A novel method for the evaluation of virus-induced gene silencing efficiency

C. Li1,2, Z.C. Zhang1, K.M. Ghebremariam1, L.H. Wang1, L. Wu1 and Y. Liang1

1State Key Laboratory of Crop Stress Biology in Arid Region, College of Horticulture, Northwest A&F University, Yangling, Shaanxi, China
2College of Life Science, Huaibei Normal University, Huaibei, Anhui, China

Corresponding author: Y. Liang
E-mail: liangyan@nwsuaf.edu.cn

Received June 25, 2013
Accepted February 13, 2014
Published November 11, 2014
DOI http://dx.doi.org/10.4238/2014.November.11.9

ABSTRACT. Virus-induced gene silencing (VIGS) is an important tool for studying gene function. However, a number of factors highly restrict the application of VIGS, such as unstable efficiency and tissue-specific silencing. We developed a novel evaluation method for improving the applicability of VIGS vectors. In this method, 4 indexes were defined and utilized to evaluate VIGS efficiency by silencing the endogenous phytoene desaturase \((PDS)\) gene with a tobacco rattle virus-based VIGS vector. To illustrate the reliability of this evaluation method, we assessed the silencing efficiency of \(\textit{SpPDS}\) and \(\textit{SpMPK1}\) in \textit{Solanum pimpinellifolium}. The silencing results of \(\textit{SpPDS}\) showed that an optical density at 600 nm of 2.0 was more suitable than 1.0 for VIGS in \textit{S. pimpinellifolium}. This suggests that the proposed evaluation method is a valid technique for optimizing the VIGS system of plants. Moreover, the \(\textit{SpMPK1}\) gene was highly silenced in the 4th-9th leaves with a 50-95% reduction in transcription levels, further demonstrating that this method can be used to select highly silenced candidates for further
experiments, particularly when the target gene shows no phenotypic change after being silenced.

Key words: Evaluation method; Virus-induced gene silencing (VIGS); Silencing efficiency; Solanum pimpinellifolium

INTRODUCTION

Virus-induced gene silencing (VIGS), an approach used for post-transcriptional gene silencing (Baulcombe, 1999), is an important tool for investigating gene function and high-throughput functional genomics in plant science by exploiting plant RNA interference-mediated antiviral defense mechanism (Liu et al., 2002b; Ekengren et al., 2003; Benedito et al., 2004; Burch-Smith et al., 2004). Viral vectors carrying sequences derived from host genes can be targeted to the mRNAs of a corresponding host using a post-infiltrated process (Bartel, 2004). Tobacco rattle virus (TRV) is one of the most frequently used vectors for VIGS because of its increased capability of spreading throughout the plant as well as its relatively mild impact on plant health relative to other viruses such as tobacco mosaic virus and potato virus X (Ratcliff et al., 2001). TRV is a bipartite, single-stranded, positive-sense RNA plant virus consisting of two types, TRV1 and TRV2; TRV2 typically harbors a coat protein and a fragment from the host gene, whereas TRV1 is responsible for encoding the viral replication and movement factors and facilitates TRV2 mobility during silencing (Liu et al., 2002b). The combination of TRV1 and TRV2 results in gene silencing in plants.

As a tool for functional analysis, TRV-based VIGS has been used in a broad range of plants, including tobacco (Ratcliff et al., 2001; Jin et al., 2002; Peart et al., 2002a,b; Ahn et al., 2004; del Pozo et al., 2004; Saedler and Baldwin, 2004; Valentine et al., 2004), tomato (Liu et al., 2002a; Ekengren et al., 2003; Brigneti et al., 2004; Ryu et al., 2004; Kandoth et al., 2007; Wu et al., 2011; Hosseini Tafreshi et al., 2012; Liu et al., 2012), pepper (Chung et al., 2004) and petunia (Chen et al., 2004; Ryu et al., 2004). However, the VIGS method has some limitations, resulting in an unstable silencing efficiency, localized tissue silencing, and variable silencing efficiencies among different genera and even within the same species caused by factors such as the length of insertion, seedling age, and culture conditions. According to previous studies, a 400-bp insertion is preferable for promoting effective gene silencing and for limiting the development of damage symptoms (Wu et al., 2011). Furthermore, previous studies have shown that 7-, 14-, or 21-day-old tomato seedlings, grown at 18-21°C are suitable for VIGS (Liu et al., 2002a; Ekengren et al., 2003; Velásquez et al., 2009; Wu et al., 2011). Studies aimed at improving VIGS efficiency have been reported in Solanum; however, silencing is not 100% efficient. Few studies have been conducted using other plants. Thus, an evaluation method is needed for assessing VIGS efficiency to optimize and select for highly silenced candidates.

In this study, we developed a novel VIGS evaluation method. To illustrate the reliability of this method, we assessed VIGS efficiency by silencing the endogenous phytoene desaturase gene (SpPDS) with a TRV-based VIGS vector in Solanum pimpinellifolium. In addition, the results were validated by successful silencing of the SpMPK1 gene in S. pimpinellifolium. Our data strongly indicate that this method is applicable for evaluating the VIGS system and highly silenced candidates. Moreover, our method can be applied for optimizing the VIGS system in other plants, thereby improving the applicability of VIGS vectors.
MATERIAL AND METHODS

Plant material and growth conditions

*S. pimpinellifolium*, commonly known as the proposed wild progenitor of domesticated tomato (van der Knaap et al., 2004), was used in this study. Seeds of *S. pimpinellifolium* (L03708) were cultivated in cell trays filled with matrix in an artificial climate incubator at 21°C ± 2°C with 50% relative humidity and a 16-h photoperiod with light intensity at 300-400 μmol·m⁻²·s⁻¹.

TRV vector construction and agroinfiltration in *S. pimpinellifolium*

The pTRV1 and pTRV2 VIGS vectors described previously (Liu et al., 2002b) were obtained from Dr. S.P. Dinesh-Kumar (University of California, Davis, CA, USA). The insertion was amplified by polymerase chain reaction (PCR) using a cDNA template synthesized using the PrimeScript RT reagent Kit (Takara, Dalian, China) from total RNA extracted from leaf tissues of *S. pimpinellifolium* L03708 using Trizol reagent (Invitrogen, Carlsbad, CA, USA). The following primer pairs were used: 5'-GGCGCGAGCTCGGCACTCAACTTTA TAAACC-3' and 5'-CGGCGCTCGAGCTTTCACTACGCA CGATTGTTC-3' (XhoI and SacI sites are underlined). A 409-bp product of *SpPDS* obtained from the digestion after PCR was cloned into the respective sites of pTRV2 plasmid digested with XhoI/SacI to produce the pTRV2: *PDS* plasmid. A 200-bp fragment of *SpMPK1*, which was PCR-amplified from the cDNA template using a pair of primers (forward: 5'-GGCGCGAGCTGATAATTGCTGAC AGATTGTTC-3' and reverse: 5'-CGGCGCTCGAGCATTTCAGTCTCAAAATCTACGCA CGATTGTTC-3'), was linked to pTRV2 to produce the pTRV2: *MPK1* plasmid.

Next, the pTRV1 and pTRV2: *PDS* or pTRV2: *MPK1* were introduced into the *Agrobacterium tumefaciens* strain GV3101 by electroporation. A colony was grown at 28°C overnight with shaking at 200 rpm in Luria-Bertani broth containing antibiotics (50 μg/mL kanamycin, 10 μg/mL rifampicin, and 50 μg/mL gentamycin). Next, secondary liquid induction media containing antibiotic and 200 μM acetosyringone was used to culture the bacteria to an optical density at 600 nm (OD₆₀₀) of 0.5-0.8. Cells were harvested and resuspended twice in suspension media (10 mM MgCl₂, 10 mM MES, 200 mM acetosyringone, pH 5.5) (Ahn et al., 2004), and the sample was adjusted to an OD₆₀₀ of 1.0 or 2.0 and incubated at room temperature for 3 h. The mixture of pTRV1 and pTRV2: *PDS*/pTRV2: *MPK1* in a 1:1 ratio was prepared and infiltrated into both cotyledons as described previously (Ahn et al., 2004), using pTRV1/pTRV2:00 as a negative control.

Evaluation of silencing efficiency

The evaluation method was composed of 4 indexes, as follows: 1) Silencing frequency of plants (SFP) was defined as the percentage of plants showing a silencing phenotype compared to the total number of plants infiltrated with pTRV2: *PDS*. We propose this as the first index in evaluating the silencing efficiency of VIGS. The following equation was used:

\[
SFP = \frac{\text{Number of plants showing silencing phenotype (bleaching or yellowing)}}{\text{Total number of plants infiltrated}} \times 100\% \quad \text{(Equation 1)}
\]
2) Silencing efficiency of a plant (SEP) was calculated by counting the number of leaves showing photobleaching on a gene-silenced plant.

\[
SEP = \frac{\text{Number of leaves showing silencing phenotype on a plant}}{\text{Total number of leaves on that plant}} \times 100\% \quad (\text{Equation 2})
\]

3) Silencing frequency of each leaf (SFL) was defined as the frequency of a particular leaf showing the silencing phenotype in all gene-silenced plants at 40 days post-infiltration (dpi). This index specifies the location of silencing.

\[
SFL = \frac{\text{The number of a certain leaf showing silencing phenotype}}{\text{Total number of plants showing phenotype}} \times 100\% \quad (\text{Equation 3})
\]

4) Silencing efficiency of gene (SEG) was defined as the quantification of gene silencing. It was computed in two ways: the reduction of gene transcription levels and the ratio of bleaching areas. The following method was used for calculation.

The reduction of gene transcription levels refers to the difference between the SpPDS gene transcription levels in control plants (defined as 100%) and those in gene-silenced plants. To quantify the abundance of SpPDS transcription in silenced and control plants, quantitative reverse transcription (RT)-PCR was performed. Total RNA was extracted from leaves, and first-strand cDNA was synthesized using the PrimeScript RT reagent Kit (Takara). Quantitative RT-PCR was performed using SYBR Premix Ex Taq II (Takara) in an iQ5 Real-Time PCR Detection System (Bio-Rad, Hercules, CA, USA). To quantify gene transcription in silenced plants, the $2^{-\Delta\Delta C_T}$ method was applied. Transcription levels of S. pimpinellifolium elongation factor 1-α were internally controlled to normalize the data for SpPDS and SpMPK1 transcripts. The primers used in experiment were forward: 5′-ATGCCACGACCAGAAG-3′ and reverse: 5′-TGCTGTAGACAAACCACCC-3′ (for SpPDS); forward: 5′-GACAGGCGTTCAAGGTAAGG-3′ and reverse: 5′-CCAATGGAGGGTATTCAGC-3′ (for elongation factor 1-α); and forward: 5′-ATCCCAAGAGGAATAACAG-3′ and reverse: 5′-ATCAACCTCAGCAACAATCT-3′ (for SpMPK1).

\[
\text{Reduction of gene transcripts} = (1 - \text{relative transcripts}) \times 100\% \quad (\text{Equation 4})
\]

The ratio of bleaching areas was defined as the bleached areas compared to the total area of detected leaves. Quantification was accomplished by sampling and photographing 3 leaves of each plant with the camera lens perpendicular to the flat leaves. Photographs were then imported into the Adobe Photoshop CS4 software (Adobe Systems Corp., San Jose, CA, USA) to calculate the total area of the 3 leaves as well as the bleaching and yellowing areas using the Magnetic Lasso and Straw tools.

\[
\text{Ratio of bleaching areas} = \frac{\text{total areas of bleaching and yellowing}}{\text{total area of the three leaves}} \times 100\% \quad (\text{Equation 5})
\]

Statistical analysis

R 2.15.2, which is an open-source software, was used for statistical analyses. Silencing
efficiency of plants was analyzed by univariate analysis of variance followed by a post hoc test. Gene silencing efficiency was analyzed using the Pearson product-moment correlation test.

RESULTS

To verify the applicability of the proposed evaluation method, 2 experiments were conducted. First, an Agrobacterium suspension was prepared and infiltrated into tomato seedlings, and then the silencing efficiency of the SpPDS gene was evaluated during 15-40 dpi. In addition, to validate the accuracy and reliability of the evaluation results, silencing of the SpMPK1 gene and the concurrent evaluation of its silencing efficiency was carried out.

Application of the evaluation method for assessing VIGS efficiency

Silencing efficiency of SpPDS at OD<sub>600</sub> of 1.0

Plants inoculated with pTRV2:PDS (OD<sub>600</sub> = 1.0) developed photobleaching symptoms at 12 dpi; experimental data were collected until 40 dpi. As shown in Figure 1A, the SFP increased rapidly from 25 to 50% from 12-15 dpi, and then gradually increased to reach a maximum of 65% at 25 dpi. Based on these results, 35 plants showing photobleaching symptoms were used to calculate the SEP (Figure 1B, box-plot). At 15 dpi, the median line was located at 40% (indicating that 40% of the total leaves were bleached on plants on average) and then reached 60% at 20 dpi.

Figure 1. VIGS efficiency of SpPDS at an OD<sub>600</sub> of 1.0 in Solanum pimpinellifolium. A. Silencing frequency of plants. The values are reported as means ± SD of 3 independent experiments (N ≥ 20) and the bars represent the standard errors. B. Silencing efficiency of plants. Thirty-five VIGS plants were used in the experiment and the results are shown as box-plots. C. Silencing frequency of each leaf at 40 dpi. The experiments were representative of 35 VIGS plants. D. Silencing efficiency of the SpPDS gene. Leaves 4-7 from 25 VIGS plants were used in the experiment and quantitative RT-PCR was performed with 3 replicates.
Unfortunately, long-ranging boxes were observed at 15 and 20 dpi, reflecting the instability of silencing efficiency among the plants. At 25 dpi, the SEP reached a maximum for the up-moved box, but then decreased at a later point, indicating that the newly emerged leaves did not show the silencing effect.

As described above, SFL reflects the silence-specific position within the plant. In this assay, photobleaching symptoms were observed in leaves 3-11, and SFL in leaves 4-7 reached 75-95% (Figure 1C). These results indicate that the SpPDS gene was highly silenced in leaves 4-7. SEG was calculated by performing quantitative RT-PCR for leaves 4-7 from 25 VIGS plants. As shown in Figure 1D, the transcription levels of the SpPDS gene ranged from 5-35%, showing a large decrease compared to control plants. Correspondingly, SEG ranged from 65-95%. These results indicate that an OD$_{600}$ of 1.0 is not the optimal concentration for VIGS in *S. pimpinellifolium* because of the lower silencing frequency of plants, unstable silencing efficiency, and shortened duration of silencing symptoms. This suggests the successful application of the evaluation method for assessing VIGS efficiency.

**Correlation between the reduction of gene transcription and the ratio of bleaching areas**

The SpPDS SEG was quantified by calculating both the ratio of bleached areas and the reduction of gene transcription levels. Leaves 3-11 of VIGS plants were analyzed and the results are shown in Figure 2. In the first method, the ratio of bleaching areas ranged from 20-97%. In the second method, the reduction in SpPDS transcript abundance was determined using quantitative RT-PCR with RNA isolated from the same samples; transcript abundance ranged from 27-98%. The results obtained using both methods showed a significant positive correlation ($r = 0.873$, $P < 10^{-9}$). This suggests that the reduction in SpPDS gene transcription levels or the ratio of bleaching areas can be used to calculate SpPDS gene silencing efficiency.

![Figure 2](image-url)  
**Figure 2.** Correlation between the reduction of gene transcripts and the ratio of bleaching areas. Leaves 3-11 were sampled randomly from silenced plants from 15-40 dpi ($r = 0.873$, $P < 10^{-9}$).
Evaluation of virus-induced gene silencing efficiency

Evaluation of SpMPK1 gene silencing efficiency at OD$_{600}$ of 2.0

To further demonstrate the general reliability of this evaluation method, we silenced the SpMPK1 gene in S. pimpinellifolium using the SpPDS gene as a positive control. Because of the lower silencing efficiency of VIGS at an OD$_{600}$ of 1.0, the infiltration concentration was increased to 2.0 in this assay.

Silencing efficiency of SpPDS at OD$_{600}$ of 2.0

Plants inoculated with an OD$_{600}$ of 2.0 showed a much higher silencing frequency. Initially (12 dpi), the SFP was 50% and then increased to 70% at 15 dpi before finally reaching a maximum (92.3%) at 25 dpi (Figure 3A and C). Thus, more than 50 VIGS plants were obtained and the SEP was calculated. As shown in Figure 3D, all box plots were relatively small throughout the experimental time, reflecting stable silencing efficiency in the plants. Median lines in box plots were located at >50%, over 70% at 25-30 dpi, and then stabilized at approximately 60% at 40 dpi. Thus, higher silencing efficiency was observed in VIGS plants.

Figure 3. VIGS efficiency of SpPDS at an OD$_{600}$ of 2.0 in S. pimpinellifolium. A. Photobleaching phenotype induced by pTRV:PDS in S. pimpinellifolium. B. Leaves used to calculate the ratio of bleaching areas. C. Silencing frequency of plants. Values are reported as means ± SD of 3 independent experiments (N ≥ 20) and the bars represent the standard errors. D. Silencing efficiency of plants. Fifty VIGS plants were used in the experiment; results are shown as box plots. E. Silencing frequency of each leaf at 40 dpi. The experiments were representative of 50 VIGS plants. F. Silencing efficiency of the SpPDS gene. SpPDS gene silencing efficiency was assessed by the ratio of bleaching areas. The experiment was performed using leaves 4-9 from 25 VIGS plants independently.
During the experimental time, photobleaching symptoms were observed in leaves 2-13, and the SFL of leaves 4-9 increased to 80-100% (Figure 3E). These results show that the SpPDS gene may be highly silenced in this position. Concurrently, 25 VIGS plants were selected to test the SEG of leaves (4-9) by calculating the ratio of bleaching areas (Figure 3B). As predicted, the SEG was 65-95% (Figure 3F). Furthermore, more VIGS plants were obtained and more persistent photobleaching symptoms were observed when the infiltration concentration was an OD\textsubscript{600} of 2.0 compared to an OD\textsubscript{600} of 1.0. These results strongly suggest that an OD\textsubscript{600} of 2.0 is better than 1.0 for VIGS in \textit{S. pimpinellifolium}, at least in the context of our experiment. Thus, the evaluation method based on \textit{PDS} is applicable for optimizing the VIGS system in plants.

\textbf{Silencing efficiency of SpMPK1 at OD\textsubscript{600} of 2.0}

The results described above show that when the infiltrated concentration was an OD\textsubscript{600} of 2.0, 92.3% of the infiltrated plants were VIGS and that leaves 4-9 were optimal samples for further experiments. To validate the accuracy of these results, we attempted to silence the \textit{SpMPK1} gene in \textit{S. pimpinellifolium}. pTRV2:SpMPK1 \textit{Agrobacterium} (OD\textsubscript{600} of 2.0) was infiltrated into cotyledons and the silencing efficiency of \textit{SpMPK1} was assessed by quantitative RT-PCR at 25 dpi. The results showed that 27 VIGS plants were obtained from 30 of the infiltrated plants (90%). Of these, 25 plants were effectively silenced, as shown by a significant reduction in \textit{SpMPK1} gene transcription levels (50-93%) in leaves 4-9 (Figure 4). These results further demonstrate that the evaluation method is applicable for studying the function of other genes by improving the overall accuracy of the results.

\textbf{Figure 4.} \textit{SpMPK1} gene silencing efficiency in leaves 4-9 from VIGS plants. The transcription levels of \textit{SpMPK1} were analyzed by quantitative RT-PCR at 25 dpi and presented as the percentages of the mean levels of control plants, which were defined as 100%.
DISCUSSION

The TRV-based VIGS vector is one of the most widely used virus vectors and has been repeatedly shown to be useful for studying gene function in plants. However, although many factors related to TRV-induced gene silencing efficiency have been studied, silencing is never 100% efficient in the tomato plant, nor in any other plant. Therefore, we developed a novel evaluation method to increase the silencing efficiency of candidates used for further experiments.

Typically, PDS is selected as a marker gene in VIGS system optimization, as a reduction in PDS gene transcripts represents the silencing efficiency of VIGS (Rotenberg et al., 2006). For some plant species in which the PDS gene sequences are unknown, the reduction of their transcript abundances within silenced plants is substituted with chlorophyll (Senthil-Kumar et al., 2007). There is currently no systematic approach for evaluating VIGS efficiency. In this study, our evaluation method included 4 indexes (SFP, SEP, SFL, and SEG). SFP and SEP were used to optimize the VIGS system in other plants. Furthermore, SFL was used to assess the gene silencing position. Lastly, SEG was used to determine that standardized samplings from gene-silenced plant were sufficient for analyzing experimental results when the gene of interest in VIGS presented without phenotypic changes. In addition, the simplicity of identifying the ratio of bleaching areas to quantify gene silencing efficiency is advantageous compared to the more difficult experimental procedures often used, such as identifying gene transcription levels and chlorophyll content.

In conclusion, because VIGS is currently applied in a growing number of plant species and is widely used in functional genomic studies of plants, it is necessary to improve the applicability of VIGS vectors. In this study, we developed a novel evaluation method and demonstrated the applicability of this model for optimizing the VIGS system in other plants as well as in selecting highly silenced candidates for further experiment when genes requiring silencing are present without phenotypic changes.

ACKNOWLEDGMENTS

We thank the Asian Vegetable Research and Development Center (AVRDC), the World Vegetable Center for supplying the experimental material. Research supported by the “13115” Technology Innovation Project of Shaanxi Province (#2009ZDKG-14) and the Science and Technology Coordination and Innovation Project of Shaanxi Province (#2011KTCL02-03).

REFERENCES


van der Knaap E, Sanyal A, Jackson SA and Tanksley SD (2004). High-resolution fine mapping and fluorescence *in situ* hybridization analysis of sun, a locus controlling tomato fruit shape, reveals a region of the tomato genome prone to DNA rearrangements. *Genetics* 168: 2127-2140.
