Cloning and characterization of a β-amyrin synthase gene from the medicinal tree *Aralia elata* (Araliaceae)


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**ABSTRACT.** *Aralia elata* is an important medicinal plant in China; it produces large amounts of oleanane type triterpene saponins. A full-length cDNA encoding β-amyrin synthase (designated as *AeAS*) was isolated from young leaves of *A. elata* by reverse transcription-PCR. The full-length cDNA of *AeAS* was found to have a 2292-bp open reading frame, encoding a protein with 763 amino acid residues. The deduced amino acid sequence of *AeAS* showed the highest identity (97%) to *Panax ginseng* β-amyrin synthase. When *AeAS* cDNA was expressed in *Escherichia coli*, an 87.8-kDa recombinant protein was detected by SDS-PAGE and Western blotting. The sequence was also heterologously expressed in the yeast *Pichia pastoris*, and production of β-amyrin was detected by HPLC. Tissue expression pattern analysis by real-time reverse transcription-PCR revealed that *AeAS* is strongly expressed in leaves and stems, and weakly expressed in roots and flowers.

**Key words:** *Aralia elata;* β-amyrin synthase; *Pichia pastoris;* PNY1; Triterpenoid saponins; Real-time RT-PCR
INTRODUCTION

Aralia elata (Miq.) Seem. is an upright deciduous small tree or shrub belonging to the Araliaceae family. It is widely distributed in China, Far East Russian, Japan, and Korea. The young shoot of A. elata is a popular edible plant in China, Japan and Korea. The stem bark and root bark are used as traditional medicine for neurasthenia, rheumatism, hepatitis, diabetes, stomach spasm, and as a tonic. It has been reported that oleanane type triterpenoid saponins were the main active components (New Medical College of Jiangsu, 1977). Numerous new oleanane type triterpene saponins were isolated from the leaves (Kim et al., 2005), stem barks (Nhiem et al., 2011), root barks (Song et al., 2001), and buds (Li et al., 2006, 2007) of this plant. These saponins were reported to show anti-diabetic activity (Kim et al., 1993), cytoprotective effect on carbon tetrachloride-induced hepatic injury (Saito et al., 1993), ethanol absorption inhibitory effect (Yoshikawa et al., 1996a), hypoglycemic activity (Yoshikawa et al., 1995, 1996b), antioxidant activity and hypcholesterolemic and hypolipidemic effect (Chuang and Jung, 2003). Recently, many studies were focused on antulcer, anti-inflammatory and anticancer activity of these oleanane type triterpene saponins from A. elata (Zhang et al., 2006; Lee et al., 2009; Nhiem et al., 2011).

Oleanane type triterpenoid saponins are synthesized starting from the isoprenoid pathway. Two molecules of isopentenyl diphosphate (IPP) are condensed with one molecule of dimethylallyl diphosphate (DMAPP) by farnesyl diphosphate synthase (FPS), creating the molecule farnesyl diphosphate (FPP). Two FPP molecules are joined by squalene synthase (SS), originating squalene. Squalene is oxidized by squalene epoxidase (SE) to 2,3-oxidosqualene. 2,3-oxidosqualene is cyclized to produce β-amyrin by β-amyrin synthase (β-AS) belonging to the oxidosqualene cyclase (OSC) family. β-amyrin is thought to be a precursor for oleanolic acid, which is the aglycone of oleanane type triterpenoid saponins. β-amyrin is produced from oxidosqualene by product specific β-AS (monofunctional) or as one of the multifunctional OSCs (Yendo et al., 2010). Up to now, a number of product specific β-AS have been cloned and their function have been identified by heterologous expression in yeast: PNY1 (Kushiro et al., 1998a) and PNY2 (Kushiro et al., 1998b) from Panax ginseng, GgbAS1 from Glycyrrhiza glabra (Hayashi et al., 2001), PSY from Pisum sativum (Morita et al., 2000), AshASI (Sad1) from Avena strigosa (Haralampidis et al., 2001), MtAMY1 from Medicago truncatula (Iturbe-Ormaetxe et al., 2003), BPY from Betula platyphylla (Zhang et al., 2003), EtAS from Euphorbia tirucalli (Kajikawa et al., 2005), LjOSC1 from Lotus japonicas (Sawai et al., 2006), SvBS from Saponaria vaccaria (Meesapyodsuk et al., 2007), BgbAS from Bruguera gymnorrhiza (Basyuni et al., 2007), AsOXAl from Aster sedifolius (Cammareri et al., 2008), AtBAS from Arabidopsis thaliana (Shibuya et al., 2009), GsASI from Gentiana straminea (Liu et al., 2009), NsbAS1 from Nigella sativa (Scholz et al., 2009).

Although extraction, structure analysis and pharmacological tests of oleanane type triterpenoid saponins from A. elata have been extensively studied, the understanding of their biosynthesis is still limited. With the aim to enhance the production of oleanane type triterpenoid saponins and to understand their biosynthesis pathway in A. elata, an AeAS gene was first cloned and characterized. Furthermore, expression patterns of AeFPS, AeSS, AeSE, and AeAS genes involved in the oleanane type triterpenoid saponin synthesis pathway were also determined by real-time RT-PCR.

β-AS, is to our knowledge the key synthase connected with oleanane type triterpenoid
β-amyrin synthase gene of Aralia elata

saponin production. Therefore, we decided to isolate the gene encoding β-amyrin towards improving saponin yield. Here, we report on the first cloning and characterization of a β-AS gene from A. elata.

MATERIAL AND METHODS

Plant materials

A. elata was cultivated in the Division of Agriculture of Jilin University in Changchun, China. Young leaves, young stems, young later roots, and flowers were collected from 3-year-old plants in August. Harvested plant materials were frozen with liquid nitrogen and stored at -80°C.

Strains, plasmids and media

Escherichia coli strain DH5α and the pGM-T vector (Tiangen, Beijing, China) were used for cloning and vector construction. E. coli strain BL21 (DE3) and the pET28a vector (Novagen, Madison, WI, USA) were used for prokaryotic expression. Bacterial cells were cultured in Luria Bertani medium [LB, 1% (w/v) tryptone, 1% (w/v) NaCl, 0.5% (w/v) yeast extract, pH 7.5]. Pichia pastoris strain GS115 (His Mut+) and the pPIC9K vector (Invitrogen, Carlsbad, CA, USA) were used for eukaryotic expression. For P. pastoris cultivations, the following media were used at appropriate steps: yeast extract peptone dextrose medium (YPD, 1% (w/v) yeast extract, 2% (w/v) peptone, 2% (w/v) dextrose), yeast nitrogen base (YNB), minimal dextrose medium (MD, 1.34% YNB, 4 x 10⁻⁴% biotin, 2% dextrose), minimal medium (MM, 1.34% YNB, 4 x 10⁻⁴% biotin, 0.5% methanol), buffered complex glycerol medium (BMGY, 1% yeast extract, 2% peptone, 100 mM phosphate buffer, pH 6.0, 1.34% YNB, 4 x 10⁻⁴% biotin, 1% glycerol), and buffered complex methanol medium (BMMY, 1% yeast extract, 2% peptone, 100 mM phosphate buffer, pH 6.0, 1.34% YNB, 4 x 10⁻⁴% biotin, 1% glycerol, 0.5% methanol). All chemicals were from Sangon (Shanghai, China).

A. elata cDNA synthesis

Total RNA was extracted from A. elata young leaves using RNAiso Plus (Takara, Tokyo, Japan), according to manufacturer instructions. The quality and concentration of RNA were assessed by ethidium bromide-stained agarose 1% (w/v) gel electrophoresis and spectrophotometry. The RNA was converted to cDNA by providing 1 µg total RNA as the template in a 20 µL reaction using M-MLV reverse transcriptase (Takara) in the presence of 1 µL 0.5 µg/µL oligo(dT)₁₅ (Takara) and 2 µL 10 mM deoxyribonucleoside triphosphate (dNTP) (Takara) at 42°C for 1 h.

Cloning of AeAS cDNA

Based on the mRNA sequence of PNY1 (GenBank accession No. AB009030) from P. ginseng, AeAS-specific primers (forward, 5'-ATGTTGAAGCTTAGAGATACGG-3' and reverse, 5'-TTAGGTCCTAGGGACCGTAAT-3') were used in the amplification reaction.
Using cDNA from young leaves of *A. elata* as the template, a 35-cycle PCR using LA Taq (Takara) with proof reading activity was conducted. The cycling parameters were 94°C for 1 min, 55°C for 1 min, 72°C for 2.5 min and a final 10-min extension at 72°C using a PTC-200 Peltier Thermal Cycler (Bio-Rad, Hercules, CA, USA). The PCR product was recovered by electrophoresis on 1% (w/v) agarose gel, cloned into pGM-T easy vector and transformed into competent *E. coli* DH5α cells for sequencing (Sangon).

**Sequence and phylogenetic analysis**

Multiple sequence alignments were generated using the CLUSTAL W program (Thompson et al., 1994). Phylogenetic analysis of deduced amino acid alignments was performed using the neighbor-joining method with the program and TreeView software (Page, 1996). Bootstrap analysis with 1000 replicates was used to assess the strength of nodes in the tree. The GenBank accession Nos. of β-AS of other plants were as follows: BAA33461 (*P. ginseng* PNY1), BAA33722 (*P. ginseng* PNY2), BAA89815 (*G. glabra* GgbAS1), BAA97558 (*P. sativum* PSY), CAC84558 (*A. strigosa* AsbAS1), CAD23247 (*M. truncatula* MtAMY1), BAB83088 (*B. platyphylla* BPY), BAE43642 (*E. tirucalli* EtAS), BAE53429 (*L. japonicas* LjOSC1), ABK76265 (*S. vaccaria* SvBS or *Vaccaria hispanica* BS), BF80443 (*B. gymnorhiza* BgbAS), AAX14716 (*A. sedifolius* AsOXA1), BAG82628 (*A. thaliana* AtBAS), ACO24697 (*G. straminea* GsAS1), and ACH88048 (*N. sativa* NsbAS1).

**Construction of AeAS prokaryotic expression vector**

The entire ORF of *AeAS* cDNA was PCR amplified using forward primer (5'-GG GAATTCATATGATGGGAGCTTAAGATAGCGG-3') (*Nde*I restriction site underlined and translation start codon in bold), and reverse primer (5'-ACGCACGCTAGCTAGTAGTGGACGCTAA-3') harboring an *Sac*I site (underlined) and a stop codon (bold). The PCR product was gel purified and cloned into pGM-T easy vector for sequencing. After the sequence was confirmed to be correct, the target fragment was digested with *Nde*I and *Sac*I, gel purified, and ligated into the *Nde*I and *Sac*I sites in pET28a. The constructed plasmid pET28a-AeAS was transformed into *E. coli* stain BL21 (DE3) according to manufacturer instructions (Novagen). The empty pET28a vector was also transformed for a negative control.

**Heterologous expression of AeAS in E. coli**

*E. coli* BL21 (DE3) cells harboring recombinant plasmid pET28a-AeAS were grown in 10 mL LB medium containing 100 µg/mL kanamycin. Cultures were shaken at 200 rpm and held at 37°C to an absorbance of 0.7 to 1.0 at 600 nm before which isopropyl thio-β-D-galactoside (IPTG) was added to a final concentration of 0.1 mM. After 4 h of additional incubation, bacteria were collected by centrifugation and resuspended in 2X sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) loading buffer (80 mM Tris-HCl, pH 6.8, 100 mM dithiothreitol, 2% SDS, 0.006% bromophenol blue, and 15% glycerol). Then, samples were boiled for 10 min and separated at 12% SDS-PAGE. Proteins were visualized by staining in Coomassie brilliant blue R-250 (Sangon). For the Western blot procedure, proteins were transferred onto nitrocellulose membrane (0.45 µm; Invitrogen) using a MiniTrans blot system (Bio-
Rad). Proteins transferred to the membrane were blocked with 5% bovine serum albumin in 20 mM Tris-HCl, pH 7.5, 150 mM NaCl (TBS buffer) for at least 2 h. The membrane was twice washed with TBS buffer containing 0.05% Tween-20 (TBST buffer) for 10 min each, and then incubated for 2 h in TBS buffer and anti-His antibody (mouse monoclonal antibody; Tiangen) diluted 1:2000. The membrane was then washed three times with TBST buffer and exposed to horseradish peroxidase conjugated goat anti-mouse IgG antibody (Sigma, St. Louis, MO, USA) diluted 1:3000 for 2 h at 37°C. Finally, the membrane was washed three times with TBST buffer for 10 min each, and color development was performed with the TMB kit (Tiangen).

**Construction of AeAS eukaryotic expression vector**

Based on the recombination plasmid pET28a-AeAS, the fusion sequence of entire ORF of AeAS cDNA and His-Tag sequence at 5′ region was PCR amplified using forward primer (5′-GCTACGTACATCATCATCATCATCACAGCAG-3′) (SnaBI restriction site underlined and His-Tag in bold), and reverse primer (5′-ATTTGCGGCCGTCGTAAGTGTCCTAGAGACGTAAT-3′) harboring an NotI site (underlined) and a stop codon (bold). The PCR product was gel purified and cloned into pGM-T easy vector for sequencing. After the sequence was confirmed to be correct, the target product was digested with SnaBI and NotI, gel purified, and ligated into the SnaBI and NotI sites in pPIC9K (Invitrogen). The constructed plasmid pPIC9K-His$_6$-AeAS was then linearized with BglII and purified for transformation. The empty pPIC9K was also linearized for a negative control.

**Transformation of the P. pastoris strain**

Transformation of the P. pastoris strain GS115 (His Mut$^+$) was performed using the electroporation method according to pPIC9K vector manual (Invitrogen). Approximately 20 µL linearized plasmid DNA (about 10 µg) was used for electroporation in 0.2 cm cuvettes, using Gene Pulse Xcell Electroporation System (Bio-Rad) at 1.5 kV, 25 µF and 200 Ω for 5 s. Immediately after pulsing, 1 mL 1 M ice-cold sorbitol was added to the cuvette cells, and then 200 to 600 µL of the mixture was spread on MD plates at 30°C for screen of His$^+$ transformants. To screen for methanol utilization, each colony on MD plates was first spotted onto an MM plate and then onto a new MD plate. After 4 h, Mut$^-$ and Mut$^+$ colonies were identified. The clone screening was performed using PCR analysis with AeAS specific primer (5′-TACTATTGCCAGCATTGCTGC-3′) and 3′AOX1 primer (5′-GCAAATGGCATTCTGACATCC-3′).

**Heterologous expression of AeAS in P. pastoris**

The screened recombinant strain (His$^+$ Mut$^+$) integrated with pPIC9K-His$_6$-AeAS was cultured in 20 mL BMGY medium with a 100-mL flask at 30°C, until an OD$_{600}$ value of 2 to 6 was attained. Yeast cells were harvested by centrifugation at 3000 g at room temperature for 5 min and then resuspended in 100 mL BMMY medium with a 250-mL flask. Yeast cells were grown at 30°C for 4 days in a shaking incubator at 200 rpm. To induce expression of the recombinant protein, the methanol at a final concentration of 0.5% (v/v) was added to the culture every 24 h. Supernatants of the culture at 0, 24, 48, and 72 h were collected for the extraction
and analysis of the recombinant protein. The procedure of SDS-PAGE and Western blot was the same as described above.

**HPLC determination of β-amyrin in *P. pastoris***

After methanol induction for 3 days, the yeast cells were collected and baked until completely dry at 80°C. The dry sample (0.2 g) was extracted with 25 mL HCl/alcohol (2:8, v/v) solution at 95°C for 4 h and concentrated to dryness under reduced temperature. The dry material was dissolved with 30 mL sterile deionized H2O and extracted by ether for 4 times (15 mL each). The mixed extraction solution was concentrated to dryness at room temperature. The crude sample was dissolved with 10 mL methanol and filtered with a 0.45-µm filter HPLC analysis. The HPLC equipment was carried out on a Shimadzu LC-20A System (Shimadzu, Kyoto, Japan) including two LC-20AT automatic rinsing pumps, an SPD-M20A UV-VIS detector, a manual sample injector 7725 valve with a 20-µL loop, and a DGU-20A, degasser. All separations were performed using a C18 column (250 x 4.6 mm, 5 µm particle size) from Phenomenex (Torrance, CA, USA). The mobile phase was methanol/water (98:2, v/v). The injection volume was 10 µL, the effluent was monitored at 254 nm, the flow rate was 1.0 mL/min and the column temperature was 40°C. A standard calibration curve assembled from a range of concentrations (25, 125, 250, 500, 1000 µg/mL) of β-amyrin (Sigma) was used as standard for HPLC analysis. Data acquisition and processing were accomplished with the Shimadzu LC solution software.

**Expression of *AeAS*, *AeFPS*, *AeSS*, and *AeSE* in *A. elata***

Total RNA extraction and cDNA synthesis from young leaves, young stems, young later roots, and flowers of *A. elata* were the same as described above. Based on the mRNA sequence of *AeFPS* (HM219226), *AeSS* (GU354313), *AeSE* (GU354314), *AeAS* (HM219225), and Panax notoginseng glyceraldehyde-3-phosphate dehydrogenase (GAPDH, DQ186631) genes in GenBank, we designed specific primers for quantitative real-time RT-PCR as follows: primers for *AeFPS* (product length of 260 bp, forward primer: 5ꞌ-CTTATTGAAATGGGAACCTA-3ꞌ, reward primer: 5ꞌ-TCATACTCGGCAAATACATC-3ꞌ), primers for *AeSS* (product length of 274 bp, forward primer: 5ꞌ-AGTCTTCAGAGGGGTAGTG-3ꞌ, reward primer: 5ꞌ-GAAGATAATAGCAATCAGGGC-3ꞌ), for *AeSE* (product length of 258 bp, forward primer: 5ꞌ-GGAAGAGAGAGAAAACTAAGG-3ꞌ, reward primer: 5ꞌ-CATTCACACAATCTCCAG-3ꞌ), for *AeAS* (product length of 197 bp, forward primer: 5ꞌ-TTCCTCAGGAGGCTAGGTG-3ꞌ, reward primer: 5ꞌ-GAACACTGTATCAAGATGCC-3ꞌ), for *AeGAPDH* (product length of 133 bp, forward primer: 5ꞌ-TTCCTCAGGAGGCTAGGTG-3ꞌ, reward primer: 5ꞌ-GAACACTGTATCAAGATGCC-3ꞌ). RT-PCR was carried out in a reaction mixture of 20 µL, containing 2 µL 1:5 (v/v) dilution of synthesized cDNA, primers to a final concentration of 0.4 µM each, 8 µL 2.5X RealMasterMix, 1 µL 20X SYBR solution (Takara). The standard cycling conditions were 95°C 30 s, followed by 40 cycles of 95°C for 5 s, 60°C 30 s, 72°C 30 s with the ABI 7500 system (Applied Biosystems Inc., Foster City, CA, USA). The data were analyzed with the 7500 Fast System SDS version 1.3.1 software (Applied Biosystems). All reactions were performed in triplicate, and relative gene-expression determinations were made with the comparative delta-delta C_t method (2^ΔΔCT; Livak and Schmittgen, 2001) by using the housekeeping GAPDH gene as the internal control.
RESULTS AND DISCUSSION

Cloning of a cDNA encoding AeAS

In order to isolate an AeAS gene, specific primers were designed from the sequence of *P. ginseng* PNY1 (Kushiro et al., 1998a). Both *A. elata* and *P. ginseng* belong to the Araliaceae family and they are highly homologous. Finally, a full-length cDNA encoding for β-AS was isolated from *A. elata* young leaves. The cDNA contained an ORF of 2292 bp and was deduced to code for a 763 amino acid protein (Figure 1). The AeAS amino acid sequence showed the high identity of 97% to PNY1 and 87% to PNY2, as would be expected, since all species belong to the Araliaceae. So we may design species-specific primers to clone unknown genes from known genes of highly homologous plants, which are the same family without degenerate primers PCR, 5'-rapid amplification of cDNA ends (RACE), and 3'-RACE. Based on this strategy, we cloned *AeFPS* (HM219226), *AeSS* (GU354313), *AeSE* (GU354314) successfully. AeAS possesses the amino acid motif DCTAE, thought to form part of the active site of OSCs (Abe et al., 1993; Abe and Prestwich 1995) and the four QW (QXXXGXW) motifs, which are thought to be involved in the stabilization of carbocationic intermediates formed during the cyclization of OSCs (Poralla et al., 1994). In addition, AeAS amino acid sequence contains the Trp residue in the MWCYCR motif that plays an important role in the formation of β-amyrin in *P. ginseng* (Kushiro et al., 2000). The second Lys in the FIKKSQ motif of AeAS is strictly conserved in monofunctional β-AS and may favor E-ring expansion to produce oleanane or ursane type triterpene saponins (Basyuni et al., 2007).

The DCTAE motif is underlined, the MWCYCR motif is double underlined, the FIKKSQ motif is underlined with a discontinued line, and the QW motifs are boxed.

Sequence and phylogenetic analysis of AeAS amino acid sequence

Alignment results showed that AeAS amino acid sequence had identity of 97% to PSN1, 87% to PSN2, 84% to BpY, 84% to BgBAS, 83% to EtAS, 82% to GgBAS1, 82% to LjOSc1, 81% to MtAMy1, 81% to PsY, 79% to SvBS, 79% to AsOXa1, 76% to AtBAS, 76% to GsAS1, 72% to NsBAS1, and 49% to AsbAS1. Fifteen dicotyledonous monofunctional β-ASs showed high similarities (72 to 97%) to each other and displayed lower similarities (48%) to AsbAS1 from the monocotyledonous plant *A. strigosa*. A phylogenetic tree was constructed from the deduced amino acid sequences of β-ASs (Figure 2). In the tree, AsbAS1 is clearly distinct from other β-ASs of dicotyledonous plants. The phylogenetic tree shows that AeAS, PNY1, together with PNY2, form one branch of the β-AS cluster. MtAMy1, PsY, LjOSc1, and GgBAS1 form the other branch, because all species belong to the Leguminosae family. Figure 3 shows that crucial motifs of β-AS are well conserved in these dicot plants, except that Gly257 instead of Trp257 in MWCYCR motif of GsAS1, and Ala448 instead of Lys448, Asn448 instead of Lys448, Asn450 instead of Lys450, and Gln452 instead of Lys452 in FIKKSQ motif of AsOXa1, GsAS1, AtBAS, and NsBAS1, respectively. Although their crucial residues are changed from other plants, they are also product specific β-AS by function establishment in *P. pastoris* (Cammareani et al., 2008; Shibuya et al., 2009; Liu et al., 2009; Scholz et al., 2009). Therefore, these observations suggested that the presence of additional protein domains act to control the reaction product of β-ASs. From the high level of amino acid sequence identity and comparison of critical motifs with β-AS of other plants, the *AeAS* cDNA may code for the product specific β-AS.
Figure 1. Nucleotide and deduced amino acid sequences of AeAS from *Aralia elata* (GenBank accession Nos. HM219225 and ADK12003). The DCTAE motif is underlined, the MWCYCR motif is double underlined, the FIKKSQ motif is underlined with a discontinued line, and the QW motifs are boxed.
The β-amyrin synthase gene of *Aralia elata* was investigated further. The full-length cDNA of *AeAS* was heterologously expressed in *E. coli* and *P. pastoris*, and product from yeast expressing the *AeAS* gene induced by methanol was analyzed by HPLC. SDS-PAGE analysis identified a protein of molecular mass about 88 kDa expressed by the *E. coli* strain BL21 when induced by IPTG (Figure 4A). Western blot of this separation revealed that this product was a His-Tag fusion protein (Figure 4B). Similarly, SDS-PAGE analysis identified a recombination protein of molecular mass about 88 kDa expressed by the *P. pastoris* GS115 when induced by methanol (Figure 5A).
recombination protein was also detected by Western blot (Figure 5B). The results showed that AeAS is successfully expressed in *E. coli* and *P. pastoris*. The product specificity of AeAS was tested by expression of the *AeAS* cDNA in yeast induced by methanol. HPLC analysis indicated that the product was not present in extracts of negative control cells carrying the empty pPIC9K vector, and the yeast cells expressing AeAS accumulated a product with a retention time of 11.7 min that was identical to that of standard β-amyrin (Figure 6). The results showed that the *AeAS* cDNA codes for a product specific β-AS, similar to other plants monofunctional β-AS. Furthermore, yeast appears to be an excellent model system not only for the functional characterization of product-specific β-AS, but also for the production of target products such as β-amyrin and other oleanane type triterpene saponins (Phillips et al., 2006; Lodeiro et al., 2007).

**Expression of *AeAS* and other genes involving in triterpene saponin synthesis in *A. elata***

*AeFPS, AeSS, AeSE,* and *AeAS* are triterpene saponin pathway genes in *A. elata*. The melting curve of the reference gene *AeGAPDH* showed a single peak, indicating that a single PCR product was present. The Ct value of the reference gene *AeGAPDH* in different organs of *A. elata* was relatively stable, indicating that it can be used as internal control (data not shown). Therefore, we used the average Ct values of *AeGAPDH* as internal standard to nor-
β-amyrin synthase gene of Aralia elata

malize the quantification of mRNA expression. Quantitative PCR analysis showed the relative expression of \( \text{AeFPS} \), \( \text{AeSS} \), \( \text{AeSE} \), and \( \text{AeAS} \) in stems, leaves, later roots, and flowers of \( A. \) elata (Figure 7). All genes could be detected in all tissues, but at different levels. The expression of \( \text{AeFPS} \) was high in stems, moderate in leaves and flowers, and very low in later roots. The expression of \( \text{AeSS} \) was high in stems, moderate in leaves and flowers, and very low in later roots. The expression of \( \text{AeSE} \) was high in stems and leaves, and very low in later roots and flowers. The expression of \( \text{AeAS} \) was high in stems and leaves, moderate in flowers, and very low in later roots. We found that the expression of all genes is very low in later roots and the result is apparently conflicting with biochemical results that showed a high content of triterpenoid saponins in \( A. \) elata roots (Song et al., 2001). A possible explanation of the contradictory results is that saponins are synthesized in green parts of the plant, and then transported to the roots just like \( \text{Calendula officinalis} \) (Hostettmann and Marston, 1995).

The housekeeping gene \( \text{AeGAPDH} \) was used to normalize the amount of sample cDNA added to the reaction. Bars indicated standard deviation from triplicate amplification.

In conclusion, we successfully cloned and characterized the \( \text{β-AS} \) gene associated with oleanolic acid biosynthesis in \( A. \) elata (Miq.) Seem. for the first time.

**Figure 5.** SDS-PAGE analysis and Western blot of \( \text{AeAS} \) expressed in \( \text{Pichia pastoris} \). A. SDS-PAGE; B. Western blot. \( \text{Lane M} = \) Protein molecular weight marker (low); \( \text{lane 1} = \) empty pPIC9K vector; \( \text{lane 2} = \) pPIC9K containing \( \text{AeAS} \) without methanol; \( \text{lane 3} = \) pPIC9K containing \( \text{AeAS} \) induced by methanol for 24 h; \( \text{lane 4} = \) pPIC9K containing \( \text{AeAS} \) induced by methanol for 48 h; \( \text{lane 5} = \) pPIC9K containing \( \text{AeAS} \) induced by methanol for 72 h.
Figure 6. HPLC analysis of yeast cell extracts. A. Empty pPIC9K vector. B. pPIC9K containing AeAS. C. Standard β-amyrin. The retention time of standard β-amyrin is 11.7 min.

Figure 7. Relative quantification of expression levels for *AeFPS* (FPS), *AeSS* (SS), *AeSE* (SE), and *AeAS* (AS) genes in stems, leaves, later roots, and flowers of *Aralia elata*, respectively.
β-amyrin synthase gene of *Aralia elata*

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