A Transgenic Lettuce Line with Resistance to Both Lettuce Big-vein Associated Virus and Mirafiiori Lettuce Virus

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Abstract. The coat protein (CP) gene of lettuce big-vein associated virus (LBVaV) in sense or antisense orientation in a binary vector pBl121 was transformed via Agrobacterium tumefaciens (Smith and Towns.) Conn. mediated transformation into lettuce (Lactuca sativa L.) to generate LBVaV-resistant lettuce. Nineteen T1 lines were produced; five to 10 plants of each T1 line were inoculated with LBVaV using Olpidium brassicae (Wor.) Dang.; and LBVaV was not detected in eight plants derived from six lines. T1 seedlings from the eight plants were tested for LBVaV resistance, and one line (line A-2) with the CP gene in antisense orientation was resistant to LBVaV while the other lines were susceptible. The transgenic line A-2 was also resistant to mirafiiori lettuce virus (MiLV) and big-vein expressions regardless of the presence or absence of LBVaV.

Lettuce big-vein disease, first described in California (Jagger and Chandler, 1934), is a soilborne disease transmitted by the fungus Olpidium brassicae (Campbell and Grogan, 1964). Infected lettuce plants develop vein bandings, and leaves become ruffled and distorted. A virus with rod-shaped particles, named lettuce big-vein associated virus (Fauquet et al., 2005), previously named lettuce big-vein virus, was first found in lettuce with big-vein disease (Kuwata et al., 1983); this virus was believed to be a causal agent of big-vein disease for nearly two decades. However, a second virus, named mirafiiori lettuce virus, was recently isolated from lettuce showing big-vein symptoms (Roggero et al., 2000), and it was reported that MiLV but not LBVaV induced big-vein symptoms in lettuce (Lot et al., 2002). Although LBVaV is not regarded as a causal agent of big-vein disease, it is possible that LBVaV is related to MiLV infections and big-vein symptom expressions because LBVaV is frequently associated with big-vein disease.

One of the most efficient methods to control big-vein disease is the use of resistant cultivars (Ryder and Robinson, 1995). Resistant cultivars have been developed by a conventional breeding method. For example, the big-vein-resistant cultivars Thompson and Pacific were developed using several resistant sources (Ryder, 1981; Ryder and Robinson, 1991). However, cultivars released so far do not exhibit high levels of resistance because breeding sources with high resistance to the disease have not been found in lettuce (Bos and Huijberts, 1990; Ryder and Robinson, 1995).

It was suggested that the expression of viral gene sequences in transgenic plants might disrupt the viral life cycle (Sanford and Johnston, 1985), and virus-derived resistance in transgenic plants was first reported using the CP gene of tobacco mosaic virus (Powell-Abel et al., 1986). After these reports, there have been numerous attempts to generate virus resistance in transgenic plants through the expression of CP genes, viral replication genes, and other viral sequences (Baulcombe, 1996). Many of these attempts have been successful, and the use of viral sequences to produce virus-resistant plants is now routine.

Nucleotide sequences of LBVaV and MiLV were recently reported (Kawazu et al., 2003; Sasaya et al., 2002, 2004; van der Wilk et al., 2002), and it is now possible to transform lettuce with LBVaV or MiLV sequences to produce transgenic lettuce with resistance to LBVaV or MiLV. In this study we introduced the LBVaV CP gene into lettuce to make LBVaV-resistant lettuce, and a transgenic lettuce line was obtained that was resistant not only to LBVaV but also to MiLV, and therefore repressed symptom expressions.

Materials and Methods

Transformation of lettuce with the LBVaV CP gene. The β-glucuronidase (GUS) gene in pBl121 was replaced by the full coding sequence of the LBVaV CP gene (1.2 kb), and the plasmids with the LBVaV CP gene in antisense and sense orientations were referred to as pYK10 and pYK11, respectively. The binary vector pYK10 or pYK11 was transformed via A. tumefaciens-mediated transformation into “Cisco” lettuce by the leaf disc method (Curtis et al., 1995). Explants were regenerated on medium containing 100 mg-L−1 kanamycin. PCR was carried out to detect the LBVaV CP gene in transgenic plants using the following primers: 5′-GATGTGATATCTCCAAGACACGTAAG-
3´ (CaMV 35S promoter region) and 5´-CTCAT AAATAACGTC ATGCA TTACA-3´ (NOS terminator region).

**Virus inoculation and detection.** *Olpidium brassicae* was obtained from lettuce showing big-vein symptoms in Kagawa prefecture, Japan. It was confirmed to contain both LBVaV and MiLV when they were detected using Western blot analysis and double-antibody sandwich enzyme-linked immunosorbing assay (DAS-ELISA) in susceptible ‘Cisco’ plants 60 days after inoculation. The methods for virus detection are described below. *Olpidium brassicae* carrying MiLV alone was kindly provided by Dr. H. Koganezawara (Kaneko Seeds Co., Gunma, Japan). *Olpidium brassicae* were maintained in ‘Cisco’ plants and determined to be viruliferous when inoculated plants showed big-vein symptoms. Roots of ‘Cisco’ plants infected with *O. brassicae* containing both viruses or MiLV alone were used for inoculation. Five grams of roots were homogenized with 300 mL of deionized water using a juicer. They were then filtered with one-layer gauze, and 10 mL of homogenized roots were poured onto the base of each seedling. Inoculated plants were kept in a plant growth chamber at 20 °C during the day (14-h photoperiod) and at 15 °C at night. Two discs were punched from randomly selected leaves of each lettuce plant using the lid of a 2-mL microtube. Each lettuce sample was digested with 0.4 mL of PBS-T (20 mM Na-phosphate buffer, pH 7.2, 150 mM NaCl, 0.05% Tween-20) and centrifuged with 6700 g, for 1 min. The supernatant was used for Western blot analysis to detect LBVaV and for DAS-ELISA to detect MiLV. 

Protein extract (10 µL) was subjected to 10% SDS-polyacrylamide gel electrophoresis, and Western blot analysis for detection of LBVaV was conducted according to a standard procedure (Gallagher et al., 1993). Mouse serum against LBVaV diluted 1:3000 was used for the primary antibody, and the alkaline phosphatase-conjugated goat antibody (Bio-Rad Laboratories, Hercules, Calif.) diluted 1:10,000 was used for the secondary antibody. A sample was considered LBVaV-positive if the CF band was observed on X-ray film.

DAS-ELISA for detection of MiLV was performed essentially as described by Clark and Adams (1977). The antibody for coating and the antibody-alkaline phosphatase (AP) conjugate (Japan Plant Protection Association Corp., Tokyo) were diluted 1:500 and 1:1500, respectively. One milligram per milliliter p-nitrophenyl phosphate (NPP) was finally added into wells of microtiter plates and incubated at 37 °C for 70 min. A sample was considered MiLV-positive if the difference of the absorbance at 405 nm between the sample and uninoculated controls was more than 0.1.

**Southern hybridization.** Genomic DNA was isolated from lettuce leaves by the CTAB method as described by Roger and Bendich (1988). Approximately 10 µg of genomic DNA was digested with Eco RI, Eco RV or Hind III, and the resulting fragments were electrophoresed in 1.5% agarose gel and transferred to a Hybond-N+ membrane (GE Healthcare Bio-Sciences Corp., Piscataway, N.J.) by capillary blotting (Sambrook and Russell, 2001). The full coding sequence of the LBVaV CP gene (1.2 kb) or the npt II gene (0.8 kb) was labeled with an alkaline phosphatase using the AlkPhos Direct Labelling and Detection System (GE Healthcare Bio-Sciences). The hybridization was performed according to the manufacturer’s recommendations.

**Results and Discussion**

**Production of transgenic lettuce.** In order to produce transgenic lettuce, 6200 leaf segments were inoculated with *Agrobacterium tumefaciens* carrying the LBVaV CP gene in antisense orientation on the binary plasmid pYK10, and 30 independent plant lines (T1 generation) were regenerated on medium containing 100 mg L−1 kanamycin. Seventeen of 30 plants were shown to be CP gene-positive by PCR analysis (data not shown), and self-pollinated. Seven of 17 plants produced seeds (T2 generation) while 10 plants did not produce seeds or died before flowering.

Another 6200 leaf segments were inoculated with *A. tumefaciens* carrying the LBVaV CP gene in sense orientation on the binary plasmid pYK11, and 36 independent plant lines were regenerated on medium containing 100 mg L−1 kanamycin. Twenty-two of 36 plants were shown to be CP gene-positive by PCR analysis (data not shown), and 12 of 22 plants produced seeds (T2 generation) while 10 plants did not produce seeds or died before flowering.

**Selection of transgenic plants for LBVaV resistance.** In order to screen resistant lines, five to 10 plants of each *T1* line (seven lines with the LBVaV CP gene in antisense orientation and 12 lines with the LBVaV CP gene in sense orientation) were inoculated with LBVaV using *O. brassicae*. LBVaV was detected using Western blot analysis. Five plants from three lines with the LBVaV CP gene in antisense orientation and three plants from three lines with the LBVaV CP gene in sense orientation were LBVaV-negative 56 days after inoculation (data not shown), and they were self-pollinated for resistance test in *T2* generation. All plants of line A-2 were LBVaV-negative while almost all plants of the other lines were LBVaV-positive 31 days after inoculation (Table 1). The presence of the CP gene in line A-2 was checked by PCR analysis, and all plants tested were CP gene-positive (Data not shown). Southern blot analysis was also performed on genomic DNA from eight plants of line A-2 (Fig. 1). The result shows the presence of one copy of the LBVaV CP gene (Panel B) and one copy of the *npt II* gene (Panel C) per genome. Two bands were detected when genomic DNA digested with *Eco RI* was hybridized with the LBVaV CP probe because of the presence of the *Eco RI* site in the LBVaV CP gene (Lane 1 in Panel B).

**Analysis of transgenic line A-2.** We examined the resistance of line A-2 not only to LBVaV but also to MiLV and symptom expressions (Figs. 2–3). All line A-2 plants were LBVaV-negative, MiLV-negative, and without symptoms while all ‘Cisco’ plants were LBVaV-positive, MiLV-positive and symptomatic 40 days after inoculation. We saw no significant difference in growth between line A-2 and ‘Cisco’ plants. As line A-2 has the LBVaV CP gene in antisense orientation, the transgene is untranslatable, suggest-

Table 1. Percentages of lettuce big-vein associated virus (LBVaV)-positive lettuce plants of *T2* generation inoculated with LBVaV using *Olpidium brassicae*.

| Line | Orientation of the coat protein gene | LBVaV-positive |
|------|-------------------------------------|----------------|
| A-1  | antisense                           | 94% (15/16)    |
| A-2  | antisense                           | 0% (0/16)      |
| A-3  | antisense                           | 81% (13/16)    |
| E-1  | antisense                           | 100% (16/16)   |
| G-1  | antisense                           | 100% (16/16)   |
| H-1  | sense                               | 100% (16/16)   |
| M-1  | sense                               | 100% (16/16)   |
| N-1  | sense                               | 100% (16/16)   |
| Cisco (control) |                              | 100% (8/8)    |

*Each lettuce line represents the progeny of selected plants of *T1* generation. ‘Cisco’ is the parental non-transformed cultivar.

*Samples were collected from lettuce leaves 31 days after inoculation. LBVaV positive total number of plants tested is indicated in parentheses.*
Fig. 1. Southern blot analysis of the transgenic lettuce line A-2 to determine the transgene copy number. (A) A map of the transferred DNA (T-DNA) region of pYK10. Solid lines, DNA probes; RB = T-DNA right border; LB = T-DNA left border; npt II = npt II gene; LBVaV CP = the coat protein (CP) gene of lettuce big-vein associated virus (LBVaV); E = Eco RI site; V = Eco RV site; II = Hind III site. Genomic DNA of line A-2 was digested with Eco RI (lane 1), Eco RV (lane 2) or Hind III (lane 3), and hybridized with the LBVaV CP probe (B) or the npt II probe (C). MW = molecular weight. The figure shows the result from one plant, but the same result was obtained from the other seven plants.

Fig. 2. Time-course of lettuce big-vein associated virus (LBVaV) detection (A), mirafiori lettuce virus (MiLV) detection (B), and symptom expressions (C) in the transgenic lettuce line A-2 plants and the non-transformed ‘Cisco’ lettuce plants. Six line A-2 plants and six ‘Cisco’ plants were inoculated with LBVaV and MiLV using Olpidium brassicae, and the numbers of LBVaV-positive, MiLV-positive or symptomatic plants were counted. LBVaV and MiLV were detected using Western blot analysis, and double-antibody sandwich enzyme-linked immunosorbent assay (DAS-ELISA), respectively. Plants were regarded as symptomatic when vein bandings were observed at least on one leaf.

Fig. 3. The non-transformed ‘Cisco’ lettuce plants (left in each picture) and the transgenic lettuce line A-2 plants (right in each picture) 40 d after inoculation with lettuce big-vein associated virus (LBVaV) and mirafiori lettuce virus (MiLV) using Olpidium brassicae.

ing that the resistance to LBVaV in line A-2 is RNA-mediated. Detection of mRNA from LBVaV CP gene has not been successful while mRNA from npt II gene was detected by Northern blot analysis (data not shown). It is possible that the failure of the detection is due to RNA silencing of the LBVaV CP gene in line A-2. On the other hand, the resistance of line A-2 to MiLV was unexpected because RNA-mediated resistance is only effective against viruses with a high degree of sequence homology to transgenes (van den Boogaart et al., 1998). LBVaV and MiLV are different species (the genera Varicosavirus and Ophiovirus, respectively) (Fauquet et al., 2005), and we can see no significant sequence similarity between them. MiLV infection was possibly inhibited in line A-2 as a result of gene silencing for LBVaV if MiLV infection was helped by LBVaV. In order to make it clear whether the inhibition of MiLV infection in line A-2 was due to the inhibition of LBVaV infection, line A-2 plants were inoculated with O. brassicae carrying MiLV alone (Fig. 4). Fifty-three and 80 d after inoculation, all ‘Cisco’ plants were LBVaV-negative, MiLV-positive and symptomatic, which is consistent with the previous report that MiLV but not LBVaV induced big-vein symptoms in lettuce (Lot et al., 2002). On the other hand, all line A-2 plants were MiLV-negative and without symptoms, indicating that line A-2 is resistant to MiLV regardless of the presence or absence of LBVaV. Further experiments are required to clarify the resistance mechanism to MiLV in line A-2.

The resistant cultivar Pacific was released about 20 years ago (Ryder and Robinson, 1991), but no cultivar with higher resistance has been released because new resistant breeding sources
have not been found in lettuce. The production of transgenic plants with virus-derived nucleotides is therefore an attractive alternative method. In this study we obtained transgenic lettuce with resistance to big-vein using the LBVaV CP gene. It is possible to use this transgenic lettuce as a resistant cultivar or as a breeding source.

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