**Foxtail Mosaic Virus-Induced Gene Silencing in Monocot Plants**

Na Liu, Ke Xie, Qi Jia, Jinping Zhao, Tianyuan Chen, Huangai Li, Xiang Wei, Xianmin Diao, Yiguo Hong, and Yule Liu*

Center for Plant Biology and MOE Key Laboratory of Bioinformatics, School of Life Sciences, Tsinghua University, Beijing 100084, China (N.L., K.X., Q.J., J.Z., T.C., X.W., H.L., Y.L.); Institute of Crop Sciences, Chinese Academy of Agricultural Sciences, Beijing 100081, China (X.D.); Research Centre for Plant RNA Signaling, College of Life and Environmental Sciences, Hangzhou Normal University, Hangzhou 310036, China (Y.H.); and Institute of Virology and Biotechnology, Zhejiang Academy of Agricultural Sciences, Hangzhou 310021, China (J.Z.)

ORCID ID: 0000-0002-4423-6045 (Y.L.).

Virus-induced gene silencing (VIGS) is a powerful technique to study gene function in plants. However, very few VIGS vectors are available for monocot plants. Here we report that Foxtail mosaic virus (FoMV) can be engineered as an effective VIGS system to induce efficient silencing of endogenous genes in monocot plants including barley (*Hordeum vulgare* L.), wheat (*Triticum aestivum* L.) and foxtail millet (*Setaria italica*). This is evidenced by FoMV-based silencing of phytoene desaturase (PDS) and magnesium chelatase in barley, of PDS and chloroplasts alteredos1 in foxtail millet and wheat, and of an additional gene *IspH* in foxtail millet. Silencing of these genes resulted in photobleached or chlorosis phenotypes in barley, wheat, and foxtail millet. Furthermore, our FoMV-based gene silencing is the first VIGS system reported for foxtail millet, an important C4 model plant. It may provide an efficient toolbox for high-throughput functional genomics in economically important monocot crops.

Virus-induced gene silencing (VIGS) provides a powerful approach for genetic and functional characterization of genes in plants. VIGS is related to post-transcriptional gene silencing, a plant antiviral defense mechanism that targets and leads to degradation of viral RNAs (Vance and Vaucheret, 2001; Baulcombe, 2004). Post-transcriptional gene silencing in plants is dependent on a relatively high degree of nucleotide homology between RNA transcripts and target endogenous gene sequences (van den Boogaart et al., 1998; Ding, 2000). The recombinant virus-carrying sequence of a host gene generates dsRNAs, which are processed by Dicer-like proteins to produce siRNAs, and the latter triggers silencing of the endogenous gene itself. This forms the basis of plant VIGS technology.

Many plant RNA and DNA viruses have been modified as VIGS vectors and they are used to dissect gene function in diverse dicot plant species (Burch-Smith et al., 2004; Senthil-Kumar and Mysore, 2011; Huang et al., 2012). For instance, *Tobacco rattle virus*-derived vector is one of the most widely used systems in dicots because of its broad host range and ability to infect meristematic tissue (Ratcliff et al., 2001; Liu et al., 2002a, Liu et al., 2002b; Padmanabhan and Dinesh-Kumar, 2009). *Bean pot mottle virus*-derived vectors have been developed to induce VIGS in soybean (*Glycine max*) and common bean (*Phaseolus vulgaris*; Zhang and Ghabrial, 2006; Zhang et al., 2010). *Cabbage leaf curl virus*-derived vector has already been developed to trigger VIGS for siRNA-mediated silencing in Arabidopsis (Turnage et al., 2002) and miRNA-mediated silencing in tobacco (*Nicotiana benthamiana*; Tang et al., 2010).

By contrast, to date only four RNA viruses and one DNA virus have been modified as vectors for VIGS in monocot species, of which, *Barley stripe mosaic virus* (BSMV)-based VIGS has been applied for functional genomics in barley (*Hordeum vulgare* L.) and wheat (*Triticum aestivum*; Holzberg et al., 2002; Tai et al., 2005; Meng et al., 2009; Facak et al., 2010; Yuan et al., 2011); *Brome mosaic virus* (BMV) in rice (*Oryza sativa* L.), barley, and maize (*Zea mays* L.; Ding et al., 2006); *Bamboo

---

1This work was supported by the National Transgenic Program of China (grants no. 2016ZX08009-001, no. 2014ZX0800104B, and no. 2016ZX08009-003); the National Natural Science Foundation of China (grants no. 31270182, no. 31470254, no. 31530059, no. 30800594, and no. 31370180); and the National Basic Research Program of China (grant no. 2014CB139400).

2These authors contributed equally to this work.

* Address correspondence to yuleliu@mail.tsinghua.edu.cn.

Y.L., N.L., and K.X. initiated the project, designed the experiments, and analyzed the data; Y.L. and N.L. wrote the article; N.L., K.X., Q.J., J.Z., T.C., H.L., and X.W. performed the experiments; X.D. helped to analyze the data; and Y.H. contributed to this work through discussions and detailed revision of the article.

1OPEN] Articles can be viewed without a subscription.

www.plantphysiol.org/cgi/doi/10.1104/pp.16.00010
mosaic virus and its satellite RNA in *N. benthamiana* and *Brachypodium distachyon* (Liou et al., 2014); and *Rice tungro bacilliform virus* in rice (Purkayastha et al., 2010). Very recently, Cucumber mosaic virus-based VIGS in maize is reported (Wang et al., 2016) BMV and BMV-based vectors are frequently used for VIGS in some monocot plants. However, both virus vectors do not work in many other monocot species and not all cultivars within a particular host species (Ding et al., 2006; Pacak et al., 2010).

**Foxtail mosaic virus** (FoMV) is a species of the genus *Potexvirus* and possesses a broad host range including 56 Poaceae species and at least 35 dicot species (Paulsen and Niblett, 1977; Short and Davies, 1987). A VIGS system based on *potato virus X* (PVX), the type member of *Potexvirus*, has been developed to silence target genes successfully in plants (Ruiz et al., 1998; Chen et al., 2015; Zhao et al., 2016). FoMV is similar to PVX and other *potexviruses* in their genomic organization. It consists of a positive-sense single-stranded RNA genome with a 5′-methylguanosine cap, a 3′-polyadenylated tail, and five major open reading frames (ORFs) and a unique 5A gene (Robertson et al., 2000; Bruun-Rasmussen et al., 2008). Each of the five major ORFs encodes a functional protein as in PVX (Bruun-Rasmussen et al., 2008). FoMV has been modified as an RNAi suppressor-dependent expression vector by deleting its coat protein (CP) and triple gene block genetic segments (Liu and Kearney, 2010). However, FoMV has not been successfully engineered as a VIGS vector, although it has been thought to be a potentially useful virus for VIGS in monocot hosts (Scофield and Nelson, 2009).

In this study, we report that a FoMV-based vector can be used to perform VIGS in monocots including barley, wheat, and foxtail millet (*Setaria italica*), and may provide a useful tool to study gene function in a broad monocot plant species.

**RESULTS**

**Development of FoMV VIGS Vector**

To develop an FoMV VIGS vector, the full-length FoMV cDNA (Fig. 1A, a kind gift from Nancy Robertson) was recloned between the duplicated 35S CaMV promoter and the nitric oxide synthase (NOS) terminator of a T-DNA binary vector. pFoMV-wt has an additional 70 adenosine residues after the viral 3′-terminus. C, Schematic of the FoMV VIGS vector, pFoMV-sg, is a pFoMV-wt derivative that contains a direct repeat of the 300-bp putative FoMV CP subgenomic promoter (GenBank accession: EF630359, nt 5202–5371), shown by arrows, and multiple cloning sites (HpaI, MluI, XhoI, and Ascl) between CP subgenomic promoters. Right- and left-pointing blue arrows indicate sense and anti-sense orientation of foreign fragments, and the predicted structure of RNA transcript is presented on the right side. D, Symptom development and RT-PCR detection of viral RNA showed that pFoMV-wt and pFoMV-sg are infectious in barley. Bar = 1 cm.

**Figure 1.** Schematic organization of FoMV genome and infectious FoMV vector. A, Genomic organization of FoMV RNA. The ORFs are indicated by boxes. The 152-kD protein is an RNA-dependent RNA polymerase. The three proteins at 26 kD, 11.3 kD, and 5.8 kD, collectively known as a triple gene block, are presumed to mediate viral cell-to-cell movement. The 5A protein was produced in vivo; the 24-kD CP is located at the 3′-terminus. B, Schematic of infectious FoMV T-DNA clone pFoMV-wt. FoMV was cloned between CaMV 35S promoter with duplicated enhancers and a nopaline synthase (NOS) terminator of a T-DNA binary vector. pFoMV-wt has an additional 70 adenosine residues after the viral 3′-terminus. C, Schematic of the FoMV VIGS vector, pFoMV-sg, is a pFoMV-wt derivative that contains a direct repeat of the 300-bp putative FoMV CP subgenomic promoter (GenBank accession: EF630359, nt 5202–5371), shown by arrows, and multiple cloning sites (HpaI, MluI, XhoI, and Ascl) between CP subgenomic promoters. Right- and left-pointing blue arrows indicate sense and anti-sense orientation of foreign fragments, and the predicted structure of RNA transcript is presented on the right side. D, Symptom development and RT-PCR detection of viral RNA showed that pFoMV-wt and pFoMV-sg are infectious in barley. Bar = 1 cm.
infectivity of pFoMV-wt, *N. benthamiana* plants were infiltrated with *Agrobacterium*-carrying pFoMV-wt. RT-PCR showed the presence of FoMV in systemic leaves at 10 d post-infiltration (dpi). Barley plants inoculated with saps of FoMV-infected *N. benthamiana* leaves, but not those with saps of mock-inoculated healthy plants, developed a mild mosaic symptom at 10 dpi. These results demonstrate that the cloned FoMV is infectious and FoMV-infected *N. benthamiana* leaves are an excellent source of FoMV for secondary viral infections in cereals such as barley. Subsequently, all plants of different monocot species were infected through mechanic inoculation with saps of FoMV or the derivatives-infected *N. benthamiana* leaves.

FoMV, similar to PVX, belongs to *Potexvirus*, and PVX has successfully been used for VIGS. We thus employed a similar strategy for construction of the PVX VIGS vector (Ruiz et al., 1998) to generate our FoMV vector. The resulting vector pFoMV-sg is a pFoMV-wt derivative that contains a direct repeat of the 170-bp putative FoMV CP subgenomic promoter (SGP, which covers the entire 5A sequence) and multiple cloning sites, including *HpaI*, *MluI*, *XhoI*, and *AscI* between duplicated CP SGP. The modified full-length FoMV cDNA in pFoMV-sg is under control of a CaMV 35S promoter with duplicated enhancers and an NOS terminator (Fig. 1C). When barley plants were inoculated with saps of pFoMV-sg-infected *N. benthamiana* leaves, the uninoculated systemic leaves of barley plants developed very mild mosaic symptoms at 15 dpi (Fig. 1D). Recombinant FoMV-sg genomic RNA was readily detected by RT-PCR (Fig. 1D).

**FoMV-Based VIGS in Barley**

To test whether pFoMV-sg can be used to induce gene silencing in barley, a 200-bp barley *phytoene desaturase* (*HvPDS*) DNA fragment was cloned into pFoMV-sg to generate FoMV-HvPDS200. It has been reported that VIGS of *PDS* resulted in a photobleached phenotype (Kumagai et al., 1995; Liu et al., 2002a). Two-leaf-stage barley plants were inoculated with saps of *N. benthamiana* leaves agroinfiltrated with FoMV-HvPDS200 or empty vector pFoMV-sg. We detected the presence of FoMV in systemic leaves of all infected barley plants. However, we only observed very mild *PDS*-silencing bleached phenotypes in FoMV-HvPDS200-barley plants, even at 30 dpi.

To improve VIGS efficiency, we inserted inverted-repeats into pFoMV-sg, because VIGS can be enhanced through viral-based production of inverted-repeats (Lacomme et al., 2003). We first silenced two barley genes, *HvPDS* and *magnesium chelatase* (*HvChlH*), using this approach. *ChlH* encodes the H subunit of magnesium chelatase that is required for chlorophyll production (Shen et al., 2006). For this purpose, we cloned 60-bp *HvPDS* and 55-bp *HvChlH* of inverted-repeat fragments into pFoMV-sg to generate FoMV-HvPDS and FoMV-HvChlH. Two-leaf-stage barley plants were inoculated with saps of *N. benthamiana* leaves agroinfiltrated with FoMV-HvPDS, FoMV-HvChlH, or pFoMV-sg. As expected, pFoMV-sg-infected barley plants showed very mild chlorotic symptoms, usually occurring in the first and second leaves above the inoculated leaves. However, all barley plants infected with either FoMV-HvPDS or FoMV-HvChlH exhibited longitudinal white patches and streaks, and developed the strongest photobleached or chlorosis VIGS phenotype in leaf 3, numbered from the bottom of the plant at 21 dpi (Fig. 2A). Real-time RT-PCR showed that the mRNA level of *PDS* and *ChlH* in phenotypic tissues was reduced by approximately 90% and 75% in barley plants infected with FoMV-HvPDS and FoMV-HvChlH compared to plants infected with pFoMV-sg, respectively (Fig. 2B). These results clearly suggest that pFoMV-sg containing inverted-repeats of host genes can induce effective silencing in barley.

**FoMV-Based VIGS in Foxtail Millet**

Foxtail millet is an important grain crop widely grown in arid regions and it is a model C$_4$ grass plant. To test whether FoMV VIGS vector is able to induce gene silencing in foxtail millet, we silenced three foxtail millet genes *SIPDS*, *Cloroplastos alterados*1...
(SiCLA1), and IspH. CLA1 encodes 1-deoxy-D-xylulose-5-P synthase (Mandel et al., 1996; Estévez et al., 2001; Tang et al., 2010) and IspH is for an isopentenyl/dimethylallyldiphosphate synthase involved in isoprenoid biosynthesis. Their null mutations or VIGS resulted in albino phenotypes (Rohdich et al., 2002; Hsieh and Goodman, 2005; Yin et al., 2015). 60-bp inverted-repeat fragments targeting SiPDS, SiCLA1, and IspH (FoMV-SiPDS, FoMV-SiCLA1, and FoMV-IspH), respectively. Three-leaf stage foxtail millet plants were inoculated with saps of N. benthamiana leaves agroinfiltrated with pFoMV-sg or its derivatives. Foxtail millet plants infected with pFoMV-sg exhibited mild chlorotic symptoms. However, these plants infected with FoMV-SiPDS, FoMV-SiCLA1, or FoMV-IspH began to develop a mottled photobleached or albino phenotype in new emerging leaves at 10 dpi to 12 dpi. All plants developed larger areas of photobleached phenotype in the newly emerging leaves (the seventh leaf, numbered from the bottom of the plant) at 24 dpi (Fig. 3); in all the cases, the silencing phenotype was maintained and appeared in upper new emerging leaves for up to 60 d (55 dpi in Fig. 3E). Moreover, we performed real-time RT-PCR to determine mRNA levels of SiPDS, SiCLA1, and IspH in the silenced leaves. The mRNA levels of SiPDS, SiCLA1, and IspH were, respectively, decreased by approximately 66 to 74%, 55 to 76%, and 56 to 73% (Fig. 3) compared with that in FoMV-sg-infected control plants. These results suggest that our FoMV-based VIGS vector can be used to silence genes in foxtail millet.

FoMV-Based VIGS in Wheat

We also tested whether FoMV is able to induce gene silencing in wheat. To achieve this, we first identified two wheat marker genes, TaPDS and TaCLA1. Then we cloned 60-bp TaPDS and TaCLA1 of inverted-repeat fragments into pFoMV-sg to generate FoMV-TaPDS and FoMV-TaCLA1, respectively. Two-leaf-stage wheat plants were inoculated with saps of N. benthamiana leaves agroinfiltrated with pFoMV-sg or its derivatives. All wheat plants infected with FoMV-TaPDS or FoMV-TaCLA1 usually appeared as small white streaks in leaf 3, numbered from the bottom of the plant at 20 dpi; the white patches and streaks or albinism fully expanded in leaf 4 at 35 dpi (Fig. 4A). As plants grew, the bleached patches or albinism spread to the newly emerging leaves. However, no FoMV-sg control plants showed any silencing phenotype. To confirm the silencing of TaPDS and TaCLA1 genes at the molecular level, we performed real-time RT-PCR. As shown in Figure 4B, mRNA levels of TaPDS and TaCLA1 in the silenced leaves were strikingly reduced compared to FoMV-sg–infected plants. Taken together, our results clearly suggest that FoMV can induce efficient gene silencing in wheat.
VIGS. First, FoMV is a rod-shaped virus, and has a single, positive-sense single-stranded RNA genome (Bancroft et al., 1991). A VIGS vector-based single viral genome is easy to handle and is a model system for understanding functional genomics because severe viral disease symptoms may complicate VIGS phenotypes in testing plants (Ramanna et al., 2013). Third, FoMV can infect a broad range of hosts among the grasses as well as dicot species (Short and Davies, 1987). Thus, our pFoMV-sg VIGS vector should have the potential for VIGS in many monocot and dicot plants although we only tested its usefulness in barley, wheat, and foxtail millet. Fourth, FoMV-sg is highly infectious, and almost 90% of tested plants induced obvious silencing phenotypes. VIGS phenotypes caused by FoMV-sg vectors were stable and persisted for at least two months. In addition, FoMV can penetrate growing points in monocot plants. It may be applied to silence genes and analyze their biological relevance in plant reproductive tissues.

PVX is the type member of the genus Potexvirus and has successfully been developed as both expression and VIGS vectors by including an additional duplicated copy of CP subgenomic promoter in the viral genome (Ruiz et al., 1998). Because FoMV is also a member of Potexvirus, we adopted a similar strategy to generate vector pFoMV-sg, which contains a duplicated copy of the 170-bp putative FoMV CP SGP and a multiple cloning site. Surprisingly, delivery of an anti-sense RNA fragment of HvPDS by pFoMV-sg did not cause any obvious silencing phenotype in barley. To trigger effective VIGS, viral replication should be high enough to produce a threshold level of silencing-trigger RNA molecules (Lacomme et al., 2003). Expression of a hairpin-loop dsRNA structure generated a stronger silencing than expression of the corresponding sense or anti-sense cDNA (Waterhouse et al., 1998; Smith et al., 2000; De Buck et al., 2001). It has been reported that insertion of direct inverted-repeat sequences of target genes that could fold as dsRNA hairpin structures strongly enhance VIGS from Tobacco mosaic virus, BSMV (Lacomme et al., 2003), and Turnip yellow mosaic virus (Pflieger et al., 2008). This is true for pFoMV-sg–mediated VIGS. Expression of inverted-repeats of PDS but not an anti-sense fragment of PDS from FoMV-sg triggered a clear photobleached phenotype in barley (Fig. 2). Similarly, FoMV-based expression of inverted-repeats of ChlH in barley (Fig. 2), of PDS and CLA1 in foxtail millet and wheat plants, of IspH in foxtail millet plants, induced efficient silencing. Moreover, silencing phenotype persisted up to 60 d (Fig. 3). These results support that an insertion of inverted-repeat sequences into the viral genome can enhance the efficiency of VIGS.

Foxtail millet is a member of the Poaceae grasses, and is a model system for understanding functional genomics and photosynthesis in grain crops and C₄ plants (Wang et al., 2010). Foxtail millet is also a close relative of switchgrass (Panicum virgatum), an important biofuel crop. Recently, the whole genome sequence of foxtail native and adaptive hosts (Robertson et al., 2000). Indeed, our pFoMV-sg VIGS vector generally induces very mild mosaic symptoms in emerging leaves of barley, wheat, and foxtail millet plants. This makes the pFoMV-sg vector extremely useful for functional genomics because severe viral disease symptoms may complicate VIGS phenotypes in testing plants (Ramanna et al., 2013).
millet has been published (Bennetzen et al., 2012; Zhang et al., 2012). Therefore this first report of an efficient VIGS system based on pFoMV-sg provides a timely toolbox for high-throughput assays of gene functions in foxtail millet and many other economically important monocot crops.

CONCLUSIONS

Our FoMV-sg vector is able to trigger efficient gene silencing in barley, wheat, and foxtail millet. FoMV-sg–mediated VIGS could possess a great potential for functional genomics in a variety of monocot crops.

MATERIALS AND METHODS

Plant Material and Growth Conditions

Tobacco (Nicotiana benthamiana), barley (Hordeum vulgare L.; Ingrid), wheat (Triticum aestivum; Zhounan30), and foxtail millet (Setaria italica; Yugul1) plants were grown in pots at 22–24°C, in a growth chamber under a 16-h light/8-h dark cycle.

Construction of Foxtail mosaic virus-Derived Vectors

The full-length Foxtail mosaic virus (FoMV) cDNA was PCR-amplified using FoMV cDNA clone P9 (Robertson et al., 2000) as a template with primer pairs P1-F/P1-R, and cloned into StuI/Xho-digested pYL44, a pBHN19-based T-vector containing a duplicated CaMV 35S promoter and nitric oxide synthase (NOS) terminator (Li et al., 2002b), to generate pFoMV-wt. Further, DNA fragment 1 containing FoMV nt 4258–5371 (GenBank accession: EF630359) and DNA fragment 2 containing FoMV nt 5202–6008 were PCR-amplified using primer pairs P2-F/P2-R and P3-F/P3-R, respectively. Sall/AscI-digested DNA fragment 1 and AscI/AvrI-digested DNA fragment 2 were inserted into Sall/AvrII-digested pFoMV-wt to generate pFoMV-sg. The resulting vector pFoMV-sg is a pFoMV-wt derivative that contains a duplicated copy of 170-bp putative FoMV coat-protein subgenomic promoter and multiple cloning sites between the duplicated coat-protein subgenomic promoters.

Barley PDS fragment (corresponding to nt 345–546 of HvPDS; GenBank accession: AY062039) was PCR-amplified using barley cDNA as template and primer pairs P4-F/P4-R, and cloned into pFoMV-sg to generate FoMV-HvPDS200 for virus-induced gene silencing.

600-bp DNA fragments of barley PDS gene (corresponding to nt 1089–1148 of HvPDS) was RT-PCR-amplified using the primers PS-F/PS-R; 55-bp DNA fragments of barley ChlH gene (corresponding to nt 24–78 of HvChlH; GenBank accession: U26545) was RT-PCR-amplified using the primers Ph-F/Ph-R, and then PCR products were digested with SpeI. The inverted-repeats of HvChlH/PDS were obtained by self-ligation of SpeI-cut PCR products, and then further cloned into the AscI site of pFoMV-sg to generate FoMV-HvPDS or FoMV-HvCHlH.

The fragments of wheat PDS and CLA1 genes were RT-PCR-amplified using a primer pair P7-F/P7-R for a PDS gene (corresponding to nt 423–482 of TaPDS; GenBank accession: F517553), and P8-F/P8-R for a CLA1 gene (corresponding to nt 1428–1487 of TaCLA; GenBank accession: AK335035). PCR products were digested with SpeI and inverted-repeats of TaPDS or TaCLA1 were obtained by self-ligation of SpeI-cut PCR products, and then cloned into the AscI site of pFoMV-sg to generate FoMV-TaPDS and FoMV-TaCLA1, respectively.

The fragments of foxtail millet PDS, CLA1, and IsplH genes were RT-PCR-amplified using primer pairs P7-F/P7-R for PDS (corresponding to nt 558–617 of SiPDS; GenBank accession: XM004985406), and P9-F/P9-R for CLA1 (corresponding to nt 1511–1570 of SiCLA; GenBank accession: XM004962054), and P10-F/P10-R for IsplH (corresponding to nt 364–423 of SiSipH; GenBank accession: XM004981826), and then PCR products were digested with SpeI. Inverted-repeats of SiPDS, SiCLA1, and SiSipH were obtained by self-ligation of SpeI-cut PCR products, and then cloned into the AscI site of pFoMV-sg to generate FoMV-SiPDS, FoMV-SiCLA1, and FoMV-SiSipH, respectively.

Agroinfiltration of N. benthamiana and Viral Inoculation of Cereals

FoMV-based VIGS was performed by using N. benthamiana as an intermediate host as described (Yu et al., 2003; Yuan et al., 2011). FoMV-based constructs were introduced into Agrobacterium tumefaciens strain GV3101. Agrobacterium was grown in Luria-Berti media containing rifampicin (30 mg l−1) and kanamycin (50 mg l−1) at 28°C for 2 d. Bacterial cells were harvested at 2500 × g for 5 min, resuspended in infiltration medium (10 mM MgCl2, 10 mM MES [2-(N-morpholino)ethanesulfonic acid], and 0.2 mM aceto-syringone) and adjusted to OD600 = 1.0, and kept at room temperature for 4 h.

Agrobacterium were infiltrated into the fully expanded leaves of N. benthamiana using a 1-mL needleless syringe. The infiltrated leaves were harvested for 7–10 d post-infiltration, ground in 20-mM Na-P buffer (pH 7.2), and saps were mechanically inoculated onto leaves of the two-leaf stages of barley and wheat, and three-leaf stages of foxtail millet plants. Each experiment was repeated at least three times, with 20 plants for each viral construct in each experiment.

RNA Extraction and Reverse Transcription-PCR

Total RNA was extracted from plant leaf tissues using the Trizol reagent (Tiangen Biotech, Beijing, China), and then treated with RNase-free DNAse I (Sigma-Aldrich, St. Louis, MO) to eliminate genomic DNA (Sigma-Aldrich). First-strand cDNA for RT-PCR was synthesized from 2 μg of RNA by RevertAid Reverse Transcriptase, with Oligo(dT), dNTP, and Ribonuclease inhibitors following the instruction of the manufacturer (Tiangen).

Real-time RT-PCR was performed using Power SYBR Green PCR Master Mix (Applied Biosystems, Warrington, Cheshire, UK) following the manufacturer’s instructions. Gene-specific primers were designed to prime outside the region targeted for silencing. For barley PDS or ChlH, primer pairs P11-F/P11-R or P12-F/P12-R were used. Barley 18S rRNA gene (GenBank accession: AK251731) was used as an internal control using primer pairs P18-F/P18-R. For wheat PDS or CLA1, primer pairs P13-F/P13-R or P14-F/P14-R were used, respectively. Wheat 18S rRNA gene (GenBank accession: AY049040) was used as an internal control using primer pairs P18-F/P19-R. For foxtail millet PDS, CLA1, or IsplH, primer pairs P15-F/P15-R, P16-F/P16-R, or P17-F/P17-R were used, respectively. Foxtail millet 18S rRNA gene (GenBank accession: KC996746) was used as an internal control using primer pairs P20-F/P20-R.

Accession Numbers

Sequence data from this article can be found in the EMBL/GenBank data libraries under the following accession numbers: FoMV (EF630359); HvPDS (AY062039); HvChlH (U26545); TaPDS (F517553); TaCLA (AK335035); SiPDS (XM004985406); SiSipH (XM004962054); SiCLA1 (XM004981826); barley 18S rRNA (AK251731); Wheat 18S rRNA (AY049040); Foxtail millet 18S rRNA (KC996746).

Supplemental Data

The following supplemental materials are available.

Supplemental Table S1. List of primers used in this work.

ACKNOWLEDGMENTS

We thank Dr. Nancy Robertson (USDA, Palmer, Alaska) for kindly providing FoMV cDNA clone P9, and Professors Qianhua Shen and Zhiyong Liu for kindly providing plant seeds.

Received January 7, 2016; accepted May 24, 2016; published May 25, 2016.

LITERATURE CITED


