Application of cucumber mosaic virus to efficient induction and long-term maintenance of virus-induced gene silencing in spinach

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Abstract Virus-induced gene silencing (VIGS) is a useful tool for functional genomics in plants. In this study, we tried to apply cucumber mosaic virus (CMV) to efficient induction of VIGS in spinach. Although VIGS for spinach had been previously developed based on two viruses (beet curly top virus and tobacco rattle virus), they still have some problems with systemic movement and long-term maintenance of VIGS in spinach. Although ordinary CMV strains infect spinach inducing distinct mosaic symptoms, using a CMV pseudorecombinant, we can modify the viral pathogenicity to attenuate viral symptoms that may mask the silencing phenotype. We here successfully demonstrated the viral ability to silence the *phytoene desaturase* (*PDS*) and the *dihydroflavonol 4-reductase* (*DFR*) genes in spinach. Because CMV could quickly induce VIGS even at 7–10 days postinoculation and the virus did not disappear even at the flowering stage, this CMV-based VIGS system would contribute to functional genomics in spinach and especially to the elucidation of molecular mechanisms for some properties unique to spinach such as plasticity of sex expression; the CMV-induced VIGS can last until the flowering stage after the virus was inoculated onto the seedling.

Key words: cucumber mosaic virus, spinach, virus-induced gene silencing.

Spinach (*Spinacia oleracea*) is a member of *Chenopodioideae*, a subfamily of the family *Amaranthaceae*, and is one of the most nutritious vegetables, rich in vitamins and minerals, being grown in over 50 countries (Fuentes-Bazan et al. 2012). It is commonly considered as dioecious species, although certain cultivars and genotypes can produce individuals with both staminate and pistillate flowers (i.e., monoecious plants) (Janick and Stevenson 1955b; Onodera et al. 2008). The dioecism and monoecism in spinach are utilized for producing commercial hybrid seeds, and hence the elucidation of the mechanisms for controlling sex expression is important for spinach breeding (Janick 1998; Onodera et al. 2011).

Spinach has long served as a model to study plant sex determination and expression as well as flavonoid biosynthesis and chloroplast function (Beerhues and Wiermann 1988; Beerhues et al. 1988; Chailakhyan and Timiriazev 1979; Ellis 1981; Janick and Stevenson 1954, 1955a, 1955b, 1955c; Sherry et al. 1993; Shimada et al. 2004; Yamamoto et al. 2014). The draft genome assemblies and the transcriptome data sets of spinach have been recently released (Okazaki et al. 2019; Xu et al. 2015, 2017), which are useful to identify loci responsible for important agronomic traits (e.g., loci for sex determination, disease resistance, etc.), enhancing the value of spinach as a model plant more and more. As the information on the genomic organization accumulate, functional analysis of spinach genes must be accelerated. However, rapid and efficient tools for studies of functional genomics are still not well established in spinach; although spinach transformation methods have been developed, they are based on callus-mediated regeneration systems, requiring much time and labor to obtain transformants (Chin et al. 2009; Nguyen et al. 2013).

Virus-induced gene silencing (VIGS) is useful for functional genomics especially in plants to which conventional transgenic methods cannot be easily applicable. For spinach, VIGS systems have been developed based on two viruses so far; one is beet curly top virus (BCTV) (Golenberg et al. 2009; Sather et al. 2010) and the other is tobacco rattle virus (TRV) (Lee et al. 2017). Although both BCTV- and TRV-based

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Abbreviations: BCTV, beet curly top virus; CMV, cucumber mosaic virus; CMV-L, CMV legume strain; CMV-Y, CMV Y strain; *DFR*, *dihydroflavonol 4-reductase*; dpi, days postinoculation; *GAPDH*, *glyceraldehyde 3-phophate dehydrogenase*; *PDS*, *phytoene desaturase*; RT-PCR, reverse transcription polymerase chain reaction; SE, standard error; TRV, tobacco rattle virus; VIGS, virus-induced gene silencing.

VIGS systems have been demonstrated to be effective in silencing some spinach genes, there is still room for improvement of VIGS in spinach. In the BCTV-based silencing, the virus vector with an insert was localized only in the inoculated leaves and could not systemically spread in the entire plant when the *ribulose bisphosphate* carboxylase small subunit, transketolase and homeotic transcription factor genes were targeted (Golenberg et al. 2009); the silenced phenotypes were observed at 6-8 weeks postinoculation (wpi). In the TRV-based silencing, when the phytoene desaturase (PDS) gene was silenced, the silencing phenotype appeared at 3 wpi but eventually disappeared after 4 wpi due to the reduction in viral accumulation in the systemically infected leaves (Lee et al. 2017). It is obvious that viral systemic infection is very important to induce efficient VIGS as demonstrated in some legume species and lily plants (Pflieger et al. 2013; Tasaki et al. 2016). Toward developing a more practical VIGS vector for spinach, we here examined the ability of the cucumber mosaic virus (CMV) vector to systemically spread in infected spinach and to sustain its systemic infection until the reproductive stage of plant. Results presented in this study demonstrate that our CMV vector targeting either the PDS gene or the dihydroflavonol 4-reductase (DFR) gene could induce efficient VIGS in spinach at most 7-10 days postinoculation (dpi) and existed in the infected spinach even after 2 months postinoculation.

CMV has been developed as a useful VIGS vector for many plants including Nicotiana benthamiana, tomato, pepper, Arabidopsis thaliana, lily, soybean, banana and maize (Hong et al. 2012; Liu et al. 2010; Otagaki et al. 2006; Tzean et al. 2019; Wang et al. 2016). Recently, it has been demonstrated that we could silence even a gene of mycorrhizal fungi that colonized the root of N. benthamiana (Kikuchi et al. 2016). In this study, we used two CMV strains: CMV Y strain (CMV-Y) and CMV legume strain (CMV-L). CMV has tripartite positivesense RNA genomes (RNA1, RNA2 and RNA3); three genomic RNAs of CMV-Y and CMV-L were named Y1-Y3 and L1-L3, respectively. Full-length cDNAs of all the genomic RNAs of both strains have been cloned into the pUC plasmids (Otagaki et al. 2006; Suzuki et al. 1991). When in vitro transcripts of three genomic RNAs from the recombinant plasmids are mixed, CMV becomes infectious; a pseudorecombinant CMV can be easily created between two CMVs (e.g., CMV-Y and CMV-L) by mixing the viral genomic RNAs. The viral vector construct based on CMV-Y RNA2 is designated A1, which has the cloning site between StuI and MluI for a foreign insert (Figure 1A). Our VIGS system for spinach consists of three sequential steps: (1) cloning of a partial sequence of the spinach gene of target into A1, (2) propagation of the recombinant A1 vector in N. benthamiana by rub-inoculating RNA transcripts, (3)



Upper leaves ('All Right') 14 dpi



Figure 1. Construction of the CMV vector for VIGS. (A) Schematic representation of the CMV vector A1, whose backbone is RNA2 encoding the 2a and 2b genes. The spinach genes (*PDS* and *DFR*) were cloned between the StuI and MluI site in the antisense orientation. (B) Symptoms on the upper leaves of spinach ('All Right') infected with the original vectors of Y1A1Y3 (left) and L1A1L3 (right).

inoculation of leaf sap of infected *N. benthamiana* onto spinach. In a preliminary experiment, we found that Y1A1Y3 induced severe mosaic symptoms in spinach although the severity varied depending on the tested cultivars. We thus used a pseudorecombinant virus, L1A1L3 that induced relatively mild symptoms on those cultivars (Figure 1B). It is important to use a mild (or symptomless) virus because severe symptoms may mask the silenced phenotype in the infected plants.

To evaluate the CMV-based VIGS system, the 130-nucleotide (nt) fragment of the PDS gene was first cloned in the A1 vector to create A1-PDS (Figure 1A). Primers used for cloning are shown in the Supplementary Table S1. In vitro transcripts of L1, A1-PDS and L3 were inoculated onto N. benthamiana, and subsequently the leaf sap of infected N. benthamiana was inoculated onto leaves of two spinach cultivars, 'All Right' (TAKII & CO., LTD., Kyoto, Japan) and 'Surprise' (TOHOKU SEED Co. LTD., Utsunomiya, Japan). In our preliminary experiments, we had found that 'Surprise' was a little more susceptible to the pseudorecombinant virus, L1A1L3 than 'All Right.' As shown in Figures 2A and 3A, distinct chlorosis was observed at 7-10 dpi along the veins in upper systemically infected leaves. As expected, the PDS mRNA levels were decreased down to less than 1/4 of those of the healthy plants at 10 dpi (Figures 2B, 3B), and photobleaching has lasted over two months (Supplementary Figure S1). There was little difference in photobleaching between the two



Figure 2. VIGS against the *PDS* gene in the spinach cultivar 'All Right'. (A) Photobleaching phenotype observed on the upper systemic leaves of spinach inoculated with L1A1-PDS2L3 at 10 dpi. The pictures of leaves of healthy plants and empty vector-infected plants are shown for comparison. (B) *PDS* expression levels in the upper non-inoculated leaves at 10 dpi. Real-time RT-PCRs were conducted using primer pairs as in Supplementary Table S1. The *glyceraldehyde 3-phosphate* (*GAPDH*) gene was used as a reference gene. Means (\pm SE) among the infected plants were analyzed for significant differences using Tukey's multiple comparison test (**p*<0.05); different letters above the bars indicate a significant difference among isolates. (C) CMV accumulation levels in upper non-inoculated leaves at 10 dpi. Real-time RT-PCR for the virus accumulation was conducted using primer pair of CMV-DET-5-340 and CMV-DET-3-340 (Supplementary Table S1). The CMV-DET-5-340 primer hybridizes to both CMV-L RNA3 (positions 1864–1884) and the subgenomic RNA from RNA3 designated RNA4, which is the mRNA for the coat protein. The CMV-DET-3-340 primer hybridizes to the 3' end of RNA3 and RNA4. The size of the PCR product is 353 bp. The *GAPDH* gene was used as a reference (Supplementary Table S1). Statistical analysis was conducted by Student's *t*-test (**p*<0.05).

cultivars when the vector contains the insert fragment, suggesting that the introduction of an insert in the vector may affect the viral pathogenicity. However, the viral accumulation level in 'Surprise' was found to be about 1/200 of the control (L1A1L3) while about 1/5 in 'All Right,' suggesting that viral accumulation levels are different in different cultivars and that a very low level of viral accumulation can still induce efficient VIGS against *PDS* (Figures 2C, 3C).

To further test whether this vector can be used for functional genomics of spinach genes, we chose the *dihydroflavonol 4-reductase* (*DFR*) gene, which is one of the key genes in the anthocyanin pathway (Shimada et al. 2004). In green spinach, anthocyanins are not normally synthesized but instead, betacyanins accumulate. However, it has been reported that some red-purple cultivars produce anthocyanins in leaves and stem although all red-purple cultivars cannot necessarily synthesize anthocyanins (Cai et al. 2018). We here cloned the 130-nt spinach *DFR* fragment into the A1 vector to produce A1-DFR using primer pair shown in the Supplementary Table S1 and induced VIGS by inoculating A1-DFR onto several red-purple spinach cultivars, whose seeds were purchased on the Japanese market: 'Kuroshio,' Kobayashi-Shubyo, Hyogo; 'Akajiku-Salad,' Musashino-Shubyo, Tokyo; 'Wase-Salad-Akari,' TAKII, Kyoto; 'Akakuki-Minster,' Nakahara-Saishujyo, Fukuoka; 'Miyabi,' TOHOKU, Utsunomiya. Plants were grown in the plant incubator at 24°C under 12h day. These inoculated plants did not show any severe symptoms but did very mild symptoms in upper leaves (Supplementary Figure S2). Although the red pigmentation was not decreased in most of the tested cultivars (five cultivars, 4 individual plants tested for each cultivar) by A1-DFR VIGS, we observed clear decoloration of the red pigment in the stem of one of the used cultivars (cultivar 'Kuroshio,' Figure 4A), suggesting that this cultivar may produce anthocyanins. We also confirmed that the DFR mRNA levels in the A1-DFR-infected plants were greatly reduced at 10 dpi in spite of the fact that the empty vector (L1A1L3) infection itself increased the DFR expression (Figure 4B). Our real-time RT-PCRs showed that the viral accumulation levels in L1A1-DFR2L3-infected plants with the silenced phenotype were about 1/1000 of the control (L1A1L3), suggesting that the high level of viral accumulation of L1A1L3 may have caused the huge increase in DFR expression (Figure 4B, C). There are



Figure 3. VIGS against the *PDS* gene in the spinach cultivar 'Surprise'. (A) Photobleaching observed on the upper leaves of spinach inoculated with L1A1-PDS2L3 at 10 dpi. The pictures of leaves of healthy plants and empty vector-infected plants are shown for comparison. (B) *PDS* expression levels in the upper non-inoculated leaves at 10 dpi. Real-time RT-PCRs were conducted using primer pairs as in Supplementary Table S1. Statistical analysis for real-time RT-PCR of *PDS* was conducted using Tukey's multiple comparison test (*p < 0.05); different letters above the bars indicate a significant difference among isolates. (C) CMV accumulation levels in upper non-inoculated leaves at 10 dpi. Real-time RT-PCR was conducted as explained in Figure 2. Statistical analysis was performed using Student's *t*-test (**p < 0.001).



Figure 4. Expected silencing phenotype (decoloration) observed in the stem of spinach inoculated with L1A1-DFR2L3 at 10 dpi. The set of pictures is VIGS against the *DFR* gene in the red-purple spinach cultivar 'Kuroshio.' The pictures of the stems from three individual plants (healthy plants and L1A1L3-infected plants) are shown for comparison. The *GAPDH* gene was used as a reference gene. (B) Real-time RT-PCR was conducted using primer pairs as in Supplementary Table S1. To maintain homogeneity of variance among the sample groups, the raw data was converted to log data for the statistical analysis. Means (\pm SE) among the infected plants were analyzed for significant differences using Tukey's multiple comparison test (p < 0.05); different letters above the bars indicate a significant difference among isolates. (C) Comparison of viral accumulation by real-time RT-PCR. Total RNA was extracted from stem. Real-time RT-PCR was conducted as explained in Figure 2. Statistical analysis was performed using Student's *t*-test (*p < 0.01).

two possible explanations for the observation that the viral accumulation level of L1A1-DFR2L3 was very low at 10 dpi in the systemic tissues. The CMV vector itself is degraded by VIGS in infected tissues but normally tolerant to VIGS to some extent because of its secondary structure. If the insertion of a target sequence in the vector alters such a secondary structure, then the viral genomic RNA would become much more susceptible to VIGS, resulting in a great decrease in viral titer. As VIGS is efficiently induced against the target gene, the virus containing the target sequence will be also efficiently degraded. Alternatively, the insertion of a foreign sequence itself sometimes directly interferes with the viral replication, depending on both the size and sequence. We assume that either or both of the two cases may have occurred on L1A1-DFR2L3. In addition, to examine whether L1A1-DFR2L3 loses the insert sequence in systemically infected upper leaves, we analyzed the leaves just beneath bolted flowers of the plants at 60 dpi. The results showed that the virus still retained the complete insert, and the accumulation level was about 1/15 of the control (L1A1L3), suggesting that the virus can maintain its systemic infection even after two months postinoculation (Supplementary Figure S3). These results altogether demonstrate that the CMV-based VIGS vector can be used for functional genomics in spinach without inducing severe symptoms.

In conclusion, we here tried to develop another VIGS system for spinach, applying the CMV vector to VIGS against spinach *PDS* and *DFR*, which would be more practical than the other two BCTV- and TRV-based VIGS systems. This CMV construct can systemically infect many spinach cultivars and efficiently accumulate in infected plants. Because we keep some CMV strains isolated directly from spinach, we can use those viral RNA genomes to adjust the viral pathogenicity and symptoms. For example, even though the initial construct does not efficiently infect a cultivar of interest due to a resistant gene, we can test other pseudorecombinant viruses using CMV spinach strains that can overcome the resistance.

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