Establishment of a Simple and Rapid Gene Delivery System for Cucurbits by Using Engineered Zucchini Yellow Mosaic Virus

Minji Kang1†, Jang Kyun Seo2†, Hoseong Choi1, Hong Soo Choi2 and Kook Hyung Kim1,3*
1Department of Agricultural Biotechnology and Research Institute of Agriculture and Life Sciences, Seoul National University, Seoul 151-921, Korea
2Crop Protection Division, National Academy of Agricultural Science, Wanju 565-852, Korea
3Plant Genomics and Breeding Institute, Seoul National University, Seoul 151-921, Korea
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The infectious full-length cDNA clone of zucchini yellow mosaic virus (ZYMV) isolate PA (pZYMV-PA), which was isolated from pumpkin, was constructed by utilizing viral transcription and processing signals to produce infectious in vivo transcripts. Simple rub-inoculation of plasmid DNAs of pZYMV-PA was successful to cause infection of zucchini plants (Cucurbita pepo L.). We further engineered this infectious cDNA clone of ZYMV as a viral vector for systemic expression of heterologous proteins in cucurbits. We successfully expressed two reporter genes including gfp and bar in zucchini plants by simple rub-inoculation of plasmid DNAs of the ZYMV-based expression constructs. Our method of the ZYMV-based viral vector in association with the simple rub-inoculation provides an easy and rapid approach for introduction and evaluation of heterologous genes in cucurbits.

Keywords: gene delivery, infectious clone, ZYMV

Zucchini yellow mosaic virus (ZYMV), which belongs to the genus Potyvirus, is a flexuous rod-shaped virus of about 750 nm in length and has a genome of a plus-sense single-stranded RNA of 9.5 kb (Gal-On et al., 1991). ZYMV is one of the major pathogens in cucurbitaceous crops worldwide (Lecoq et al. 2009; Simmons et al., 2013). ZYMV causes severe stunting and yellowing symptoms on leaves, and stream and deformation in fruits of cucurbitaceous crops (Kwon et al., 2005). ZYMV possesses a covalently linked viral protein (VPg) at the 5′-terminus and a poly (A) tail at the 3′-terminus (Arazi et al., 2001). The genome of ZYMV encodes a large polyprotein and a putative pretty interesting potyviral protein (PIPO) at the N terminus of the P3 region (Chung et al., 2008). The large polyprotein is cleaved by self-encoded proteases into at least ten functional proteins, including P1, helper component protease (HC-Pro), P3, 6K1, cylindrical inclusion protein (CI), 6K2, VPg, NIa-Pro, Nlb, and coat protein (CP) (Arazi et al., 2001; Kwon et al., 2005).

Cucurbitaceous crops, which belong to the family Cucurbitaceae, are an important part of diverse nutrition diet worldwide. The cucurbitaceous crops mostly cultivated in Korea include watermelon, cucumber, squash, and oriental melon (Kwon et al., 2005). However, research tools capable of introducing and evaluating valuable genes into cucurbitaceous crops are very limited. The plant virus-based vectors have been developed to express various foreign proteins in a wide range of host plants, since plant viruses accumulate high levels of virus-encoded proteins in infected plants (Gleba et al., 2014; Hefferon, 2012). The polyprotein expression system of potyviruses is advantageous to express foreign proteins as a part of viral proteins (Arazi et al., 2001; Seo et al., 2009). So far, various potyviruses, including soybean mosaic virus, turnip mosaic virus, tobacco etch virus, potato virus A, have been already engineered as vectors for expression of heterologous genes (Beauchemin et al., 2005; Dolja et al., 1992; Kelloniemi et al., 2008; Seo et al., 2009). Commonly, two sites between P1 and HC-Pro cistrons and between Nlb and CP cistrons have been used to insert target genes in potyvirus-based vectors (Beauchemin et al., 2005; Kelloniemi et al., 2008).
In this study, we constructed the infectious full-length cDNA clone of ZYMV and engineered it as a viral vector for systemic expression of heterologous proteins in cucurbits. Upon simple rub-inoculation of plasmid DNAs of the ZYMV-based viral vector, we successfully expressed two reporter genes including gfp (the green fluorescence protein gene) and bar (the herbicide-resistance gene) in zucchini plants.

The ZYMV isolate PA (ZYMV-PA) obtained from pumpkins at Andong city (Kwon et al., 2005) was propagated in Cucumis melo L. (oriental melon) in a greenhouse and used for construction of infectious full-length cDNA clone of ZYMV. Total RNAs were extracted from the C. melo L. leaves infected with ZYMV-PA using the TRI Reagent (MRC, USA) according to the protocols provided by the manufacturer. The extracted total RNA was used for cDNA synthesis of ZYMV-PA. The full-length cDNA clone of ZYMV-PA was constructed by assembling three partial amplified fragments of the ZYMV-PA cDNA. The first fragment including the ZYMV nucleotides from 1 to 1,807 (ZYMV 1–1807) was amplified by RT-PCR using specific primers (ZYPA-F, 5′-AAAATTGAAACAAATCACAAAGACTATAAG-3′ and ZYPA-1807-BamHI-R, 5′-ACGGATCCAACTTACGGGTACCATTT-3′). The resulting amplicon was digested with BamHI and inserted into between the StuI and BamHI sites of pCass-RZ, which is a modified binary vector (de Wispelaere and Rao, 2009). The resulting clone was named pZYMVF1. The second fragment (ZYMV 1788–5938) was amplified by RT-PCR using specific primers (ZYPA-1788-F, 5′-CAAATGGTACCCGTAAGTTGG-3′ and ZYPA-5938-R, 5′-CCAGTGAGAGBAMHI-3′). The resulting product was digested with KpnI and BamHI and ligated with pZYMVF1, which was opened with KpnI and BamHI. The resulting clone was named pZYMVF1F2. For amplification of the last fragment (ZYMV 5911–9593), cDNA was synthesized using a tagged oligo(dT) primer (ZYPA-dT-BamHI-RT, 5′-GCTGGGATCCGTGGTAAATTTTTTTTTTTTTTTTTTTT-3′) to hybridize to the ZYMV poly(A) tail. The cDNA was then amplified by PCR using specific primers (ZYPA-5911-F, 5′-CATCAGATTTGTGGATCCTCTC-3′ and ZYPA-BamHI-R, 5′-GCTGGGATCCGTGGTAA-3′). The resulting product was digested with BamHI and ligated with pZYMVF1F2, which was opened with BamHI. This final clone designated as pZYMV-PA contained full-length cDNA of ZYMV-PA with 25 adenines at the 3′ end (Fig. 1).

To examine the infectivity of the cloned ZYMV-PA, purified plasmid DNAs of pZYMV-PA was inoculated onto the leaves of zucchini plants (C. pepo L.) by direct rub-inoculation with carborundum. Ten μg of pZYMV-PA prepared using the Plasmid Maxi Kit (QIAGEN, USA) was mixed with inoculation buffer (50 mM potassium phosphate, pH 7.5) to a total volume of 80 μl for the inoculation of two leaves per plant. At 12 days post-inoculation (dpi), zucchini plants inoculated with pZYMV-PA showed symptoms of yellowing, deformation, and mosaic, similar to those caused by the original ZYMV-PA (Fig. 2A). To confirm ZYMV infection of the inoculated plants, total RNAs extracted from the upper un-inoculated leaves were subjected to RT-PCR using the ZYMV-specific primers (5′-CTTTGAAACAGAAGACTATAAG-3′ and 5′-GCCGTAAAATATTAGAATTAC-3′) that detect the CP region of ZYMV (Fig. 2B). A band of the predicted size (approximately 837 bp) was observed in all inoculated samples (Fig. 2C). Therefore, the results demonstrated that pZYMV-PA is fully infectious.

To develop pZYMV-PA as a viral vector for expres-

![Fig. 1. Schematic representation of construction of the infectious full-length cDNA clone of zucchini yellow mosaic virus. The pCass-RZ vector contains, in sequential order, a left border of T-DNA (LB), a double CaMV 35S promoter (35S X 2), a cis-cleaving ribozyme sequence (RZ), a 35S terminator (Ter), and a right border of T-DNA (RB). The restriction enzyme cleavage sites used to construct the full-length clone are shown in gray boxes.](image-url)
Fig. 2. Infection of zucchini plants upon mechanical rub-inoculation of plasmid DNAs of pZYMV-PA. (A) Wild-type (wt) virus of ZYMV-PA (ZYMV-PA wt) and the cloned ZYMV-PA (pZYMV-PA) were mechanically inoculated on the leaves of zucchini plants. At 12 days post inoculation (dpi), ZYMV-PA wt (a2) and pZYMV-PA (a3) caused same symptoms of yellowing, leaf deformation (arrowheads) and mosaic. (B) The primer pair (the marked arrows) spanning the CP region was used for RT-PCR detection of ZYMV. (C) Infectivity of pZYMV-PA was confirmed by RT-PCR detection of ZYMV replication. Arrow indicates PCR bands of the predicted size.

Fig. 3. Schematic representation of ZYMV-based viral vector constructs. (A) ZYMV genome organization and introduction of a gene insertion cassette between the NIb and CP cistrons. The shaded box indicates the gene insertion cassette containing multiple cloning sites (MCS: ApaI, SpeI, and SalI) and a Nla-Pro cleavage site (/S---DTVMLQ/). (B) Diagrammatic representation of insertion of the gfp and bar genes into the ZYMV-based vector. The gfp and bar genes were inframe-inserted into the pZYMV-MCS vector utilizing the SpeI site.

Expression of heterologous genes in cucurbit plants, cloning sites (ApaI, SpeI and SalI) and an additional Nla-Pro cleavage site (/S---DTVMLQ/) were inserted into the polyprotein ORF between the NIb and CP cistrons (Fig. 3A). To this end, two SacI sites in pZYMV-PA were utilized (Fig. 3A). The NIb region downstream
of the first SacI site was amplified using a primer pair (ZYPA-Nlb-F, 5'-CCTGTGGAATGGATCTTTAA
AGGCTGAGCTCAGAC-3' and ZYPA-Nlb(Nla_MCS)-R,
5'-GACTGTGCACTAATGTTGCCCGACTGGAG
CATCACAGTGCCTCTCTGGTCAAGAAATG-3'; the cloning sites are underlined) and subcloned into
pGEM-T Easy Vector (Promega, USA). This intermediate clone was named pPA-MCS1-T. Next, the CP
region upstream of the second SacI site was amplified using a primer pair (ZYPA-(MCS_Nla)CP-F,
5'-AGTCGTCGACACTGTGATGCTCCAGTCGG
GCACCTCAGAAAATGTGGG-3' and pCass-SalI-
R, 5'-GACTGTCGACTGATTTCAGCGTACCGAG
TTCGAGCTC-3') and digested with SalI. The resulting
fragment was inserted into pPA-MCS1-T that was opened with SalI. The resulting clone was named pPA-MCS2-T. Then, the ZYMV Nlb/CP region harboring the cloning sites and a Nla-Pro cleavage site was cut out from pPA-MCS2-T by digestion with SalI and inserted into pZYMV-PA, which was opened with SalI. This final clone was named pZYMV-MCS (Fig. 3A). The infectivity of pZYMV-MCS was investigated by mechanical inoculation of zucchini plants using purified plasmid DNAs of pZYMV-MCS as described above. At 12 dpi, zucchini plants inoculated with pZYMV-MCS showed similar disease symptoms when compared with those induced by
pZYMV-PA (Fig. 4A). Infection of the inoculated plants with ZYMV conformed by RT-PCR using the primer pair (5'-CCATCAAGCAATGTTGGTTGATTG-3' and 5'-CGACTCCCATCTTGATTGAC-3') that detects the
Nlb/CP region (Fig. 4B and C).

To evaluate the expression of foreign genes from the ZYMV vector, two reporter genes, gfp and bar, were amplified by PCR using appropriate primers (GFP-SpeI-F, 5'-GACTGTGCACTAATGTTGCCCGACTGGAG
CATCACAGTGCCTCTCTGGTCAAGAAATG-3' and GFP-SpeI-R, 5'-GACTGTGCACTAATGTTGCCCGACTGGAG
CATCACAGTGCCTCTCTGGTCAAGAAATG-3' and GFP-SpeI-R, 5'-GACTGTGCACTAATGTTGCCCGACTGGAG
CATCACAGTGCCTCTCTGGTCAAGAAATG-3') and inserted into pZYMV-MCS utilizing the SpeI cloning site. The resulting constructs were designated as pZYMV-GFP and pZYMV-Bar (Fig. 3B). The plasmid DNAs of pZYMV-GFP and -Bar were inoculated onto the cotyledons of zucchini seedlings by direct rub-inoculation. At 12 dpi, both of the constructs induced typical yellowing and mosaic symptoms of ZYMV (data not shown). In addition, strong signals of green fluorescence emitted by GFP were observed on the systemic leaves of the plants inoculated with pZYMV-GFP when the plants were visualized under illumination with a hand-held UV-light source (Dark Reader Hand Lamp HL28T; Clare Chemical Research, Dolores, CO) (Fig. 5). At 12 dpi, the zucchini plants infected by
Fig. 5. Detection of fluorescence signals emitted by GFP under UV light on leaves of zucchini plants inoculated with pZYMV-GFP. (A) Plasmid DNAs of pZYMV-PA and pZYMV-GFP were mechanically inoculated on the leaves of zucchini plants. At 7 dpi, systemic leaves of zucchini plants inoculated with pZYMV-GFP (a3) showed intense fluorescence. No fluorescence was detected on leaves of healthy (a1) and pZYMV-PA-inoculated plants (a2). (B) The closed view of the systemic leaves photographed under UV light at 12 dpi.

Fig. 6. Acquiring herbicide resistance of zucchini plants infected by pZYMV-Bar. The zucchini plants were treated with a non-selective herbicide (0.54% glufosinate-ammonium (w/v)) at 12 day post inoculation of the indicated inoculums. (A) The representative photographs of the zucchini plants inoculated with pZYMV-PA (a2) and pZYMV-Bar (a3) at 7 days after herbicide treatment. (B) Magnified images of the zucchini plants inoculated with pZYMV-PA (b1) and pZYMV-Bar (b2). Asterisks indicate leaves treated with herbicide and arrowheads indicate upper leaves that grew after herbicide treatment.
pZYMV-Bar were treated with a non-selective herbicide (0.54% glufosinate-ammonium (w/v)). At 7 days after herbicide treatment, while the healthy and pZYMV-PA-inoculated plants were significantly damaged, the plants inoculated with pZYMV-Bar were still healthy, indicating the expression of *bar* gene from the ZYMV viral vector is sufficient to confer herbicide resistance in zucchini plants (Fig. 6).

To evaluate the stability of the foreign gene insertion in the ZYMV viral vector, viruses obtained from the plant infected with either pZYMV-GFP or -Bar were passed three times from plant to plant by mechanical sap-inoculation. Total RNA was extracted from upper systemic leaves of each individual plant and analyzed for stable gene insertion in the viral genome by RT-PCR using the primer pair (5′-CCATCAAGCAATGTTGGTTGAT-3′ and 5′-CGACTCCCATCTGTTGATCATT-3′) that detects the NIb/CP region (Fig. 7A). A unique band with the predicted size (approximately 1.5 kbp for *gfp* insertion and 1.3 kbp for *bar* insertion) was observed for every plants infected by either pZYMV-GFP or -Bar (Fig. 7B and C). The results suggest that foreign genes inserted in the ZYMV viral vector could be maintained stably during serial passages.

With recent advances in plant biotechnology, plants have been targeted to be engineered as a mass production system for recombinant proteins (Hefferon, 2012). The interest in using plants as a cost effective means for the production of valuable proteins has led to the development of various plant virus-based vectors (Gleba et al., 2014; Hefferon, 2012). Virus-mediated gene expression in plant is a superior system to the transgenic approach because plant viruses systemically infect plants and accumulate large quantities of viral proteins. Thus, various plant virus-based gene delivery vectors have been developed for the overproduction of recombinant proteins within a short period of time in plants (Gleba et al., 2014; Hefferon, 2012). Since the turnaround time for introduction of new traits into plants is short, the plant virus-based gene delivery system can facilitate large-scale analysis of gene functions based on gain-of-function studies.

In this study, we have developed the ZYMV-based gene delivery vector for efficient expression of heterologous proteins in cucurbit plants. When using potyvirus-based vectors, multiple foreign proteins can be expressed from one construct by utilizing the different genomic regions (for example, two sites between P1 and HC-Pro and NIb and CP cistrons for insertion of foreign genes (Beauchemin et al., 2005; Kelloniemi et al., 2008). A previous study has developed a bombardment-based gene expression system by engineering ZYMV as a viral vector (Arazi et al., 2001). However, in general, bombardment-based methods require expensive devices and materials and are less-efficient. On the other hand, we established a simple, easy and low cost method for inoculation of the ZYMV-based viral vector by direct rub-inoculation of purified plasmid DNAs of the vector. To this end, we utilized viral transcriptions and processing signals, including the CaMV 35S promoter, a self-cleaving ribozyme sequence and a poly(A) signal, to produce infectious *in vivo* transcripts resembling the native ZYMV genomic RNA. This DNA-mediated rub-inocula-
tion method has an advantage of not requiring expensive and complicated steps or special equipments to deliver a viral vector containing foreign genes. It is highly desirable to identify and evaluate valuable traits for improving productivity, environmental and disease resistance, and the commercial value of cucurbit plants. The availability of the ZYMV-based gene delivery system developed in this study will be helpful to begin rapid in planta screening and evaluation of valuable candidate genes.

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