Alternative mRNA splicing of 3′-terminal exons generates ascorbate peroxidase isoenzymes in spinach (Spinacia oleracea) chloroplasts

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We have isolated two cDNA clones encoding spinach (Spinacia oleracea) stromal and thylakoid-bound ascorbate peroxidase isoenzymes [Ishikawa, Sakai, Yoshimura, Takeda and Shigeoka (1996) FEBS Lett. 384, 289–293]. The gene (ApxII) encoding both chloroplastic ascorbate peroxidase isoenzymes was isolated and the organization of the gene was determined. Alignment between the cDNAs and the gene for chloroplastic ascorbate peroxidase isoenzymes indicates that both enzymes arise from a common pre-mRNA by alternative splicing of two 3′-terminal exons. Genomic Southern-blot analysis supported this finding. The gene spanned nearly 8.5 kbp and contained 13 exons split by 12 introns. The penultimate exon 12 (residues 7376–7530) for the stromal ascorbate peroxidase mRNA consisted of one codon for Asp586 before the TAA termination codon, and the entire 3′-untranslated region, including a potential polyadenylation signal (AATAAA). The final exon 13 (residues 7545–7756) for the thylakoid-bound ascorbate peroxidase mRNA consisted of the corresponding coding sequence of the hydrophobic C-terminal region, the TGA termination codon and the entire 3′-untranslated region, including a potential polyadenylation signal (AATATA). Both exons were interrupted by a 14 bp non-coding sequence. Northern-blot and reverse transcription-PCR analysis showed that the transcripts for stromal and thylakoid-bound ascorbate peroxidase are present in spinach leaves.

INTRODUCTION

In plant tissues the chloroplasts are potentially the most powerful source of oxidants and sites within the cell most at risk from photo-oxidative damage. The photoreduction of molecular oxygen in chloroplasts is unavoidable and leads to the production of a superoxide radical (O2−), which is followed by formation of H2O2 via the disproportionation of O2− in enzymic and non-enzymic reactions. The production rate of O2− and H2O2 is estimated to be about 160 and 80 μM·s−1 in chloroplasts under normal conditions [1]. Because 10 μM H2O2 is sufficient to inhibit the photosynthetic assimilation of CO2 by 50% [2], photosynthesis is significantly impaired within a fraction of a second if H2O2 is not promptly scavenged. To rid themselves of excess H2O2, therefore, chloroplasts of higher plants develop two ascorbate peroxidase (AsAP) isoenzymes which exist as stromal soluble (sAsAP) and thylakoid-bound (tAsAP) forms [3–5]. Moreover, other AsAP isoenzymes have been known as a soluble (sAsAP) and thylakoid-bound (tAsAP) forms [3–5].

Miyake et al. [5,11] have reported that the enzymological properties of tAsAP are very similar to those of sAsAP with respect to high specificity for ascorbate (AsA), Km values for H2O2 and AsA, inhibition by cyanide, thiol-modifying reagents, thiols and suicide inhibitors, such as hydroxyurea, and inactivation in AsA-depleted medium. The sole difference in terms of properties between tAsAP and sAsAP is the higher molecular mass of the membrane-bound enzyme compared with the soluble enzyme. Thus tAsAP seems to be bound to thylakoid membranes in such a form that the active site of the enzyme is exposed to the stroma for the access of the substrate. In the previous study, the first complete cloning and molecular characterizations of sAsAP and tAsAP from spinach have shown that the nucleotide sequence encoding the tAsAP isoenzyme is identical with that of sAsAP through the coding region up to amino acid position 364, where the remainder of the C-terminal coding region is substituted by a different sequence that encodes 50 amino acids which constitutes a hydrophobic thylakoid-membrane-binding domain [12]. These data and the presence of different polyadenylation tracts in cDNAs encoding the sAsAP and tAsAP variants suggest that both enzymes are generated by a common pre-mRNA from an identical gene by alternative splicing of the 3′-terminal exons. The cDNAs encoding cAsAP isoenzymes have been isolated and characterized from several plant sources, including spinach (Spinacia oleracea) [7,13,14]. The genes from pea (Pisum sativum) [15] and Arabidopsis thaliana (thale cress) [16] have been isolated and characterized. Accordingly, it seems likely that the AsAP of spinach is a multigene family and there are at least four AsAP genes. In order to elucidate the occurrence of an identical nuclear gene by alternative splicing of the 3′-terminal exons and to understand the gene regulation of chloroplastic AsAP isoenzymes in higher plants, we sought to clone and characterize the nuclear gene (ApxII) from spinach.

MATERIALS AND METHODS

Materials

Spinach seeds were germinated on moist gauze at 15 °C in the dark. The cotyledons from seedlings grown for 10–14 days in the dark were transferred to illumination (140 μE/s per m²) for 24 h.

Abbreviations used: O2−, superoxide radical; AsAP, ascorbate peroxidase; sAsAP, stromal ascorbate peroxidase; tAsAP, thylakoid-bound ascorbate peroxidase; cAsAP, cytosolic ascorbate peroxidase; mAsAP, microbody-bound ascorbate peroxidase; AsA, ascorbate; poly(A)+, polyadenylated; SSC, standard saline/citrate.

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The nucleotide sequence data reported in this paper have been submitted to the DDBJ, EMBL and GenBank Nucleotide Sequence Databases under the accession number AB002467.
to obtain the greening cotyledons as previously described [12]. Restriction enzymes and modifying enzymes were purchased from Takara Shuzo Co., Ltd. (Kyoto, Japan). The cDNAs coding for spinach chloroplastic AsAP isoenzymes were originally cloned into plasmid pBluescript SK(+) [7,12]. The Escherichia coli strain Y1090r and DH5α F' were obtained from Amersham (Amersham, Bucks., U.K.). All other chemicals were of analytical grade and were used without further purification.

Genomic DNA extraction
Genomic DNA was isolated from mature green spinach leaves. The leaves (20 g) were frozen in liquid nitrogen and ground to a slurry using a pestle and mortar while adding 20 ml of extraction buffer [50 mM Tris/HCl (pH 7.8)/100 mM NaCl/10 mM EDTA/200 µg/ml proteinase K/2 % SDS]. The slurry was incubated at 37 °C for 1 h. The solution was phenol-extracted three times, and the DNA was collected by precipitation with 0.1 vol. of 3 M ammonium acetate and 2 vol. of 100 % ethanol. The DNA was further purified by ethidium bromide/CsCl density-gradient centrifugation at 100000 g for 16 h at 20 °C.

Genomic Southern hybridization
Total DNA (20 µg) from spinach leaves was digested to completion with various restriction enzymes, separated by agarosegel electrophoresis (1 % gels) and transferred to a Hybond N + filter (Amersham) using a model 785 vacuum blotter according to the manufacturer's (Bio-Rad, Richmond, CA, U.S.A.) instructions. After transfer of the DNA to a nylon membrane, the genomic DNA fragments for chloroplastic AsAP isoenzymes were detected by probing with radiolabelled full-length tAsAP cDNA using the a random-primed DNA labelling kit (Takara, Shuzo). Washing the membrane in 1 x standard saline citrate (SSC)/0.1 % SDS at 65 °C was referred to as 'low stringency' and 0.1 x SSC/0.1 % SDS at 65 °C as 'high stringency'.

Construction and screening of partial genomic library
Spinach genomic DNA (40 µg) was digested with restriction endonuclease EcoRI and fractionated by electrophoresis in 1 % agarose gel. The 3.0-6.0 kbp fragments, which were identified by probing with radiolabelled full-length tAsAP cDNA using the a random-primed DNA labelling kit (Takara, Shuzo), were obtained from the genomic DNA fragments for chloroplastic AsAP isoenzymes by alternative splicing [12]. The genomic organization of the gene for chloroplastic AsAP in spinach leaves was examined by Southern-blot analysis. When the genomic DNA was digested with several restriction endonucleases and the fragments were probed with tAsAP cDNA, one to three hybridization signals were detected, depending on the restriction enzymes (results not shown). When the membrane was washed under low-stringency conditions, the result showed the same hybrid pattern as that shown). When the membrane was washed under low-stringency conditions, the result showed the same hybrid pattern as that shown.)

RESULTS

Genomic Southern-blot analysis
The finding that the cDNAs encoding chloroplastic AsAP isoenzymes either contained or lacked a sequence with a part of the intact reading frame strongly suggests the possibility that only one gene generates pre-mRNA for chloroplastic AsAP isoenzymes by alternative splicing [12]. The genomic organization of the gene for chloroplastic AsAP in spinach leaves was examined by Southern-blot analysis. When the genomic DNA was digested with several restriction endonucleases and the fragments were probed with tAsAP cDNA, one to three hybridization signals were detected, depending on the restriction enzymes (results not shown). When the membrane was washed under high-stringency conditions, the result showed the same hybrid pattern as that under high-stringency conditions (results not shown). These observations suggest that there was one copy of the gene for chloroplastic AsAP in the genome of spinach. To determine whether this hypothesis was compatible with the structure of the gene for chloroplastic AsAP, the genomic DNA fragments
Figure 1 Organization of the spinach chloroplastic AsAP gene (ApxII)

Horizontal thin lines indicate sequentially overlapping clones derived from the spinach partial genomic library. The shaded box indicates the proposed region of spinach genomic DNA. Important restriction-enzyme sites used for mapping and subcloning are indicated. The locations of primers to detect the vicinity of connection of each clone are shown by arrows. Exons are represented by open (untranslated) and closed (translated) boxes and numbered at the bottom.

spanning the coding sequence for chloroplastic AsAP isoenzymes were cloned from a partial genomic library of spinach leaves. Three positive clones, PAsAP1 (4.6 kbp), 2 (3.1 kbp), 3 (5.6 kbp) hybridized with tAsAP cDNA and were found to contain the distinct portion of the entire chloroplastic AsAP gene (ApxII). Therefore the three fragments contained a full-length, 8.5 kbp ApxII gene in addition to 0.3 and 4.5 kbp respectively of the 5′-upstream and 3′-downstream regions. Furthermore, PAsAP4, one of the clones harbouring 10.2 kbp restriction-endonuclease-HindIII fragments, was found to overlap the whole region of PAsAP1 and the upstream region of PAsAP2 by restriction-enzyme mapping and partial sequencing of PAsAP4. Synthetic primers corresponding to the vicinity of connection of each EcorI fragment shown in Figures 1 and 2 were used to identify the connection sites of the AsAP gene. The restriction map and cloning strategy of genomic clones encoding the chloroplastic AsAP isoenzymes are shown in Figure 1.

Organization of exon and intron structure

By alignment between the cDNAs for chloroplastic AsAP isoenzymes and the gene, the ApxII nuclear gene was found to contain 12 introns (Figures 1 and 2). The coding region of the chloroplastic AsAP gene consisted of 13 exons, referred to as 1 to 11, 12 (penultimate) and 13 (final) in order from 5′ to 3′ (Figures 1 and 2). The TATA box was found in positions −43 to −46, and two putative CCAAT boxes were found 42 and 67 bp upstream from the TATA box. Significant potential regulatory elements were not found upstream of the transcription start site. Primer-extension analysis identified the transcription start site within exon 1 (Figure 3); the 5′-terminus of exon 1 contained the long 5′-untranslated region and the ATG initiation codon. The cDNA clones encoding chloroplastic AsAP isoenzymes were more than 200 nucleotides shorter than expected from the primer extension analysis [12]. It is worth noting that three small upstream open reading frames (residues 16–156, 201–236 and 233–277), which may influence expression from the authentic one [17], are present in the 5′-leader sequence. Exons 1–11 encoded the common amino acid sequence for stromal and thylakoid-bound AsAP isoenzymes. The important point to note is the two 3′-terminal exons. Two splice acceptor sites occur in exons 12 and 13 separated by a 14 bp-encoding sequence (residues 7531–7544). The penultimate exon 12 (residues 7376–7530) consisted of one codon for Asp before the TAA termination codon, and the entire 3′-untranslated region including a potential polyadenylation signal (AATAAA) of the sAsAP mRNA. The final exon 13 (residues 7545–7756) consisted of the corresponding coding sequence of the hydrophobic C-terminal region, the TGA termination codon, and the entire 3′-untranslated region, including a potential polyadenylation signal (AATATA) of the tAsAP mRNA. These results show that the mRNAs for the chloroplastic AsAP isoenzymes arose from only one gene by alternative usage in the final two exons. For all of the coding exons, the intron–exon splice junctions were readily identifiable, which conformed to the consensus sequences GT at the donor site and AG at the acceptor site (Table 1), which was in agreement with those of general genomic genes [18]. The 5′-splice site at 7275 was used in combination with the 3′-splice site at 7544, generating a spliced transcript for tAsAP in which 270 nt of the intron 12 were removed.

Detection of chloroplastic AsAP isoenzyme mRNAs

To analyse the expression of mRNA for chloroplastic AsAP isoenzymes, the total RNAs from spinach leaves under normal conditions (non-stress conditions) were subjected to Northern blotting using labelled tAsAP cDNA as a hybridization probe. As a result, the mRNAs of chloroplastic AsAP isoenzymes were expressed under normal conditions with a length of 1.4 kb (results not shown). However, we could not distinguish the level of sAsAP mRNA from that of the tAsAP mRNA because of the similar size of the mRNAs for the chloroplastic AsAP isoenzymes. To provide the expression of both isoenzymes in detail, oligo(dT)-primed cDNAs were synthesized by reverse transcriptase from
Figure 2 For legend see opposite page
Figuure 2 Nucleotide and deduced amino acid sequences of the gene (ApxII). The deduced amino acid sequence from pumpkin mature tAsAP isoenzymes may provide a good example of a new type of gene regulation mechanism in higher plants. Recently, the cDNA gene both contain their own termination codons and potential polyadenylation signals respectively. The production of polyadenylation in the amino acid sequences by regulation systems such as alternative splicing, 3'-end cleavage and polyadenylation of mRNA precursors is well documented in animal systems [19,20]. In higher plants, heterogeneous C-terminal production by alternative splicing has been shown in spinach ribulosebisphosphate carboxylase/oxygenase (rubisco) activase [21]. In this case, the alternative usage of a part of the intron results in the synthesis of two rubisco activases that are different only in the presence or absence of their 37 additional amino acids at the C-terminal regions. Another example is revealed by pumpkin (Cucurbita sp. cv. Kurokawa Amakuri nankin) hydroxy-pyruvate reductase isoenzymes with or without a putative C-terminal signal for targeting to microbodies [22]. Although these cases are similar to spinach chloroplastic AsAP isoenzymes with respect to the production of C-terminal regions by alternative splicing, the generation of two different polypeptides from the resulting in-frame shift caused by insertion of part of their introns is different in the case of the ApxII gene. Two spinach chloroplastic AsAP isoenzymes generate the resulting alternative usage of the independent 3'-end nucleotide sequences which are located in the penultimate exon 12 and the final exon 13 of the gene (Figure 4). This is the first report describing the addition of a peptide sequence to a protein C-terminus by alternative 3'-end cleavage and polyadenylation. Therefore spinach chloroplastic AsAP isoenzymes may provide a good example of a new type of gene regulation mechanism in higher plants. Recently, the cDNA sequence encoding tAsAP from pumpkin has been reported [23]. The deduced amino acid sequence from pumpkin mature tAsAP showed a high degree of identity (83.7%) and a hydrophy profile similar to those of tAsAP from spinach. Interestingly, antibodies raised against the C-terminal 82-residue polypeptide of the pumpkin tAsAP cross-reacted with both tAsAP and

DISCUSSION
Here the first entire genomic structure (ApxII) encoding chloroplastic AsAP isoenzymes from spinach was identified. The ApxII gene consisted of essentially 13 exons interrupted by 12 introns. In previous studies, genomic structures encoding cAsAP isoenzymes (ApxI) have been characterized from pea [15] and Arabidopsis [16]. The structure of the ApxII gene for the chloroplastic AsAP isoenzymes from spinach was considerably different from those of the ApxI genes for cAsAP. The ApxI genes are interrupted by 9 introns, the first of which was located in the 5'-untranslated regions of the mature transcript. The intron in the 5'-noncoding region was not observed in the gene of ApxII (Figure 1).

From the results of a comparison with the ApxII genomic structure and cDNA sequences and genomic Southern blot, it is clear that the mRNAs of spinach chloroplastic AsAP isoenzymes are produced from a single gene by alternative exons 12 and 13 and share the same open reading frame, except for the hydrophobic thylakoid-membrane-binding domain. How the chloroplastic AsAP isoenzymes are produced by different selection of their 3'-ends is an interesting problem. One possibility is that primary transcripts with 3'-end heterogeneity arise from cleavage at different alternative polyadenylation sites. Intact exons 12 and 13 of the ApxII gene both contain their own termination codons and potential polyadenylation signals respectively. The production of heterogeneity in the amino acid sequences by regulation systems such as alternative splicing, 3'-end cleavage and polyadenylation of mRNA precursors is well documented in animal systems [19,20]. In higher plants, heterogeneous C-terminal production by alternative splicing has been shown in spinach ribulosebisphosphate carboxylase/oxygenase (rubisco) activase [21]. In this case, the alternative usage of a part of the intron results in the synthesis of two rubisco activases that are different only in the presence or absence of their 37 additional amino acids at the C-terminal regions. Another example is revealed by pumpkin (Cucurbita sp. cv. Kurokawa Amakuri nankin) hydroxy-pyruvate reductase isoenzymes with or without a putative C-terminal signal for targeting to microbodies [22]. Although these cases are similar to spinach chloroplastic AsAP isoenzymes with respect to the production of C-terminal regions by alternative splicing, the generation of two different polypeptides from the resulting in-frame shift caused by insertion of part of their introns is different in the case of the ApxII gene. Two spinach chloroplastic AsAP isoenzymes generate the resulting alternative usage of the independent 3'-end nucleotide sequences which are located in the penultimate exon 12 and the final exon 13 of the gene (Figure 4). This is the first report describing the addition of a peptide sequence to a protein C-terminus by alternative 3'-end exons in higher plants. Therefore spinach chloroplastic AsAP isoenzymes may provide a good example of a new type of gene regulation mechanism in higher plants. Recently, the cDNA sequence encoding tAsAP from pumpkin has been reported [23]. The deduced amino acid sequence from pumpkin mature tAsAP showed a high degree of identity (83.7%) and a hydrophy profile similar to those of tAsAP from spinach. Interestingly, antibodies raised against the C-terminal 82-residue polypeptide of the pumpkin tAsAP cross-reacted with both tAsAP and

**Table 1** Splice sites (emboldened), length and percentage A + T of introns in the gene encoding chloroplastic AsAP isoenzymes

<table>
<thead>
<tr>
<th>Intron</th>
<th>Length (bp)</th>
<th>5’-Splice site</th>
<th>3’-Splice site</th>
<th>A + T (%)</th>
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<tr>
<td>1</td>
<td>92</td>
<td>CAG GTTCT</td>
<td>TGCAG AGA</td>
<td>63.1</td>
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<tr>
<td>2</td>
<td>2521</td>
<td>ATG GAT</td>
<td>TGCAG GTT</td>
<td>63.2</td>
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<tr>
<td>3</td>
<td>527</td>
<td>CAG GTAAG</td>
<td>TGCAG GTC</td>
<td>67.4</td>
</tr>
<tr>
<td>4</td>
<td>93</td>
<td>GAG GCTTT</td>
<td>TTTAG GAG</td>
<td>71.0</td>
</tr>
<tr>
<td>5</td>
<td>575</td>
<td>CTG GTAAG</td>
<td>TTTAG ATG</td>
<td>64.0</td>
</tr>
<tr>
<td>6</td>
<td>89</td>
<td>AAG GTAAG</td>
<td>GATAG GAT</td>
<td>64.1</td>
</tr>
<tr>
<td>7</td>
<td>958</td>
<td>ACG GCTCT</td>
<td>ACTAG AAA</td>
<td>67.1</td>
</tr>
<tr>
<td>8</td>
<td>91</td>
<td>AAG GCTTT</td>
<td>TGCAG GAC</td>
<td>72.5</td>
</tr>
<tr>
<td>9</td>
<td>96</td>
<td>AAG GTAAG</td>
<td>TTTAG GTA</td>
<td>67.7</td>
</tr>
<tr>
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<td>753</td>
<td>GAG GTCTG</td>
<td>GACAG GGT</td>
<td>65.3</td>
</tr>
<tr>
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<td>101</td>
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<td>72.3</td>
</tr>
<tr>
<td>12</td>
<td>270</td>
<td>GAG GTCTG</td>
<td>TGCAG ACA</td>
<td>71.1</td>
</tr>
</tbody>
</table>

The primer 3'-1r was labelled with ³²P and annealed with 150 µg of total RNA from spinach leaves. The analysis was performed as described in the Materials and methods section. PE, sequencing ladder (A, G, C and T). The arrow indicates the major transcription initiation site.
sAsAP from pumpkin. It seems likely that the alternative-splicing mechanism at the 3′-end for the chloroplastic AsAP isoenzymes occurs in higher plants, including pumpkin. Recently, cDNA sequences of sAsAP (accession No. X98925) and tAsAP (accession no. X98926) from Arabidopsis have been appeared in the GenBank database. A comparison of nucleotide sequences of cDNAs for both sAsAP and tAsAP from Arabidopsis showed 66.1 % identity, suggesting that both isoenzymes may be encoded by different genes. However, mature native sAsAP and tAsAP proteins in Arabidopsis chloroplasts are not identical.

The existence of only one gene encoding the chloroplastic AsAP isoenzymes and the one TATA box in the 5′-end flanking region of the ApxII gene suggests that these enzymes are transcribed from the same promoter that is active. Preliminary evidence for transcriptional levels of ApxII products using reverse transcription PCR showed that the levels of mRNA for both chloroplastic AsAPs are in almost equal quantities throughout the spinach leaves under normal conditions (results not shown). The activity of tAsAP was almost the same as that of sAsAP in spinach chloroplasts [5]. These results clearly support the hypothesis that the localization of the AsAP isoenzymes in both the thylakoid membrane and the stroma soluble fraction has important implications in the protection from photo-oxidative stress caused by active oxygen species, including H₂O₂, in the chloroplasts of higher plants.

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REFERENCES

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