18</td>
<td>1.18 ± 0.09</td>
<td>N/A</td>
</tr>
</tbody>
</table>

**Figure 5.** Localization of PpRPOT-GFP fusion proteins in stably transformed P. patens. Localization of PpRPOT1-GFP fusion protein in a part of a gametophore (a–c) and a protonema (d–f), and of PpRPOT2-GFP fusion protein in a part of a gametophore (g–i) and a protonema (j–l). a, d, g, and j, Fluorescence of GFP; b, e, h, and k, fluorescence of chlorophyll; c, f, i, and l, Nomarski differential interference image. Fluorescence of GFP (green) and chlorophyll (red) was observed using a fluorescence microscope BX-60 (Olympus) equipped with cubes U-MNIBA and U-MWU, respectively. Bar = 5 μm.
Figure 6. Comparison of various plastid and mitochondrial RPOTs in plants. A, Phylogenetic analysis and alignment of the N-terminal sequences of RPOT homologs. The following sequences were obtained from the GenBank database (in parentheses): PpRPOT1 and PpRPOT2 (P. patens; AB055214 and AB055215); OsRpoT2 (Oryza sativa; AB096015); AtRpoT;3 (Arabidopsis; Y08463); NsRpoT-C (N. sylvestris; AJ302020); ZmRpoT2 (Zea mays; AF127022); Wheat-C (Chenopodium album; Y08067); ZmRpoT1 (Z. mays; AF127021); OsRpoT2 (O. sativa; AB096014); Wheat-G (T. aestivum; AF091838); AtRpoT1;2 (Arabidopsis; AJ001037); and NsRpoT-B (N. sylvestris; AJ302019). Left, Phylogenetic tree constructed by the neighbor-joining method. Amino acid sequences were used in this analysis (Kabeya et al., 2002; ALIGN_000281 in the EMBL-Align database). The numbers on the branches show bootstrap confidence levels obtained with 1,000 bootstraps. Right, Alignment of the N-terminal sequences of RPOT homologs. Asterisks indicate the conserved sequence block. B, Summary of localization of various RPOTs. If there are two putative initiation codons, (1) indicates the polypeptide translated from the first Met and (2) indicates the polypeptide translated from the second one. Results of computer prediction on the targeting by TargetP and on the probability of initiation site by NetStart are listed along with experimental results. Abbreviations for the localizations: Pt and Mt indicate plastids and mitochondria, respectively. (Pt ?), Potentially targeted to plastids but no evidence for translation of such polypeptide. The following numbers are used to show experimental evidence: 1, GFP fusion; 2, in vitro import; and 3, immunoblot. ND, No available data. References cited in the figure image are as follows: Hedtke et al. (1997, 1999, 2000); Chang et al. (1999); Ikeda and Gray (1999); Kobayashi et al. (2001a, 2001b, 2002); Kabeya et al. (2002); Richter et al. (2002); Kusumi et al. (2004); and this study.


**Figure 7.** Localization of AtRpoT2-GFP fusion proteins in stably transformed Arabidopsis. A, Arabidopsis was stably transformed with pBI-AtRpoT2-GFP. Localization of AtRpoT2-GFP fusion protein in cotyledon (a–c), leaf (d–f), and root (g–i). B, Arabidopsis was stably transformed with pBI-AtRpoT2-1+1GFP. Localization of AtRpoT2-GFP fusion protein in cotyledon (j–l), leaf (m–o), and root (p–r). In the root, no fluorescence of chlorophyll was observed. Fluorescence of MitoTracker is shown to locate mitochondria, although cell wall was also densely stained. a, d, g, j, m, and p. Fluorescence of GFP; b, e, k, and n, fluorescence of GFP and chlorophyll; h and q, fluorescence of MitoTracker; c, i, l, o, and r, Nomarski differential interference image. Fluorescence of GFP, chlorophyll, and MitoTracker was observed using a fluorescence microscope BX-60 (Olympus) equipped with cube U-MNIBA, U-MWU, and U-MWIG, respectively. Bar = 10 μm.

AtRpoT2 affected the light-induced accumulation of several plastid gene transcripts during early seedling development. However, the fluorescence of AtRpoT2-GFP was not detected in plastids under any conditions tested (data not shown). These results are essentially equivalent to those of the in vivo targeting experiments of PpRPOTs described above, and suggest that AtRpoT2 may also contain a formal (or, more precisely, unused) plastid targeting sequence in its N terminus, as in the case of the two PpRPOTs.

**DISCUSSION**

In this study, we showed that the PpRPOT1 and PpRPOT2 proteins in the moss *P. patens* are immunologically detected in the mitochondrial fraction but not in the plastid fraction, and that the transcription activity in the plastids is nearly completely inhibited by tagetitoxin, an inhibitor of PEP. We then confirmed the mitochondrial localization of PpRPOTs-GFP fusion proteins in transiently and stably transformed *P. patens*. We further investigated the translation efficiency of the 5′-upstream sequences of the first or the second AUG codon with GUS fusions. Our data indicated that PpRPOTs are translated from the second AUG codon, and such protein is targeted only to mitochondria in vivo, although the proteins are capable of targeting to plastids when translation is forced to start from the first AUG. The exclusive mitochondrial localization was confirmed in various tissues of stably transformed *P. patens*. Therefore, all available evidence indicates that the two PpRPOTs are targeted to mitochondria but not to plastids. Accordingly, a nuclear-encoded plastid RNA polymerase similar to mitochondrial RPT in flowering plants, called NEP, does not exist in the plastids in *P. patens*. Plastid-encoded enzyme is likely the only RNA polymerase in *P. patens* plastids. Although we examined various cells and tissues of the transgenic moss expressing PpRPOT1-GFP and PpRPOT2-GFP, we did not detect a cell in which GFP fluorescence is localized to the plastids. The reproductive organs were not examined because they arose sporadically and rarely. They are to be examined in the future. Nevertheless, as far as the dual targeting of PpRPOTs, as originally proposed in the protonemal cells, is concerned, we can clearly say that PpRPOTs are not present in the chloroplasts of protonemal cells. Unfortunately, we have no clear answer to the question of why two AUG codons are present in these genes and why the first AUG is not used in vivo. The sequence upstream of the first AUG might not inhibit translation initiation, but simply the AUG is inactive as the initiation codon as all other AUGs within the transcript. An evolutionary view on this point is described below.

We should emphasize that the effects of the 5′-upstream sequence or translational context should be considered in experiments to examine the localization of polypeptide using GFP-fusion proteins. In a number of targeting experiments with GFP-fusion proteins, translation was forced to start from the first AUG codon without the 5′-upstream sequence. The presence or absence of the 5′-upstream sequence strongly influences the translation efficiency. In the case of PpRPOTs, the 5′-upstream sequence of the first AUG does not promote translation initiation at this site. It is generally difficult to predict the selection of the translation initiation site merely on the basis of the nucleotide sequence. Different consensus sequences are known in different organisms, e.g. UAAAUGANAU in protozoa (*Yamauchi*, 1991), C(A/G)CCAUAG in vertebrates (*Kozak*, 1987), (A/C)(A/G)(A/C)-CAUGGC in monocots, and AA(A/C)AAUGGC in dicots (*Joshi* et al., 1997). However, some software that predicts functional initiation codon was developed, such as NetStart (*Pedersen* and *Nielsen*, 1997), for specific species of plants. The prediction for AtRpoT2 is in agreement with our experimental data. In addition, we developed a matrix based on information content calculated for available *P. patens* data obtained.

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**Kabeya and Sato**
from GenBank entries. This approach seemed partly successful in that the score for the second AUG was higher than the score for the first AUG in both PpRPOT1 and PpRPOT2. However, the score for the first AUG of PpRPOT2 is higher than the score for the second AUG of PpRPOT1. This suggests that this method should be refined with more data. In vitro translation experiments often have been used to examine the translation initiation site, but this also is difficult since Lütcke et al. (1987) reported that the selection of translation initiation codons differs in wheat germ and reticulocyte. Thus, a heterologous system does not provide conclusive evidence. In fact, Richter et al. (2002) observed products that were translated from the first AUG and the second AUG in their in vitro translation experiments using reticulocytes, whereas we showed that the translation initiation from the first AUG is negligible in vivo. After all, in vivo experiments with constructs containing the native 5'–upstream sequences appear to be essential for identifying the translation initiation site and subcellular localization, although other experiments that have been done conventionally are also valuable.

Localization of AtRpoT2 is an important issue in analyzing the transcription in both mitochondria and plastids in Arabidopsis. This protein possesses the N-terminal extension, as do PpRPOTs, and has so far been regarded as a dually targeted protein, while AtRpoT1 and AtRpoT3 are targeted to mitochondria and plastids, respectively (Hedtke et al., 2000). However, our results with GFP-fusion proteins suggested that the AtRpoT2 transcript is translated from the second AUG but not from the first AUG in the natural context and its product uniquely localized to mitochondria in many Arabidopsis tissues (Fig. 7). Tagetitoxin test cannot be applied in Arabidopsis because the isoforms 1 and 3 are localized in mitochondria and plastids, respectively, and the effect of additional localization of isoform 2 is difficult to assess. Our results lack critical data on immunological analyses of the three RpoT isoforms in Arabidopsis organelles; however, the present situation necessarily raises questions about the regulation of translation and localization of AtRpoT2.

Recently, Baba et al. (2004) reported that both plastid and mitochondrial transcription was affected in an AtRpoT2 mutant, which was isolated from a population of activation-tagged T-DNA insertion lines. This mutant exhibited short roots, reduced hypocotyl length, delay in greening, and defect in light-induced accumulation of several plastid mRNAs as well as atp1 of mitochondrial mRNA. Their finding seemed consistent with the traditional idea that AtRpoT2 is localized to both plastids and mitochondria. They analyzed organellar gene expression mostly in leaves, and, therefore, we still need some intricate explanation that compromises our data (AtRpoT2 is localized to mitochondria in leaves, stems, and roots) and the data of the overexpressing line. It is quite probable that AtRpoT2 in the activation line is involved in plastid transcription in some way. The AtRpoT2 protein could be imported to plastids as well due to side effect of high level expression. Another possibility might be an indirect effect resulting from complex mitochondrion-chloroplast interactions. A mutant in the mitochondrial genome is known to cause variegation (Sakamoto et al., 1996), although no detailed mechanism is known. However, a more probable explanation for the solution of these apparent discrepancies might be that AtRpoT2 targets to plastids as well in developing leaf cells or leaf primordia. There, this enzyme could trigger a cascade of reactions leading to normal gene expression in mature leaves. The overexpression of AtRpoT2 could change the development of leaf cells and affects the level of gene expression in chloroplasts in mature leaves, although this enzyme is not localized in chloroplasts in mature leaves. This possibility may be solved by a strategy that develops a system that can sensitively detect changes in targeting of a given protein (see below).

In a previous study, we proposed that the creation of the NEP occurred in angiosperms after their separation from gymnosperms (Kabeya et al., 2002). The creation of the NEP in angiosperms occurred by gene duplication. The results of this study suggest that the plastid targeting sequence of the NEP might have been acquired before this gene duplication, even though it is not really used. There are two possible hypotheses on the origin of the angiosperm NEP. In one hypothesis, the plastid targeting sequence was present already before the separation of vascular plants and mosses but had remained nonfunctional (or formal) until the plastid targeting sequence was really used by activating the translation from the first AUG in angiosperms. In angiosperm NEPs such as AtRpoT3, the second Met has been changed and no longer acts as an initiation site. The mitochondrial RPOTs, such as AtRpoT1, lost the N-terminal extension either by mutation of the first AUG or deletion of entire extension. In another hypothesis, a formal plastid targeting sequence was added independently at various stages of evolution. In this case, the formal plastid targeting sequences of the moss are an example of an unsuccessful attempt to create a plastid protein. The NEP is a successful example, while the dually targeted RPOTs such as AtRpoT2, are an example of ongoing evolution by the addition of plastid targeting sequence.

A biologist’s intuition favors that retention of the N-terminal extension that could serve as a plastid targeting sequence in many RpoT proteins is meaningful. This is an opinion shared by many of our colleagues. There are several solutions to this philosophical question. (1) The presence of the N-terminal extension is found in many RpoT proteins, but the sequence and length are not highly conserved. There is no selection pressure in this respect. In particular, the extension sequences in the moss are very different from those in flowering plants. (2) We do not have enough data on the presence of such N-terminal extension in plant proteins or eukaryotic proteins in general. If the presence of N-terminal extension is specific to RpoT
proteins, then we will have to consider a specific role of the N-terminal extension. However, the exact N terminus of most proteins has not been determined experimentally, even though we sometimes encounter two or three Met residues in the N-terminal segment of an ORF, which is the longest reading frame that can be estimated for a given genomic sequence. (3) We will be able to challenge this hypothesis based on biologists’ intuition by experimental approach. If we can detect sensitively a change in intracellular localization of a protein, we will be able to answer such a question. A possible method is to use an enzyme that functions normally in mitochondria but that causes serious damage when targeted to plastids. Various versions of such a system can be imagined, and we should try to demonstrate whether targeting to plastids of AtRpoT2 occurs in some special types of cells during the development of plant.

In conclusion, available data suggest that the two PpRPOs are targeted only to mitochondria due to exclusive translation from the second AUG codon and that the same is apparently true for AtRpoT2.

MATERIALS AND METHODS

Plant Material

Grenzlebn strain of Physcomitrella patens (Hedw.) Bruch & Schimp subsp. patens Tan was grown in the minimal medium supplemented with 5 mM diammonium (+)-tartrate as described previously (Hashimoto and Sato, 2001). Agar (0.8%) plates were used for maintaining the stock culture at 25°C. Light was provided by a bank of fluorescent lamps at a fluence rate of about 35 μmol m⁻² s⁻¹.

Isolation of Plastids

Seven-day-old protonemal cells were digested in Solution 1 (2.0% Driserase [Kyowa Hakko, Tokyo] and 8% mannitol) at 25°C for 30 min. Protoplasts were recovered by filtration through a 70-μm nylon mesh. The suspension of broken protoplasts was centrifuged at 10,000 g for 10 min at 4°C. The pellet was suspened in Grinding buffer.

Isolation of Mitochondria

Mitochondria were isolated from the broken protoplasts (see above). The supernatant after the centrifugation at 10,000 g for 10 min was then centrifuged at 18,000 g for 10 min at 4°C. The pellet was suspended in Grinding buffer.

Percoll was added to the suspension to a final concentration of 20%. Mitochondria were purified by Percoll density gradient centrifugation (20%/33%/80%, v/v) at 18,000g for 60 min at 4°C. A yellowish turbid band that formed at the 33%/80% interface was collected, washed three times with the Grinding buffer at 18,000g for 10 min at 4°C, and then diluted three times with TAN buffer. The mitochondria were suspended in a small volume of TAN buffer containing 33% glycerol, flash frozen in liquid nitrogen, and stored at −80°C. Purity of the organelles was checked by examination under fluorescence microscope as described above.

Effects of Tagetitoxin on Organellar Transcription

Transcription activity of plastids and mitochondria was measured as incorporation of [3H]UTP. In organellar assay, a 60-μL reaction contained transcription buffer [40 mM Tris-HCl, pH 8.3, 25 mM MgCl₂, 9 mM MgSO₄, 30 mM (NH₄)₂SO₄, 5 mM dithiothreitol, 0.01% Nonidet P-40, 180 μM ATP, 180 μM GTP, 180 μM CTP, 5 μM [3H]UTP (at a specific radioactivity 0.16 GBq μmol⁻¹ for plastids and 0.51 GBq μmol⁻¹ for mitochondria), 5.7 unit μL⁻¹ RNAguard (Amersham Bioscience, Piscataway, NJ), and 60 μg of protein plastids or 24 μg of protein mitochondria. Transcription activity of recombinant PpRPOs was measured as incorporation of [3H]UTP. For in vitro assay of purified enzymes, a 60-μL reaction contained transcription buffer (see above), 5 mM dithiothreitol, 0.01% Nonidet P-40, 180 μM ATP, 180 μM GTP, 180 μM CTP, 5 μM [3H]UTP (at a specific radioactivity 0.16 GBq μmol⁻¹), 5.7 unit μL⁻¹ RNAguard, 3 μg of calf thymus DNA as the template, and 1.5 μg of recombinant PpRPO or PpRPO2. The reaction mixtures were incubated for 30 min at 25°C. Tagetitoxin (Epigentec, Madison, WI) was added to a final concentration of 10 μM in the inhibition experiments. After the reaction, 5-μL aliquots were spotted onto DEAE paper (DE-81; Whatman, Clifton, NJ). After successive washing with 5% Na₂HP0₄, water, and ethanol, radioactivity was determined by liquid scintillation counting.

Antibody Preparation

The DNA fragments corresponding to the amino acids 121 to 500 of PpRPO1 and 114 to 499 of PpRPO2 were amplified from pZL-1 and pZL-2 using the primers 1 and 2, or 3 and 4 (Table I). These PCR products were digested with SfiI and BamHI and BglII, and inserted into BamHI and SfiI sites of the expression vector pGEX-4T-2 (Amersham Bioscience), respectively. The resulting plasmids were named pGEX-1p and pGEX-2p, and transformed into Escherichia coli XL-1 Blue cells. The overexpression and purification with glutathione-Sepharose 4B (Amersham Bioscience) of the GST-PpRPO1 and GST-PpRPO2 fusion proteins were performed according to the manufacturer’s directions. The fusion protein eluted from the column was further purified by gel filtration with the Superdex 75 column (Amersham Bioscience) that had been equilibrated with PBS buffer. Purified proteins were used to immunize guinea pigs. Polyclonal antisera were obtained, and the IgG fraction (after ammonium sulfate fractionation) was used in the immunoblot analysis. For immunoblot analysis, intact plastids and mitochondria were isolated from P. patens as described above. SDS-PAGE and immunoblotting were performed using a 7.5% or 20% polyacrylamide gel as described in a previous paper (Sato et al., 1998).

GFP-Fusion Plasmids

Either of the plasmids pZL-1 or pZL-2 that contained cloned cDNA encoding PpRPO1 and PpRPO2 (Kabayda et al., 2002), respectively, were used as templates for PCR amplification. TP1-GFP and TP2-GFP containing the 5’-upstream sequence of the first AUG codon were amplified, respectively, using primers 5 and 32, or 13 and 32. TP1M2-GFP and TP2M2-GFP containing the 5’-upstream sequence of the second AUG codon (but downstream the first AUG codon) were amplified using primers 5 and 32, or 13 and 16. TP1M1-GFP and TP2M1-GFP, having TL in place of the 5’-upstream sequence of the second AUG codon, were generated using primers 5 and 32, or 13 and 16. TP1M1-GFP and TP2M1-GFP, having TL from the pea (Pisum sativum) RBCS 3A gene in place of the 5’-upstream sequence of the first AUG codon, were generated using primers 5 and 7, or 13 and 15. TP1M2-GFP and TP2M2-GFP, having TL in place of the 5’-upstream sequence of the second AUG codon, were generated using either primers 7 and 10, or primers 5 and 9. TP2M36-GFP, containing
TL and Met-36-to-Ile mutation, was generated likewise by two successive PCR reactions, using primers 13, 15, 17, and 18. TP1 +1-GFP, containing no 5′-upstream sequence of the first AUG codon, was amplified using primers 5 and 11. TP1 –10-GFP, containing 10-nucleotide sequence in the 5′-upstream region of the first AUG codon, was amplified using primers 5 and 12. These PCR products were digested with Ncol and Sall and inserted into the Sall-Ncol sites of sGFP56ST (Chiu et al., 1996), which contains a synthetic GFP gene with S6ST mutation and optimized codon usage for plants under the control of the CaMV 35S promoter. The plasmids pPPmADS-TP1GFP and pPPmADS-TP2GFP were constructed by inserting the 5′-upstream sequence of the first AUG codon plus the full-length transit peptide of PpRPOT1 or PpRPOT2 and a DNA fragment containing the GFP coding sequence into an expression vector pPPmADS2-7133 with E7133 promoter (Mitsuhashi et al., 1996).

The DNA fragments related to AtRpoT2 were amplified from Arabidopsis (cv Columbia) genomic DNA (laboratory stock). To construct AtRpoT2-GFP, the 5′-upstream sequence of the first AUG codon plus the full-length transit peptide sequence was amplified using primers 19 and 21. To construct AtRpoT2–1+GFP, the sequence coding for the full-length transit peptide was amplified using primers 20 and 21. These PCR products were digested with Ncol and Sall and inserted into the Sall-Ncol sites of sGFP56ST. These constructs were used in transient expression experiments (data not shown). The DNA fragments beginning from the CaMV 35S promoter and ending at the NOS terminator were obtained from the plasmids AtRpoT2-GFP or AtRpoT2–1+GFP, and were inserted into the HindIII-EcoRI sites of pBlI01. These plasmids were named pBlI-AtRpoT2-GFP and pBlI-AtRpoT2–1+GFP, respectively.

**Transient Expression of GFP-Fusion Constructs in the Moss**

Thirty micrograms of the GFP-fusion plasmids were introduced into the protoplasts of *P. patens* by polyethylene glycol-mediated transformation (Schafer, 1994; Nishiyama et al., 2000).

**GUS-Fusion Plasmids and Measurement of Translation Activity with GUS**

To generate the DNA fragment PpRPOTu1u1-GUS and PpRPOTu2u1-GUS containing the 5′-upstream sequence of the first AUG codon, the fragment containing the 5′-upstream sequence of the first AUG codon was amplified with primers 23 and 32 using the pZL-2 plasmid as a template, or with primers 27 and 32 using the pZL-2 plasmid as a template. The fragment containing suDA gene was amplified with primers 22 and 31, or 26 and 31, using the pBlI01 plasmid as a template. Then the two fragments in each combination were combined by a second PCR with primers 31 and 32. To generate the DNA fragment PpRPOTu2u2-GUS and PpRPOTu2u2-GUS containing the 5′-upstream sequence of the second AUG codon (but downstream the first AUG codon), the fragment containing the 5′-upstream sequence of the second AUG codon was amplified with primers 6 and 25 using the pZL-1 as a template or with primers 14 and 29 using the pZL-2 as a template, while the fragment containing the suDA gene was amplified with primers 24 and 31, or 28 and 31, using the pBlI01 plasmid as a template. Then, the two fragments in each combination were connected by amplification with primers 6 and 31 or primers 14 and 31. The DNA fragment TL-GUS containing the TL from the pea RBCS 3A gene and the GUS coding sequence was amplified with the primers 30 and 31 using the pBlI01 plasmid as a template. These PCR products were digested with Sall and Nof and inserted in place of the GFP coding sequence of sGFP56ST, keeping the CaMV 35S promoter and the NOS terminator unchanged.

One of these plasmids (15 μg) and sGFP56ST as an internal standard were introduced into the moss protoplasts as described above. One day after transformation, GUS activity of the protoplasts was determined according to Jefferson et al. (1987), while the amount of GFP protein was determined by immunoblot with anti-GFP antibody (Innogenetix, Carlsbad, CA).

A two-sample *t* test comparing GUS activity of pUC18 and various GUS constructs was used to calculate *P* value. A *P* value of less than 0.05 was considered significant.

**Stable Transformation**

*P. patens* was transformed according to Schafer (1994). To obtain stable transformants, pPPmADS-TP1GFP digested with Nof1 was introduced into the protoplasts. Transformed protoplasts were incubated for 4 d on BCDAT medium and then transferred to BCDAT medium containing 50 mg L−1 G418 (Sigma, St. Louis) for 3 weeks. The selected plants were transferred onto a medium without G418 and allowed to grow for 7 d. Then, they were transferred again onto the selection medium. After the second selection, stably transformed *P. patens* was confirmed by PCR analysis.

Arabidopsis was transformed according to Bechtold et al. (1993) using Agrobacterium tumefaciens strain EHA105 containing pBlI-AtRpoT2-GFP or pBlI-AtRpoT2–1+GFP. Transformed Arabidopsis were selected on the Murashige and Skoog medium (Murashige and Skoog, 1962) containing 1.5% Suc, 50 mg L−1 kanamycin, and 100 mg L−1 carbenicillin.

**Computational and Phylogenetic Analysis**

Database sequences and alignment files were used the SIEVE package version 1.30 (Sato, 2000). A total of 266 database entries for moss sequences were retrieved directly from the glnprn seq files in the GenBank (version 141) release using the getent command. The UTR sequence (20 bases) was extracted for each entry by the cdsnuc command. We finally used 214 UTR sequences for further calculation. SequenceLogo (Schneider and Stephens, 1990) was prepared using the alpro and makelogo programs, which were downloaded from Tom Schneider’s web site (http://www.cebcb.nl/seq/seqlogo/) and compiled locally for Power PC G5 running under MacOS X 10.3. The information content values in the resulting logo file in ASCII postscript format were used as a scoring matrix (the values were in fact multiplied by 2.5 for drawing as SequenceLogo with a height of 5.0 cm). The score was calculated by adding the value for the corresponding nucleotide at each position from −10 to −1 with respect to A of initiation codon.

For phylogenetic analysis, alignments of amino acid sequences were constructed by ClustalX program version 1.81 (Thompson et al., 1994) with final manual adjustment. The N-terminal part was excluded from the alignment by the getclu command of SIEVE because it was highly variable.

The programs used for constructing phylogenetic trees by the neighboring-joining method were PROTDIST, NEIGHBOR, SEQBOOT, and CONSENSE of the PHYLIP package (Felsenstein, 1988). Graphical representation of phylogenetic tree was made by the njplot program (Perriere and Gouy, 1996).

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