Full Length Research Paper

Cloning and differential expression of 1-aminocyclopropane-1-carboxylate synthase gene in different floral tissues of *Dendrobium* ‘anna’ flowers

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Cloning of the DenACS from *Dendrobium* hybrid cultivar Anna was performed by RT-PCR and nucleotide sequence analysis revealed that the open reading frame of this gene was 1,308 bp in length, encoded for a protein of 435 amino acid residues. The calculated molecular mass of the deduced polypeptide is 48.5 kDa and the predicted isoelectric point is 5.84. The deduced amino acid sequence of the DenACS-encoded protein showed a high degree of identity to those of the ACS from petunia, geranium, carnation and rose. The expression of DenACS was examined by RT-PCR and the results revealed that this gene was highly expressed in flower stage 2 (partially opened flower) and stage 3 (full opened flower) during flower development. It was also expressed in all floral tissues and organs including petal, sepal, pedicel, labellum, stigma, young leaves and roots, but the maximum expression was observed in petal, sepal and pedicel. These results indicated that the orchid DenACS is involved in the flower opening, flower senescence as well as in vegetative cell growth and development.

Key words: Ethylene biosynthesis, 1-aminocyclopropane-1-carboxylic acid (ACC), ACC synthase (ACS), gene expression, orchid (*Dendrobium* spp.).

INTRODUCTION

The phytohormone ethylene plays a key role in various aspects of plant growth and development, e.g., seed germination, stem elongation, leaf and flower senescence and fruit ripening and abscission (Abeles et al., 1992). It also acts as an important signaling molecule in plants responses to a range of both biotic and abiotic stimuli such as pathogen attack, mechanical damage (Johnson and Ecker, 1998; Bleecker and Kende, 2000), flooding (Olson et al., 1995) and chilling (Lelievre et al., 1997).

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Abbreviations: PCR, Polymerase chain reaction; RT-PCR, reverse transcriptase -PCR; ACC, 1-aminocyclopropane-1-carboxylic acid; ACS, ACC synthase; Met, methionine; AdoMet, S-adenosylmethionine; EFE, ethylene-forming enzyme; ORF, open reading frame; CDP, cytidine diphosphate; dNTP, deoxyribonucleoside 5’-triphosphate; PLP, pyridoxal 5’-phosphate.

Ethylene is synthesized in plant tissues via the conversion of methionine (Met) to S-adenosylmethionine (AdoMet), catalyzed by the enzyme S-adenosylmethionine synthase. AdoMet serves as an intermediate in a number of biosynthetic pathways, including the production of polyamines. AdoMet is then transformed to 1-aminocyclopropane-1-carboxylate (ACC) by the ACC synthase (ACS, S-adenosyl-L-methionine methylthioadenosine-lyase, EC 4.4.1.14), which is most likely a cytoplasmic enzyme. In the last step, ACC is converted to ethylene, CO₂ and HCN by the ACC oxidase (ACO, EC 1.4.3), formerly known as the ethylene-forming enzyme (EFE) (Kende, 1993). Ethylene biosynthesis is mainly regulated through the expression of ACS genes (Yang and Hoffman, 1984), but in some cases the regulation can also be through the expression of ACO genes, e.g., in melon (Yamamoto et al., 1995; Lasserre et al., 1996), tomato (Barry et al., 1996), marsh dock (Vriezen et al., 1999), peruvian lily (Wagstaff et al., 2005) and plum (Fernandez-Otero et al., 2006).

ACC synthases have been cloned from various plant
species, including winter squash (Nakajima et al., 1990), tomato (Olson et al., 1991; Rottmann et al., 1991), zucchini (Huang et al., 1991), apple (Dong et al., 1991), Arabidopsis (Liang et al., 1992; Van Der Straeten et al., 1992), mung bean (Botella et al., 1992), tobacco (Bailey et al., 1992), carnation (Park et al., 1992), rice (Zarembinski and Theologis, 1993), orchid (Bui and O'Neill, 1998) and rose (Wang et al., 2004; Ma et al., 2006). It has been shown in several cases that ACC synthase genes are encoded by a multigene family and they are differentially expressed in response to various signals such as wounding, fruit ripening, flower senescence and auxin (Nakajima et al., 1990; Van Der Straeten et al., 1990; Dong et al., 1991; Olson et al., 1991; Rottmann et al., 1991; Bui and O'Neill, 1998; Ma et al., 2006). The expression of ACC synthase genes in some systems is also regulated in a tissue-specific manner during plant growth and development, e.g., in carnation, the CARACC3 mRNA is mostly abundant in the petals while the CARAS1 mRNA is mostly abundant in the styles (Park et al., 1992; Henskens et al., 1994). In rose, three ACC synthase genes have been isolated and the Rh-ACS1 is transcribed specifically in response to wounding whereas the Rh-ACS2 is detectable only in senescent petals. With respect to the Rh-ACS3, formerly known as RKacc7, it is specifically expressed in petals, ovary and sepals and its expression is increased dramatically as the flower matured to senescence (Wang et al., 2004; Ma et al., 2006). According to the orchid flower, little is known about tissue and cell specificity of expression of ACC synthase gene during flower development.

Previously, we have cloned and analyzed the expression of ACO gene, DenACO, from Dendrobium hybrid cultivar Anna, one of the most important cut flower for ornamental industrial of Thailand and found that DenACO is highly expressed in petal, labellum and stigma during flower opening, suggesting that it plays a crucial role in the flower development process (Nag tong et al., 2009). To gain a better understanding of the ethylene response in different stages of flower development as well as in different floral tissues, cloning and expression of ACC synthase gene was investigated in this study.

**MATERIALS AND METHODS**

**Plant materials**

Orchid flowers (Dendrobium hybrid cultivar Anna) were harvested from a local commercial greenhouse (Nakhon Pathom, Thailand). The flowers were immediately put in tap water after harvested and then transported to the laboratory within 2 h. After being cut to 20 cm under water, the flowers were placed in deionized water (DW). Flowers at different stages and floral tissues including sepal, petal, pedicel, labellum and stigma were collected, immediately frozen in liquid nitrogen and stored at -20°C for further processing.

**Total RNA extraction**

For cDNA cloning, total RNA was extracted from petals using RNeasy Plant Mini Kit (QIAGEN, Germany). The extraction of RNA was carried out essentially as recommended by the manufacturer's instruction except that RNase-free DNase I was added to a final concentration of 0.1 mg/ml to remove any contamination genomic DNA.

**Gene amplification, cloning and sequencing**

Amplification of the Dendrobium ACS gene was performed by Reverse Transcription-PCR (RT-PCR) using degenerated oligonucleotide primers designed based on the ACS sequences of the Dendrobium hybrid cultivar Pompadour (EF488013), Karen (EF488014), Dendrobium crumenatum (U64031) and Cattleya spp. (AY504664). An internal fragment of the ACS gene was amplified using the forward ACSFor1 (5'-TCCTAGCCCGATCGCGCAGA-3') and reverse ACSRor1 (5'-ACACA(G)/C)AAATTTCTGCTCTCT-3'). The RT-PCR reaction was carried out using OneStep RT-PCR Kit (QIAGEN, Germany). The reaction mixture (50 µl) consisted of 10 µl of 5×QIAGEN OneStep RT-PCR buffer, 400 µM dNTP, 0.6 µM of each primer (forward and reverse primers), 2 µl of QIAGEN OneStep RT-PCR enzyme mix and 1 µl of RNA template. The RT reaction was carried out at 45°C for 45 min. Following an initial denaturation of template cDNA at 95°C for 15 min, 40 cycles of the following temperatures were used: denaturation at 94°C for 1 min, annealing at 55°C for 2 min and extension at 72°C for 3 min. The amplified product was separated on 0.7% agarose gel and purified using the NucleoSpin Extract II Kit (Macherey-Nagel, Germany). The purified PCR product was cloned into pGEM T-easy vector (Promega, Madison, WI, USA) and was transformed into the competent E. coli JM109 using the Rapid DNA Ligation and Transformation Kit (Fermentas, USA). After screening, target DNA was sequenced by the dideoxy nucleotide chain termination method using the MegaBACE 1000 automated DNA sequencer (Pharmacia Biotech, Sweden). In order to obtain the sequence of 5'- and 3'-end of the ACS gene, the SMART™ RACE cDNA Amplification Kit (ClonTech, USA) was used. The reaction was carried out essentially as recommended by the manufacturer's instruction. The complete full length open reading frame (ORF) of the ACS gene was confirmed by RT-PCR using the forward ACSFor2 (5'-ATGTCACAAAGATTGGGGATG-3') and reverse ACSRor2 (5'-CTTATATATTCTGCTTCTCTT-3'). After amplification reaction, the amplified product was separated on the gel, purified, cloned and sequenced as previously described. The sequence of the ACS gene and deduced amino acid sequence was analyzed using GENETYX (Software Development, Tokyo, Japan), while homology searching was performed using FASTA and BLAST program in the GenBank and DDBJ databases.

**Southern hybridization analysis**

The ACS gene(s) in the orchid genome was investigated by Southern blotting. Genomic DNA was extracted from petals of orchid flowers using NucleoSpin Extract II Kit (QIAGEN, Germany). The extraction of genomic DNA was carried out essentially as recommended by the manufacturer's instruction. DNA samples (20 µg) were digested with BamHI, EcoRI, HindII or PstI, electrophoresed on 0.7% agarose gel and transferred to Hybond N* positively charged nylon membrane (Schleicher and Schuell) by overnight capillary transfer. The DNA was cross-linked to the membrane by UV irradiation and baked at 80°C for 2 h. All procedures were performed according to the standard methods as described by Sambrook and Russell (2001).

PCR fragment (approximately 650-bp in length) containing part of the coding region of ACS gene of orchid was used as a DNA probe. Labeling of the DNA probe with alkaline phosphatase for use in conjunction with chemiluminescent detection with CDP-Star was
performed using Gene Images AlkPhos Direct Labeling and Detection System (AlkPhos Direct™, Amersham). All procedures for DNA labeling were carried out as recommended by the manufacturer. The Hybond N⁺ positively charged nylon membrane with cross-linked DNA was prehybridized at 55°C for 2h in hybridization oven (Hybrid limited equipment class I) with gently rotation. After incubation, the labeled-DNA probe was added and subsequently incubated at 55°C an overnight. The membrane was washed according to the standard procedure for Southern blot analysis. After washing, the CDP-Star detection reagent (AlkPhos Direct™, Amersham) was added onto the membrane and the membrane was left at room temperature for 5 min. The membrane was then exposed to X-ray film (hyper film, Amersham) for 1 h, thereafter the film was developed and fixed with developer and fixer solution (Kodak), respectively.

RT-PCR analysis

The expression level of the orchid ACS gene was determined by RT-PCR. Total RNA was isolated from petals of orchid flowers at different developmental stages (stages 1 - 3) (Figure 1A) and from different floral tissues including sepal, petal, pedicel, labellum, stigma (Figure 1B), young leaves and roots. RT-PCR was carried out using OneStep RT-PCR Kit (QIAGEN, Germany) with the forward ACSFor2 and reverse ACSRor2 primers synthesized base on the 5'- and 3'-region of the orchid ACS gene. The reaction mixture (50 μl) consisting of 10 μl of 5xQIAGEN OneStep RT-PCR buffer, 400 μM dNTP, 0.6 μM of each primer (forward and reverse primers), 2 μl of QIAGEN OneStep RT-PCR enzyme mix and 1 μg of RNA template. As a control, 10 μg samples of total RNA were subjected to agarose gel electrophoresis (0.9% agarose) and stained with ethidium bromide. Actin gene was used as an internal control. A thermocycler was used to perform 1 cycle of 45 min at 50°C for reverse transcription followed by 40 cycles of 1 min at 94°C, 1 min at 55°C and 2 min at 72°C. The amplified products at the 28th, 31st, 34th, 37th and 40th cycle were electrophoresed on a 0.9% agarose gel. After staining with ethidium bromide, the relative amounts of the products were compared using the Gel Image Master (Pharmacia Biotech). The experiment was repeated at least twice. Under these conditions, the OneStep RT-PCR was able to specifically detect mRNA because no band was observed when reverse transcriptase was omitted.

RESULTS AND DISCUSSION

Cloning and sequencing analysis of the ACS gene from Dendrobium 'Anna' flower

The ACS gene of the Dendrobium hybrid cultivar Anna was cloned as described in materials and methods. After sequencing, a complete full length open reading frame encoding the entire amino acid sequence of the ACS gene product was obtained. The nucleotide sequence of this open reading frame, designated as DenACS, its flanking region and the deduced amino acid residues were shown in Figure 2. The DenACS sequence has been deposited in the GenBank nucleotide sequence database under the accession number GU138671. The DenACS contained 1,308 bp of the coding sequence, encoded a protein of 435 amino acid residues. The calculated molecular mass of the deduced polypeptide is 48.5 kDa and the predicted isoelectric point is 5.84. The highly conserved regions among all ACS sequences
Figure 2. Nucleotide and deduced amino acid sequences of the orchid DenACS. Nucleotides are numbered from the first nucleotide from 5’ end of the sequence. Amino acids are indicated below the nucleotide sequence in single-letter codes. Translation stop codon is indicated by an asterisk. Seven conserved regions observed in ACS are shaded, and the residues in the boxes are the conserved residues in all ACS and amino acid transferases. The active site lysine which binds to pyridoxal 5’-phosphate (PLP) and AdoMet is indicated as a bold letter in the box.

were identified in the DenACS (Henskens et al., 1994). One of the conserved regions contained the active site lysine residue (YSLSKDLGLPGFRVG) which binds pyridoxal 5’-phosphate (PLP) and AdoMet (Yip et al., 1990). The amino acids conserved among amino-transferases and ACC synthases (Huang et al., 1991; Rottmann et al., 1991) were also found in the DenACS gene product (Figure 2).

The deduced amino acid sequence of the Dendrobium hybrid cultivar Anna, DenACS, showed 95, 94, 92, 88 and 85% amino acid identity to the ACS from Dendrobium hybrid cultivar Karen (EF488014), Pompadour (EF488013), Dendrobium crumenatum (U64031), Cattleya spp. (AY504664) and Phalaenopsis spp. (AF004663), respectively.
Expression analysis of the DenACS gene

It has been reported in several plant species that the expression of ACS gene occurs during different developmental stages of flowers (Henskens et al., 1994; Bui and O'Neill, 1998; Wang et al., 2004; Ma et al., 2006). Therefore, the DenACS gene expression in orchid ‘Anna’ flowers was compared during three developmental stages from unopened bud to fully opened flower (Figure 1). As shown in Figure 6, the accumulation level of the DenACS transcripts in the flower stage 2 (partially opened flower) and stage 3 (full opened flower) was approximately 3 - 4 times higher than that observed in the flower stage 1 (unopened bud). In the previous study, we also found that the expression level of ACC oxidase gene, DenACO, is dramatically increased in the flower stages 2 and 3 (Nag tong et al., 2009). These findings suggested that DenACS and DenACO play a crucial role in flower opening and senescence, as previously described by Yamamoto et al. (1994), Shibuya et al. (2000) and Nukui et al. (2004). Our results are consistent with that of Wang et al. (2004) who reported that the accumulation of the rose ACS gene, RKacc7, is increased at the onset of flower senescence. The major accumulation of the ACS transcripts has also been observed at the start of flower senescence in carnation (Park et al., 1992) and Arabidopsis (Van Der Straeten et al., 1992).

Various expression levels of the ACS gene in the floral tissues have been reported in several plant species. In carnation, for example, the accumulation of the CARACC3 transcripts is mostly abundant in the petals whereas that of the CARAS1 transcripts is mostly abundant in the styles (Park et al., 1992; Henskens et al., 1994). In rose, the Rh-ACS1 is transcribed specifically in response to wounding whereas the Rh-ACS2 is detectable only in senescent petals. In addition, the Rh-ACS3 is specifically expressed in petals, ovary and sepal and its expression is increased dramatically as the flower matured to senescence (Wang et al., 2004; Ma et al., 2006). With respect to the present study, the expression of the DenACS was detected in all floral tissues including petal, sepal, pedicel, labellum and stigma, but the maximum expression was illustrated in petal, sepal and pedicel. The accumulation level of the DenACS transcripts in these tissues was approximately 3 times higher than those observed in labellum and stigma (Figure 7). The induction of the DenACS in these floral tissues may be triggered by the aging and senescence of the orchid flowers, like that observed in carnation (Henskens et al., 1994).

The accumulation of the DenACS transcripts was detected in both young leaves and roots of the orchid plants, but their accumulation levels were not significantly different (Figure 8). Our results are similar to that of Rodrigues-Pousada et al. (1993) who reported that the expression of ACS is high in young tissues. Therefore, we speculated from these results that the DenACS is involved in cell growth and development process.

There is a diverse group of factors that have been described in numerous plant species that modulate differential expression of the ACS genes, e.g. flooding, drought, wounding, pathogen attack, flower pollination and senescence and auxin (Nakajima et al., 1990; Van Der Straeten et al., 1990; Dong et al., 1991; Olson et al., 1991; Rottmann et al., 1991; Abeles et al., 1992; Woeste et al., 1999; Bui and O'Neill, 1998; Ma et al., 2006). Whether the DenACS expression is regulated by these factors remains to be investigated.

The opening and senescence of the orchid flowers are complex processes, involving several internal developmental factors such as ethylene (Yamamoto et al., 1994; Van Doorn, 2002). Based on data studies carried out in our laboratory, we hypothesize that ethylene (or unknown internal developmental factors) induces the expression of DenACS, leading to increased ACC synthase activity. Newly formed ACC is then translocated throughout the floral organs, including petal, sepal, pedicel, labellum and stigma, resulting in the induction of DenACO in these organs. As a result, ACC is finally oxidized to ethylene by

(Figure 3). Although Dendrobium hybrid cultivar Anna is belonged to the same genus with Dendrobium hybrid cultivar Karen, Pompadour and D. crumenatum, 5 - 8% difference in amino acid sequences was detected. This might be due to a difference in the plant genotype, since all these cultivars are hybrid lines. A comparison of the ACS from other plant species in the GenBank database with DenACS showed that DenACS had 55% amino acid sequence homology to ACS from petunia (AF049711), 54% to ACS from geranium (U88971) and carnation (X66605) and 39% to ACS from rose (AY378152) (Figure 4). The amino acid sequences of the highly conserved active site region of ACS from various species were found. This region contained the lysine residue responsible for pyridoxal 5'-phosphate binding which is probably essential for functioning of the ACS protein (Yip et al., 1990). However, four amino acid changes (G/S, L/I, L/M and L/F) were observed between the species, suggesting that the ACS(s) from these species are different (Figure 4) (Kim et al., 1992).

A multigene family for ACS sequences has been reported in zucchini (Huang et al., 1991), tomato (Olson et al., 1991), Arabidopsis (Liang et al., 1992), muangbean (Botella et al., 1992), rice (Zarembinski and Theologis, 1993), carnation (Henskens et al., 1994; Ten Have and Woltering, 1997), orchid ‘Phalaenopsis’ (Bui and O’Neill, 1998), melon (Ishiki et al., 2000) and rose (Ma et al., 2006). With respect to the ACS gene in Dendrobium hybrid cultivar Anna, one to two hybridization bands were observed after Southern hybridization analysis (Figure 5). These results raise the possibility that a multiple gene family may also encode for ACS in Dendrobium hybrid cultivar Anna.

Expression analysis of the DenACS gene
Figure 3. Comparison of the deduced amino acid sequence of DenACS from Dendrobium hybrid cultivar Anna with ACS from Dendrobium hybrid cultivar Karen (EF488014), Pompadour (EF488013), D. crumenatum (U64031), Cattleya spp. (AY504664) and Phalaenopsis spp. (AF004663). Amino acids residues identical and similar to each other are illustrated by asterisks and dots, respectively. Gaps introduced for alignment are indicated by a horizontal dash. The conserved active site regions are underlined and the active site lysine that bind to pyridoxal 5’-phosphate (PLP) and AdoMet is indicated by bold letter.
Figure 4. Comparison of the deduced amino acid sequences of the DenACS with ACS from petunia (*Petunia hybrida*) (AF049711), geranium (*Pelargonium hortorum*) (U88971), carnation (*Dianthus caryophyllus*) (X66605) and rose (*Rosa roxburghii*) (AY378152). Amino acids residues identical and similar between each other are shown by asterisks and dots, respectively. Gaps introduced for alignment are indicated by a horizontal dash. The conserved active site regions are underlined and the active site lysine that binds to pyridoxal 5’-phosphate (PLP) and AdoMet is indicated by bold letter.
Figure 5. Southern hybridization analysis of DenACS. Genomic DNA (20 μg) isolated from petal of the orchid flowers was digested with BamHI (B), EcoRI (E), HindIII (H) and PstI (P), electrophoresed on 0.7% agarose gel and transferred to nylon membrane. The membrane was hybridized with a 0.65-kb internal fragment of ACS gene.

Figure 6. RT-PCR analysis of the orchid DenACS expression in different stages of flowering. Total RNAs were prepared from orchid flowers at stages 1, 2 and 3 and subjected to RT-PCR analysis with primers specific for the DenACS as described in materials and methods. Actin gene was used as an internal control. The numbers above the lanes represent the number of PCR cycles. Ethidium bromide-stained rRNA was used as a control to normalize the amount of total RNA (10 μg).
the ACC oxidase activity. These data would be expected to be beneficial for future work such as the creation of the new cultivars that have strong resistance to senescence caused by ethylene through the modification of the DenACS or DenACO gene using the modern biotechnological techniques, such as antisense or RNAi technology.

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