Characteristics of a Lettuce mosaic virus Isolate Infecting Lettuce in Korea

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Lettuce mosaic virus (LMV) belonging to the genus Potyvirus in the family Potyviridae is one of the most important pathogens of lettuce (Revers et al., 1997b). In most cases, LMV is seed-borne and transmitted by aphid in a non-persistent transmission manner (Bos et al., 1994; Nebreda et al., 2004). LMV has a single positive-stranded genomic RNA of 10,080 nucleotides in length and a poly A tail at its 3’ end. The viral genomic RNA contains a single open reading frame (ORF) which encodes a single large polyprotein of 3,255 amino acids (Revers et al., 1997a). The genomic RNA of LMV is covalently linked to VPg, viral protein genome linked, at its 5’ end and is encapsidated in capsid protein to form flexuous virions.

Worldwide, LMV is important disease of plants in the family Asteraceae, especially lettuce (German-Retana et al., 2008). Lettuce infected with LMV shows several symptoms such as mosaic, dwarfism, yellowing, wilting and failure to form heads. In order to protect lettuce crops, both genetic resistance and virus-free seed are generally used in the field. The recessive allelic resistance genes \(\text{mo}1\) and \(\text{mo}2\) correspond to alleles encoding the eukaryotic translation initiation factor elF4E (eIF4E) in lettuce crops (Tavert-Roudet et al., 2012). However, some LMV isolates have been overcoming the resistance induced by \(\text{mo}1\) alleles in the last 20 years.

LMV isolates have previously been clustered in three main groups, LMV-Yar, LMV-Greek and LMV-RoW (Rest of the World) (German-Retana et al., 2008; Krause-Sakate et al., 2002). LMV-Yar has only one LMV isolate from Yemen and LMV-Greek has 10 LMV isolates from Greece. These two groups are non-seed-borne and non-resistance-breaking. Except for these 11 LMV isolates, all of the remaining isolates distributed around the globe are included in LMV-RoW containing two large subgroups,
LMV-Common and LMV-Most (Krause-Sakate et al., 2004). Both subgroups are seed-borne, but only LMV-Most has the ability to overcome resistance triggered by recessive allelic resistance genes $mo_1^1$ and $mo_1^2$ (Bos et al., 1994; Pink et al., 1992; Revers et al., 1997a, b; Tavert-Roudet et al., 2012). Accordingly, LMV-Most was named after $mo_1$-breaking and seed transmitted. Classification of LMV isolates is based on nucleotide sequence divergence of the nuclear inclusion protein b (NIb)-coat protein (CP) junction between positions 8,936 and 9,151 (Krause-Sakate et al., 2004).

So far, no Korean LMV isolate has been classified and characterized. In this study, we report the characteristics of LMV-Muju, Korean LMV isolate, infecting lettuce. In addition to LMV-Muju, we classified all the rest of the LMV isolates not included in previous report showing dendrogram of LMV isolates.

In order to analyze characteristics of the Korean LMV isolate, we collected lettuce showing pale green and mottle symptoms in emerging crops in Muju county, Jeollabuk-do, Korea in August 2011 (Fig. 1a).

The reverse transcription (RT) polymerase chain reaction (PCR) assay was conducted for the detection of LMV. Total RNA was extracted from LMV-infected leaves with TRI Reagent (MRC, USA) according to the manufacturer’s instructions. The forward primer 08894p (5’- CCG TAC ATA GCI GAR TGT GCT -3’) and the reverse primer 09171m (5’- GCG TTG ATG TCG TCA TCY TT -3’) giving product size of 278 bp were used as diagnostic PCR primers (Krause-Sakate et al., 2002; Peypelut et al., 2004). RT reaction using random N25 primer was performed with one cycle at 42°C for 1 h. PCR was performed with 32 cycles using the procedure (95°C, 30s; 50°C, 30s and 72°C, 1 min), followed by additional extension cycle at 72°C for 5 min. Amplified RT-PCR product was cloned into RBC TA Cloning Vector (RBC Bioscience, Taiwan) and sequence was determined by GenoTech Corp. sequencing service. We detected LMV from lettuce showing pale green and mottle symptoms by using RT-PCR. The sequence result showed sequence homology of up to 98% with characterized LMV isolates, so we therefore named

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**Table 1.** The primer sets used for complete genome of LMV-Muju in RT-PCR

| Primer name | Locus       | Sequence                      | Expected size |
|-------------|-------------|-------------------------------|---------------|
| LMV 1-F     | 1-25        | AAAATAAAAACACACCAAACAAACTC    | 1,700 bp      |
| LMV 1-R     | 1700-1681   | GTACATGCACGAGGCCTGGA          |               |
| LMV 2-F     | 1481-1500   | ATACTCCCATACCTGACATC          | 1,215 bp      |
| LMV 2-R     | 2695-2676   | AGTCTGGATTGAACCATATG          |               |
| LMV 3-F     | 2521-2540   | CGAGATAGCAGTGAACACGC          | 1,260 bp      |
| LMV 3-R     | 3780-3761   | GTGCGTATGGCCTACGTAG           |               |
| LMV 4-F     | 3571-3590   | GGTGCGAGTGCAAGTGCAAG          | 1,440 bp      |
| LMV 4-R     | 5010-4991   | CCAATTTCTATGTGTGGT            |               |
| LMV 5-F     | 4941-4960   | TGGAAATGAGAAGAAGTGGGC         | 1,250 bp      |
| LMV 5-R     | 6190-6171   | CTGTCTTTGCGGCTTACCTT          |               |
| LMV 6-F     | 6111-6130   | CATGGAATGTGTTCAGACG           | 1,550 bp      |
| LMV 6-R     | 7660-7641   | GCTCTCTGAGGTGCTCCCA           |               |
| LMV 7-F     | 7561-7580   | CACAGTCAAGGAGTGAAGTGA         | 1,180 bp      |
| LMV 7-R     | 8740-8721   | ATCCCCACTCTAGGTGGACA          |               |
| LMV 8-F     | 8651-8670   | TGGTTCATGTTCACATTGC          | 1,430 bp      |
| LMV 8-R     | 10080-10061 | GTCTCCGACTGAAAACCAGA          |               |

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*Fig. 1.* Disease symptoms induced by LMV. (A) Systemic mild mosaic symptoms of lettuce naturally infected with LMV. (B) *N. benthamiana* test plant was inoculated with sap of LMV-infected lettuce and showed systemic mild mosaic symptoms.
the isolate LMV-Muju after its origin. In order to prove pathogenicity of LMV-Muju, we inoculated test plants with extracts of infected leaf tissue based on Koch’s postulates. Seedlings of *N. benthamiana* were mechanically inoculated with crude sap of leaf sample infected with LMV-Muju in 1X phosphate buffered saline, pH 7.4, and grown in insect-free incubation room maintained at 23°C with 16 h light period. Disease symptoms of the test plants inoculated with LMV were observed at 10 days post-inoculation (dpi). At 21 dpi, each plant showed systemic mosaic symptoms (Fig. 1b). We also detected LMV from test plants by RT-PCR.

**Fig. 2.** Schematic representation of the complete LMV genome. LMV isolates are classified based on nucleotide sequence divergence of the Nlb-CP junction between positions 8,936 and 9,151. HC-Pro protein of LMV is regarded as a pathogenicity regulator. Amino acid residue 621 of the CI of LMV is involved in overcoming resistance triggered by *mol* alleles.

**Fig. 3.** Phylogenetic tree showing nucleotide sequence divergence of the Nlb-CP junction of LMV isolates (nt 8,936–9,151). The length of the branches and percentage of bootstrap frequency are indicated. LMV-Muju was categorized to the LMV-Common subgroup within the LMV-RoW group, and all the rest of the LMV isolates not included in previous reports were also sorted into LMV-RoW.
In order to determine the complete LMV genome sequence, 8 primer sets were designed from well conserved regions of four complete LMV sequences (GenBank accession no. X97704, X97705, AJ278854, and AJ306288) obtained from NCBI GenBank database (Table 1). Total RNA was extracted from LMV-Muju with the TRI Reagent (MRC, USA) and cDNA was synthesized using random N25 primer with RevertAid Reverse Transcriptase (Thermo scientific, USA) according to the manufacturers’ instructions. PCR was performed with 32 cycles using the procedure (95°C, 30s; 53°C, 30s and 72°C, 2 min), followed by additional extension cycle at 72°C for 5 min. Amplified RT-PCR products were cloned and sequenced as above. Assembly of all the sequences was conducted using the DNAMAN version 5.0.10 program. The complete genome of LMV-Muju (GenBank accession no. KF955619) has 10,080 nt in length, including non-translated sequences at both termini, as for previously characterized isolates (Fig. 2).

Phylogenetic analysis was conducted using CLC Main Workbench 6.1.1 program. LMV-Muju was classified based on nucleotide sequence divergence of the Nib-CP junction between positions 8936 and 9151, and it was sorted into LMV-Common belonging to LMV-RoW (Fig. 2, 3). This result suggested that LMV-Muju may be seed-borne and a non-resistance-breaking isolate. Evidence from previous studies indicates that LM isolates categorized in the same group share similar characteristics (German-Retana et al., 2008; Krause-Sakate et al., 2002). In addition to LMV-Muju, all the rest of the LM isolates not included in previous studies were also classified (Fig. 3). As in previous studies, all of the remainder fall within the LMV-RoW group except for the 11 LMV isolates in LMV-Yar and LMV-Greek.

The large single polyprotein of LMV is processed by three viral proteinases, P1, helper component proteinase (HC-Pro) and nuclear inclusion protein a (Nla)-Pro, into ten mature proteins playing roles in replication and survival of the virus in the host cell (Fig. 2). Several studies reported that HC-Pro proteins of the genus Potyvirus are symptom determinant(s) for pathogenicity (Atreya and Pirone, 1993; Atreya et al., 1992; Dolja et al., 1992; Klein et al., 1994). Ballut et al. (2005) reported that HC-Pro protein of LMV is a pathogenicity regulator which interferes with some of the catalytic functions of the 20S proteasome. LMV-O induces relatively mild mosaic, but LMV-E induces severe mosaic with malformation, stunting and/or necrosis (German-Retana et al., 2008; Pink et al., 1992; Redondo et al., 2001). Redondo et al. (2001) reported that HC-Pro protein of LMV-E is crucial for development of the severe symptoms in susceptible lettuce Trocadéro. Although the symptoms induced by LMV-0 and LMV-E are different, amino acid residues of HC-Pro proteins of the two isolates are quite similar (German-Retana et al., Revers et al., 1997a). They differ at only seven amino acid residues between positions 35 and 287 of HC-Pro proteins (Fig. 2). LMV-Muju was identical to LMV-O at these seven amino acid positions and in inducing relatively mild mosaic symptoms. There were only two different positions at amino acids 72 and 83 of LMV-Muju HC-Pro; however, LMV-O is identical to LMV-E at amino acids 72 and 83. These results suggested that LMV-Muju is similar to LMV-O in having the symptom determinants causing relatively mild mosaic as shown in Fig. 1a, and that the variant residues at positions 72 and 83 do not influence symptom development.

There are some LMV isolates having an ability to overcome resistance triggered by recessive allelic resistance mol gene (mol’ and mol’1), encoding the elF4E (Krause-Sakate et al., 2002; Revers et al., 1997a, b; Tavert-Roudet et al., 2012). Previous studies reported that the C terminus of the cylindrical inclusion (CI) protein and the VPg interacting with lettuce elF4E are involved in mol resistance-breaking (Abdul-Razzak et al., 2009; Tavert-Roudet et al., 2012). Mutations in VPg allowed LMV-O to overcome mol’ resistance, but mutations in C-terminal region of CI protein allowed it to overcome both alleles (Abdul-Razzak et al., 2009). In other words, a mutation at amino acid residue 621 of the CI was sufficient to cause virulence on resistance lettuce (Fig. 2). LMV-E has threonine at amino acid 621 of the CI protein, but LMV-O and LMV-Muju have serine. This result is consistent with the classification shown schematically in Fig. 3. Taken all together, LMV-Muju belonging to LMV-Common is expected to be a seed-borne and non-resistance-breaking isolate.

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