Isolation of 1-aminocyclopropane-1-carboxylate synthase gene from *Oncidium*
Gower Ramsey

G.H. Yang1,2 and J.P. Liu1

1Key Laboratory of Protection and Development Utilization of Tropical Crop Germplasm Resources, Ministry of Education, College of Agronomy, Hainan University, Haikou, Hainan Province, China
2Sanya Science & Technology Academy for Crop Winter Multiplication, Sanya, China

Corresponding author: J.P. Liu
E-mail: liu3305602@163.com

Received September 26, 2013
Accepted February 21, 2014
Published October 20, 2014
DOI http://dx.doi.org/10.4238/2014.October.20.24

ABSTRACT. A full-length cDNA of a 1-aminocyclopropane-1-carboxylate synthase (ACS) family member from *Oncidium*, named *OnACS1* (GenBank accession No. JQ822087) was cloned and characterized by reverse transcription polymerase chain reaction and rapid amplification of cDNA ends technology. The full-length cDNA was 1586 bp, including a 1308-bp open reading frame, a 105-bp 5′ untranslated region (UTR), and 173-bp 3′ UTR, encoding 436 amino acids. The deduced amino acid sequence of *OnACS1* shares 85, 84, and 83% homology with ACS proteins of *Cattleya bicolor*, *Dendrobium crumenatum*, and *Phalaenopsis* hybrid, respectively. Prokaryotic expression and sodium dodecyl sulfate polyacrylamide gel electrophoresis showed that a specific band was produced and was consistent with the predicted protein size. A tissue-specific manner of *OnACS1* expression was observed, and it was predominantly expressed in the gynostemium. The *OnACS1* expression in the sepals and
Isolation of the OnACS1 gene from Oncidium Gower Ramsey

... was upregulated by 1% ethephon treatment.

**Key words:** 1-Aminocyclopropane-1-carboxylate synthase; Oncidium; Ethylene; Floral senescence

**INTRODUCTION**

Ethylene, a small gaseous phytohormone, plays an important regulatory role in various aspects of plant growth and development such as ripening and senescence (Abeles et al., 1992; Yang and Hoffman, 1984). In the biosynthesis of ethylene in higher plants, the last two steps are the conversion of S-adenosylmethionine to 1-aminocyclopropane-1-carboxylic acid (ACC), which is catalyzed by ACC synthase (ACS), and the conversion of ACC to ethylene, which is catalyzed by ACC oxidase (Adams and Yang, 1979; Yang and Hoffman, 1984; Kende, 1993). Because ACS is generally considered as the rate-limiting step in the committed ethylene biosynthetic pathway, a significant effort has been directed at the isolation and characterization of ACS. ACS is composed of a larger gene family (Zarembinski and Theologis, 1994; Lin et al., 2009), and genes encoding ACS have been isolated and characterized from a number of flowering plants such as carnation (Park et al., 1992; Henskens et al., 1994) and Phalaenopsis orchid (Bui and O’Neill, 1998).

The Oncidium Gower Ramsey is a commercially important orchid cut flower. We are interested in the regulation of ethylene biosynthesis in Oncidium Gower Ramsey flowers during senescence. As a first step toward this goal, we report that an ACS gene was isolated from Oncidium Gower Ramsey flower tissue, and we show that it was differentially expressed in different organs.

**MATERIAL AND METHODS**

**Plant material**

Greenhouse-grown Oncidium Gower Ramsey ‘Gold2’ plants were obtained from a commercial grower. Roots, stems, leaves, and floral parts (labella, petals, sepals, and gynandria) were collected in liquid N\(_2\), pulverized into a fine powder, and stored at -80°C until subsequent use in RNA isolation. The effects of ethephon on ACS gene expression in sepals and gynandria were tested. Floral parts were smeared with 1% ethephon and sealed in plastic bags for different lengths of time (0, 6, 12, 24, and 48 h). Tissue samples of sepals and gynandria were then frozen in liquid nitrogen and stored at -80°C until RNA extraction.

**RNA extraction and OnACS1 gene isolation**

Total RNA was extracted from frozen tissue of floral parts using a previously described method (Yang and Liu, 2011). The first strand of cDNA was synthesized using the RevertAidTM First Strand cDNA Synthesis Kit (MBI Fermentas, JINGMEI BIOTECH CO., LTD., Shenzhen, China) according to the manufacturer protocol. To amplify the cDNA fragment of OnACS1, we used specific primers ACS-F and ACS-R (Table 1) that were designed from the given nucleotide sequences registered on the database (accession Nos. AY504664, EF488013, and AF004663). Reactions were subjected to 30 cycles of 95°C for 30 s, 54°C for 1 min, and 72°C for 1 min.
Table 1. Primer sequences used in this study.

<table>
<thead>
<tr>
<th>Primer name</th>
<th>5'-3' Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACS-F</td>
<td>ACAGCTCCAAATGGCCTCAAACGTTCTTCTCTCTC</td>
</tr>
<tr>
<td>ACS-R</td>
<td>GAAACAGAGCTGAAACTGACTGAGGAGGGTTCTCCTT</td>
</tr>
<tr>
<td>Mi13-F</td>
<td>GACCCGCGTAAAGCTATTTCTAGTGAACGGAGG</td>
</tr>
<tr>
<td>Mi13-R</td>
<td>CGCCAGGGTTTTCCAGAGTTGAC</td>
</tr>
<tr>
<td>3'-RACE-1</td>
<td>TCTCTGATCTCATACAAGTCTGCTCCG</td>
</tr>
<tr>
<td>3'-RACE-2</td>
<td>CAGATGTGGTGCTCTGATTTTCAATGACAGAG</td>
</tr>
<tr>
<td>5'-RACE-1</td>
<td>GAAGAGAAATTGTAGCAGTGGTGCGAGAGG</td>
</tr>
<tr>
<td>5'-RACE-2</td>
<td>GAGAGAGTGAGTTGGAAGCCATTGGAGC</td>
</tr>
<tr>
<td>ACS_orf-F</td>
<td>CGCGAGCAGATGCAAGTGTCTTTCTTCAGAAGAGG</td>
</tr>
<tr>
<td>ACS_orf-R</td>
<td>CGCGAGCAGATGCAAGTGTCTTTCTTCAGAAGAGG</td>
</tr>
<tr>
<td>Actin-F</td>
<td>GCACACTTCTCTGCTAGCAGGAGG</td>
</tr>
<tr>
<td>Actin-R</td>
<td>TCTCTGATCTCATACAAGTCTGCTCCG</td>
</tr>
<tr>
<td>ACS-RT-F</td>
<td>CGCGAGCAGATGCAAGTGTCTTTCTTCAGAAGAGG</td>
</tr>
<tr>
<td>ACS-RT-R</td>
<td>CGCGAGCAGATGCAAGTGTCTTTCTTCAGAAGAGG</td>
</tr>
</tbody>
</table>

The polymerase chain reaction (PCR) products were purified with the TaKaRa Agarose Gel DNA Purification Kit Ver.2.0 [TaKaRa Biotechnology (Dalian) Co., Ltd., Dalian, Liaoning Province, China] and ligated into the pMD 18-T vector according to the manufacturer instructions. The ligation products were transformed into *Escherichia coli* DH5α cells. Recombinant bacteria were selected and the DNA sequences were determined. In order to obtain the full-length nucleotide sequence of OnACS1, rapid amplification of cDNA ends (RACE)-PCR was performed using the 3'-full RACE Core Set and the 5'-full RACE Core Set (Takara, China) according to the user manual. The 3'-end fragments were amplified using specific primers 3'-RACE-1 and 3'-RACE-2, and the 5'-end fragments were amplified using 5'-RACE-1 and 5'-RACE-2. Each primer was designed based on the nucleotide sequences of the cDNA fragments obtained from the reverse transcription (RT-PCR) described above (Table 1). The PCR products were purified, ligated into the pMD 18-T vector, introduced into *E. coli* DH5α, and sequenced. The full-length sequence of OnACS1 was obtained with 5'-end fragments, the cDNA fragments, and 3'-end fragments using the DNAMan software. Amino acid sequence analysis and multiple sequence alignment were accomplished using the DNAMan and Bioedit software. Phylogenetic analysis of ACS amino acid sequences was conducted using the MEGA software.

Expression of recombinant proteins

The open reading frame (ORF) of OnACS1 was analyzed with the ORF finder software. The coding region of the OnACS1 gene was amplified by PCR using the following two primers (Table 1): ACS_orf-F (sense) contains a BamHI restriction site (underlined) immediately upstream of the ATG translation initiation codon, and ACS_orf-R (antisense) was constructed with a Sall restriction site (underlined) located downstream of the stop codon of the OnACS1 ORF. The PCR product was verified by sequencing, digested with the appropriate restriction enzymes, and directionally ligated into the corresponding sites of the pGEX-6P-1 expression vector system. The resulting plasmid was introduced into *E. coli* BL21 competent cells, and the transformed cells were cultured at 37°C in liquid Luria-Bertani medium with 100 mg/mL ampicillin. Isopropyl-1-thio-b-D-galactoside (IPTG) was added to the shaking culture to a final concentration of 1 mM. The cells were harvested for sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) analysis when the cell density reached an optical density at 600 nm of 0.6-0.8.
Isolation and characterization of the OnACS1 gene

Using the total RNA isolated from the floral tissues as a template, the first strand of cDNA was amplified. Using the conservative sequence of the registered ACS genes of Cattleya (accession No. AY504664), Dendrobium (accession No. EF488013), and Phalaenopsis (accession No. AY504664), the OnACS1 transcript levels were determined by fluorescence quantitative PCR analysis of the total RNA fractions that were obtained from the different organs of Oncidium Gower Ramsey “Gold2” plants and the sepals and gynandria that were treated with ethephon. The RT reaction was conducted in a reaction mixture containing 2 μL total RNA, 1 μL 50 μM Oligo dT Primer, 4 μL 5X PrimeScript Buffer, 1 μL PrimeScript RT Enzyme Mix I, 1 μL 100 μM Random 6-mers, and 11 μL RNase-free dH2O in a total volume of 20 μL. The RT reaction was incubated for 15 min at 37°C and for 5 s at 85°C. The real-time PCR was performed in a reaction volume of 20 μL containing 2.0 μL RT reaction product, 0.4 μL each of the upstream and downstream primers (ACS-RT-F and ACS-RT-R, 10 μM), 10 μL 2X SYBR Premix Ex Taq II, 0.4 μL 50X ROX Reference Dye or DYE II, and 6.8 μL dH2O with a 7500 Real-Time PCR System (Applied Biosystems, USA). The PCR was carried out for 30 s at 95°C, followed by 40 cycles of 5 s at 95°C and 34 s at 60°C, and supplemental incubation for 15 s at 95°C, 60 s at 60°C, and 15 s at 95°C. Absolute values for transcript abundance from RT-PCR were quantified using the 2^-ΔΔCT method by Kenneth and Thomas (2001).
sion No. AF004663), the gene-specific primers ACS-F and ACS-R were designed. Following 30 cycles of PCR amplification using the first strand of cDNA that was synthesized as a template, a product of the expected size (400 bp) was easily visualized with 1% agarose gel electrophoresis (data not shown). Nested PCR primers were used to amplify ACS sequences in two rounds of PCR using 3'-RACE-1 and 5'-RACE-1 as the first-round primers and 3'-RACE-2 and 5'-RACE-2 as the second-round primers. Then, 3' fragments of 720 bp and 5' fragments of 700 bp were obtained (data not shown). The resulting PCR products described above were sequenced and spliced to obtain the full-length cDNA with the DNAman software (DNAMAN 6.0, USA). The full-length cDNA was verified with amplification of the coding region of the *OnACS1* gene, agarose gel electrophoresis, and subsequent sequencing (data not shown). The *OnACS1* gene sequence is 1586 bp and contains a 1308-bp ORF encoding 436 amino acids (Figure 1). The ORF is preceded by a 5' UTR of 105 bp and followed by a 3' UTR of 173 bp. A comparison of the deduced OnACS1 amino acid sequence and amino acid sequences for ACSs from other plants revealed that the *Oncidium* ACS protein shares 85, 84, and 83% homology with ACS proteins of *Cattleya bicolor* (accession No. AAR88653), *Dendrobium crumenatum* (accession No. AAB67882), and *Phalaenopsis* hybrid (accession No. AAR00507), respectively. Phylogenetic analysis of amino acid sequences of plant ACSs (Figure 2) shows that the *Oncidium* ACS protein is clustered with the ACS proteins of *Cattleya bicolor*, *Dendrobium*, and *Phalaenopsis* orchids. Therefore, we designed the *Oncidium* ACS as OnACS1 (GenBank accession No. JQ822087).

Figure 1. Nucleotide sequence and predicted protein sequences of OnACS1. The open reading frame is indicated by upper case letters. The 5' and 3' untranslated regions are indicated by lower case letters. The deduced amino acid sequence is given in upper case letters under the nucleotide sequence. The translation stop codon is indicated by an asterisk.
Isolation of the OnACS1 gene from Oncidium Gower Ramsey

Expression analysis of recombinant OnACS1 was conducted with a translational fusion of the OnACS1 ORF into E. coli using the prokaryotic expression vector pGEX-OnACS1. SDS-PAGE analysis showed specific expression that was induced by 1 mM IPTG at 37°C for 4 h. The molecular weight of the protein agreed with the size of the predicted protein (Figure 3). The gas chromatographic results revealed that a small peak appeared, indicating that ethylene production could be detected, and the enzymatic activity of the recombinant OnACS1 protein was determined (data not shown).

Analysis of OnACS1 prokaryotic expression and enzymatic activity

Figure 2. Phylogenetic analysis of amino acid sequences of plant ACC synthase family. The phylogenetic tree constructed from an optimal alignment of proteins using the MEGA software. Numbers represent amino acid sequence identity between members of the subfamily.

Figure 3. SDS-PAGE analysis of recombinant OnACS1. Lane 1 = pGEX-OnACS1 proteins from cells induced by IPTG (1 mM) at 37 °C for 4 h. Arrow indicates the expected band for the pGEX-OnACS1 proteins; lane 2 = pGEX-OnACS1 proteins from uninduced Escherichia coli; lane 3 = pGEX-OnACO1 from cells induced by IPTG (1 mM) at 37°C for 4 h; lane 4 = pGEX-OnACO1 proteins from uninduced E. coli; lane 5 = Proteins from cells transformed with the empty vector pGEX induced by IPTG (1 mM) at 37 °C for 4 h; lane 6 = Proteins from uninduced transformed with the empty vector pGEX; lane M = molecular weight marker.
Analysis of \textit{OnACS1} gene expression with fluorescence quantitative PCR

The expression of the \textit{OnACS1} gene in various tissues of \textit{Oncidium} Gower Ramsey “Gold2” was investigated with fluorescence quantitative PCR analysis. The expression of the \textit{OnACS1} gene predominantly occurred in gynandria, and lower levels of \textit{OnACS1} gene expression were detected in roots, stems, leaves, lip, petals, and sepals, and the \textit{OnACS1} expression was slightly higher in the sepals than in the other tissues (Figure 4). The effects of ethephon treatment of the floral tissues on \textit{OnACS1} gene expression in the gynandria and sepals were studied. \textit{OnACS1} gene expression increased in the sepals with the duration of ethephon treatment, and expression slightly increased in the gynandria in response to ethephon (Figure 5).

\textbf{Figure 4.} Expression of \textit{OnACS1} gene in various tissues of \textit{Oncidium} Gower Ramsey “Gold2” determined by fluorescence quantitative PCR analysis. Values are reported as means of three replicates ± SE.

\textbf{Figure 5.} Changes in \textit{OnACS1} gene expression in gynandria and sepals after treatment of the floral tissues with 1% ethephon for different time. Values are reported as means of three replicates ± SE.
DISCUSSION

In this report, the full-length cDNA of OnACS1 was isolated and sequenced from Oncidium flowers by the RT-PCR and RACE methods. The ORFs of the reported ACS genes are generally 1200-1300 bp and encode about 430 amino acids, including the Dendrobium ACS gene (AAB67882) encoding 435 amino acids and the Cattleya ACS gene (AAR88863) encoding 434 amino acids. The cloning and characterization of the OnACS1 gene indicated its ORF is 1308 bp and encodes 436 amino acids. Alignments of deduced amino acid sequences of ACC synthases from Oncidium with those of Cattleya bicolor, Dendrobium crumenatum, and Phalaenopsis hybrid showed that these sequences share high identity and are highly conserved. These results indicate that the cloned gene is a typical member of the Oncidium ACS gene family. Analysis of OnACS1 prokaryotic protein expression and enzymatic activity further confirmed this.

Many studies reported that the ACS genes display distinct spatial and temporal expression patterns (ten Have and Woltering 1997; Trebitsh et al., 1997; Bui and O’Neill, 1998; Jones and Woodson, 1999; Yamasaki et al., 2001; Yamagami et al., 2003; Tsuchisaka and Theologis, 2004; Peng et al., 2005; Salman-Minkov et al., 2008). The expression of the OnACS1 gene in all tissues tested suggests that it is constitutively expressed but at different levels: high levels in gynandria, moderate levels in sepals, and low levels in other organs. Ethephon, a chemical that can be converted into ethylene upon metabolism by the plant, significantly promoted OnACS1 gene expression in the sepals and slightly promoted expression in the gynandria. These results demonstrated that it was differentially regulated in a tissue-specific manner. Moreover, OnACS1, as an ethylene biosynthetic gene, was induced by ethephon treatment, indicating that ethylene is the main regulator of its expression. A detailed study of the role of OnACS1 in ethylene production and floral senescence of Oncidium is underway.

ACKNOWLEDGMENTS

Research supported by the Hainan University Fund (#2012hekled-7).

REFERENCES


