Rose Phytoene Desaturase Gene Silencing by Apple Latent Spherical Virus Vectors

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Abstract. We have succeeded in establishing a virus-induced gene silencing (VIGS) of rose using Apple latent spherical virus (ALSV) vectors. An ALSV infection on rose did not cause any symptoms like those observed on other plant species and grew healthy. We have cloned and sequenced the gene responsible for the PDS phenotype, which is highly uniform photo-bleached phenotype with PDS inhibitions were observed on the upper leaves of primary shoots and on a secondary shoot of R. rugosa for more than 5 months. ALSV vectors seemed useful for analyzing gene function and for the molecular breeding of rose.

Roses are one of the economically most important ornamental plants, and modern commercial roses consisting of several thousand cultivated varieties are found widespread around the world. Because consumers always expect new varieties with high ornamental value such as flowers of a new shape and color, breeders and breeding companies are continuously producing new varieties by cross-hybridization and/or mutation breeding techniques. However, because only eight to 10 species among more than 150 have contributed to modern commercial roses (Matsumoto et al., 1998), it has become difficult to produce new epoch-making varieties because their genetic background is limited.

A genetically modified blue rose was introduced into the market in 2011 (Kato et al., 2011). Because roses have no blue varieties because of the absence of the flavonoid 3’,5’-hydroxylase (F3’,5’H) gene, which is essential to synthesizing the delphinidin-based anthocyanins, a transgenic blue rose has been produced by the expression of an introduced pansy F3’5’H gene (Kato et al., 2008). Besides the F3’5’H gene producing the blue rose, the introduction of an Ace-AMP1 and basic (Class 1) chitinase gene caused resistance to powdery mildew and blackspot, respectively, and the rooting ability of rose was enhanced by the introduction of a ROL gene from Agrobacterium rhizogenes (Marchant et al., 1998a, 1998b; van der Salm et al., 1997). However, such traits proved unattractive to consumers as did other genetically modified roses introducing the reporter or marker genes such as the β-glucuronidase gene, luciferase gene, neomycin phosphotransferase gene, and green fluorescent protein gene (Firoozabady et al., 1994; Kim et al., 2004; Li et al., 2002, 2003; Vergne et al., 2010). These agrobacterium-mediated genetic or biolistic transformation systems require much effort as a result of their low efficiency of transformation and also need a long timespan for making and keeping the somatic embryos and the generation of transgenic plants. Moreover, because many rose varieties are known to be recalcitrant to regeneration and hence to genetic transformation, somatic embryos of limited rose varieties such as ‘Royalty’, ‘Carefree Beauty’, ‘Tineke’, ‘Glad Tidings’, ‘Moneyway’ and ‘Old Blush’ were used as sources for genetic transformation (Firoozabady et al., 1994; Kim et al., 2004; Li et al., 2002, 2003; Marchant et al., 1998a, 1998b; van der Salm et al., 1997; Vergne et al., 2010). Because all the varieties used (except for ‘Old Blush’) are tetraploid, they are not useful for studying the gene function in rose for complicated dose effects among alleles.

Recently, although we have established a plantlet regeneration system of the diploid wild rose, Rosa rugosa, we could not succeed in its genetic transformation, probably as a result of its low transformation frequency (Matsumoto, 2009). R. rugosa distributed throughout northern Japan has contributed to the development of some modern commercial roses, including hybrid rugosas, hybrid kordesii, and shrub roses (Cairns, 2000). This species possesses unique and valued traits, including large fragrant flowers, a long flowering period, disease resistance, and winter-hardiness (Svejda, 1974, 1977). It is one of the few recurrently flowering species, which has been critically important to establishing its widespread use in the ornamental trade.

VIGS is a powerful tool for the fast and efficient functional analysis of genes in plants (Burch-Smith et al., 2004; Purkayastha and Dasgupta, 2009; Robertson, 2004). Because VIGS circumvents the plant transformation system, it might be possible to use it for rose varieties that are hard to transform. An ALSV of ≈25-nm diameter particles belonging to the genus Cheravirus contains two single-stranded RNA species (RNA 1 and RNA 2) and three capsid proteins (VP25, VP20, and VP24) (Li et al., 2000). It consists of two components, M and B, containing two molecules of RNA 2 and a single molecule of RNA 1, respectively (Li et al., 2000). RNA 1 and RNA 2 of 6813 nt and 3385 nt length, respectively, excluding the 3’ poly(A) tail, have a single open reading frame encoding to a replication-associated protein of 243 K and a 42 KDA movement protein on the N-terminal side and three capsid proteins on the C-terminal side, respectively (Li et al., 2000; Yoshikawa et al., 2006). Recently, a VIGS vector based on ALSV has been developed that succeeded in silencing endogenous genes such as PDS, magnesium chelatase, a ribulose-1,5-bisphosphate carboxylase small subunit, an alpha subunit of chloroplast chaperonin, an elongation factor 1, a terminal flower 1, and an iso flavone synthase in a broad range of plants including tobacco, tomato, Arabidopsis, cucurbits, legumes, apple, pear, and Japanese pear (Igarashi et al., 2009; Sasaki et al., 2010). Moreover, the ALSV-based vector expressing a FLOWERING LOCUS T gene has induced rapid flowering in apple, suggesting that the vector might enter and replicate in the meristematic tissue of apple trees (Yamagishi et al., 2011).

In this study, as a first step for developing a functional analysis of genes in rose and a new breeding system in rose, we investigated the availability of ALSV vectors in rose.

Materials and Methods

Plant materials. Wild roses (Rosa rugosa Thunb. ex Murray, R. multiflora Thunb. ex Murray, and R. bracteata Wendl.) were grown in a field at the Experimental Farm of the Graduate School of Bioagricultural Sciences, Nagoya University, Japan. Young leaves were collected, frozen dry at –80 °C, and stored until use. Their seeds were stored at 4 °C, and germinated seedlings with one to three true leaves were used for further experiments.

Construction of ALSV vectors and viral inoculation. Total RNAs of R. multiflora leaves were extracted essentially as described by Chang et al. (1993), and reverse transcription was performed with Oligo (dT) using the RNA PCR™ kit (AMV) Version 3.0 (TAKARA BIO Inc., Shiga, Japan) according to the manufacturer’s instructions. To clone the rose PDS gene, we used Rosa PDS-F (5’-TGCA...
GATGCAGGT(CATAAA-3') and tPDS (5'-CTTCCATTTTCTGTCAACC-3') as forward and reverse primers from alignment sequences of the PDS genes in R. × hybridra (F1B35397.1), Nicotiana tabacum (AF161742.1), and Fragaria × ananassa (FJ795342.1). A fragment of ≈900 bp corresponding to the R. multiflora ortholog of PDS (mulPDS) was amplified by polymerase chain reaction using the primers as mentioned, and PrimeSTAR® GXL DNA Polymerase (TAKARA BIO Inc., Shiga, Japan) under the following conditions: 35 cycles of 10 s at 98 °C, 15 s at 55 °C, 1 min at 72 °C, and a final extension of 7 min at 72 °C. The fragment was cloned using the DNA ligation Kit <Mighty Mix> and/or the Mighty TA-cloning Kit (TAKARA BIO Inc., Shiga, Japan). The sequences of the positive clones were then determined by dye deoxy chain termination on an Applied Biosystems 3130 Genetic analyzer (Life Technologies Japan Co., Ltd.) using a Big Dye Terminator Version 3.1 Cycle Sequencing Kit (Life Technologies Japan Co., Ltd.).

A 162-bp fragment corresponding to the nucleotide positions 189–351 of the mulPDS gene was amplified from R. multiflora cDNAs obtained by RT with Oligo (dT) by PCR using the Xho I-RmPDS-F (5'-GAATTCAGGCGA AACAAGCCAGGAGATTT-3') and Xho I-RmPDS-R (5'-AAGGATCCAGCTGTC ACCAAGAATG-3'), including the BamHI site (underlined) as forward and reverse primers, respectively, for making the mulPDS-ALSV vector. The PCR was carried out with KOD-Plus-Ver.2 DNA polymerase (TOYOCO Co., Ltd., Osaka, Japan) under the following conditions: 2 min at 94 °C, 30 cycles of 10 s at 98 °C, 30 s at 54 °C, 30 s at 68 °C, and a final extension of 7 min at 72 °C. The fragment corresponding to the 162 bp was cloned and sequenced as mentioned, and the 162-bp fragment was obtained by Xho I and BamHI double digestion at 37 °C for 2 h and then ligated to the BamHI-Xho I sites of the pEALSR2-based vector (Li et al., 2004).

We have named the vector pEALSR2L5-R5-mulPDS, *Chromopodium quinoa* plants were wounded and inoculated with combinations of two cDNA infectious clones comprising the viral genome (pEALSR1 and pEALSR2L5-R5, pEALSR1 and pEALSR2L5-R5-mulPDS) using a mechanical method (Li et al., 2004). After 2 to 3 weeks, leaves with symptoms of viral infection were crushed in three volumes of extraction buffer (0.1 M Tris-HCl, pH 7.8, 1 mM EDTA) and then the crude extract containing viral RNA was reinoculated to the *C. quinoa* plants. Total RNAs were purified from infected *C. quinoa* leaves using a TRizol reagent (Life Technologies Japan Ltd., Tokyo, Japan) and then adsorbed onto gold particles as follows. Fifteen milligrams of gold particles (1.0 μm in diameter; Bio-Rad Laboratory, Inc., CA) was sonicated in RNase-free distilled water (50 μL) for at least 1 min using the ULTRASONIC CLEANER VS-F100 (As One Co., Ltd., Osaka, Japan). After 5 min shaking of gold particles with 150 μL of total RNA extracted from infected leaves (150 μg), 45 μL of 5 M ammonium acetate was added and shaken for 5 min and then 990 μL of 2-propanol was added and shaken for 5 min. The mixture was placed at −20 °C for at least 1 h. Gold particles coated with total RNAs were washed three times with 1.0 mL of ethanol (95.5%) and resuspended in an appropriate amount (≈1.8 mL) of ethanol (95.5%). The suspension was used for preparing the gold-coat tubing according to the manufacturer’s instructions. The tube coated with gold particles was cut into 30 cartridges, each containing 5 μg of total RNAs from infected leaves.

The rose seedlings were wounded and inoculated with combinations of gold particles coated with total RNAs at 1650 psi pressure using the PDS-1000/HeTM Particle Delivery System (Bio-Rad Laboratory, Inc.) or 80–100 psi pressure using the Helios Gene Gun System (Bio-Rad Laboratory, Inc.). After particle bombardment, the plants inoculated wtALSV or ALSV vectors were grown in a growth chamber BIOTRON (Nippon Medical and Chemical Instruments Co. Ltd., Tokyo, Japan) and set at 23 °C under long-day conditions (16 h light, 8 h dark).

**RNA analyses.** Total RNAs of the newly generated upper leaves after inoculation of wtALSV or ALSV vector on rose seedlings were extracted using the kit RNA-suisui-R (Bio-Rad Laboratory, Inc.) or 80–100 psi pressure using the Helios Gene Gun System (Bio-Rad Laboratory, Inc.). After particle bombardment, the plants inoculated wtALSV or ALSV vectors were grown in a growth chamber BIOTRON (Nippon Medical and Chemical Instruments Co. Ltd., Tokyo, Japan) and then set at 23 °C under long-day conditions (16 h light, 8 h dark).

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symptoms on either inoculate or upper leaves of *R. rugosa*, and they grew healthy as observed in ALSV-infected plants of the family *Amaranthaceae*, *Brassicaceae*, *Leguminosae*, *Scrophulariaceae*, and *Solanaeae* (Igarashi et al., 2009). Based on those results, we thought that ALSV could infect rose without symptoms.

**Cloning of the PDS gene from rose and PDS fragment for silencing.** Because PDS is one of the widely used VIGS markers, we tried to silence the PDS gene in rose. We have cloned and determined a 838-bp cDNA sequence of the PDS gene in *R. multiflora* (deposited under the accession number AB705444), because only 491 bp of that in *R. × hybrida* was available on the database. We compared the cDNA sequence with that of *R. × hybrida* at the corresponding region and clarified a 97.8% and a 98.0% similarity at nucleotide and a deduced amino acid level, respectively. We searched the PDS gene in *R. multiflora* and *R. × hybrida* for inducing the PDS silencing for different wild roses. Because the ALSV vectors carrying 200 bp PDS sequences induced the stable VIGS efficiently in *Nicotiana* species (Igarashi et al., 2009), we have selected the 162 bp (54 deduced amino acids) cDNA fragment having identical sequences.

**Virus-induced silencing of PDS gene in the seedlings of rose.** The 162-bp fragment of the PDS gene in rose was inserted into an ALSV-RNA2-based vector (Fig. 2). The resulting virus was designated as pEALSRL5R5-mulPDS (mulPDS-ALSV), and total RNAs isolated from mulPDS-ALSV-infected *C. quinoa* leaves were inoculated to the 71 *R. rugosa* and 48 *R. bracteata* seedlings because the stage of the second true leaf appeared by particle bombardments. We observed a photo-bleached phenotype in four independent seedlings of *R. rugosa*, and the ratio of mulPDS-ALSV infection was 5.6% (Table 1). The first one showed the photo-bleached phenotype as a fourth true leaf, and the ratio of mulPDS-ALSV infection was 5.6%.

**Silencing of the PDS gene in *Rosa rugosa* primary shoot.** (A) *R. rugosa* seedlings used for infection. (B-C) *R. rugosa* seedling infected with pEALSRL5R5-mulPDS (mulPDS-ALSV). The photo-bleached phenotype was observed on newly generated fourth leaves 16 d after inoculation. (D) Detection of a polymerase chain reaction fragment corresponding to 464 bp derived from pEALSRL5R5-mulPDS. GL3 = third true leaf (green); WL4 = fourth true leaf (white); GL = green leaf from secondary shoot; WL = photo-bleached leaf from secondary shoot.

**Expression analyses of PDS gene in *Rosa rugosa* primary and secondary shoots.** Semiquantitative reverse transcription–polymerase chain reaction (PCR) analyses of phytoene desaturase and β-actin gene in the leaves of *R. rugosa* primary (A) and secondary shoots (B). 29–31 represents a PCR cycle number. C = control (no inoculation); WT, wtALSV inoculation; GL3, third true leaf (green) from primary shoot; WL4 = fourth true leaf (white) from primary shoot; WL = sixth true leaf (white) from primary shoot; GL = green leaf from secondary shoot; WL = photo-bleached leaf from secondary shoot.

**Green fluorescent protein (GFP-ALSV) infection on transgenic tobacco expressing GFP (GFP-tobacco) at the third to fifth true leaves of the sixth true leaf stage induced VIGS at the 10th to 12th leaves (Yaegashi et al., 2007).** Based on those results, the amount of mulPDS-ALSV seemed to be insufficient for VIGS at the stage of third green leaf generation. The
second and third seedlings also showed the photo-bleached phenotypes at the fifth true compound leaf after 14 and 16 d from inoculation, respectively (data not shown). These seedlings also stopped growing, probably as a result of a scarcity of nutrients caused by insufficient photosynthesis. One died at the fifth stage of true leaf expansion, but the photo-bleached leaves of the other one turned green, then started re-growing after 1 month, and newly generated leaves also showed a photo-bleached phenotype (data not shown). The fourth seedling exhibited the most severe phenotype (Fig. 4). Although the primary shoot showed no photo-bleached phenotype, the secondary shoot derived from the base of the second true leaf showed a completely photo-bleached phenotype. The secondary shoot began growing at the stage of the eighth true leaf’s appearance on a primary shoot, whereas the photo-bleached phenotype continued growing for more than 5 months. We have detected a PCR fragment corresponding to the expected length of 464 bp using primers specific for mulPDS-ALSV RNA2 from the white leaves of a second shoot but not detected from the green leaves of a primary shoot (Fig. 5). The expression levels of the PDS gene were distinctly reduced at the white leaves of the secondary shoot compared with those in the green leaves of the primary shoot (Fig. 4B). Thus, we speculated that an infection of mulPDS-ALSV occurred in the axially bud existing at the base of the second true leaf. Actually, ALSV propagated at the host cells moved to a secondary shoot; MP = pEALSR2LR5-mulPDS. GL = green leaf from primary shoot; WL = photo-bleached leaf from secondary shoot; MP = pEALSR2LR5-mulPDS (mulPDS-ALSV) plasmid DNA.

In the present study, we demonstrated that the ALSV vector could act as an inducer for the silencing of the endogenous gene in rose. One advantage of ALSV-mediated VIGS in rose is that ALSV infects rose without showing any symptoms. This may be preferable depending on the phenotype of the target gene. On the other hand, at present it is necessary to bombard a large amount of rose seedlings as a result of the low efficiency of ALSV infection. Although we have not succeeded in inoculating onto cotyledons of germinated rose seeds, it might be possible to develop a particle bombardment procedure optimized for those tissues by changing the bombardment pressure or gold particle size. Also, because susceptibility to virus could be different between species or genotypes, it is worth testing whether other genus Rosa and/or genotypes of R. rugosa show improved infection efficiency. Nonetheless, our VIGS-inducing system in rose reported here might be a powerful tool for functional validation of genes associated with horticulturally important traits such as flower morphology, recurrent flowering, and density or number of thorns.

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