Rapid Detection of *Lily mottle virus* and *Arabis mosaic virus* Infecting Lily (*Lilium spp.*) Using Reverse Transcription Loop-Mediated Isothermal Amplification

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The *Lily mottle virus* (LMoV) impedes the growth and quality of lily crops in Lanzhou, China. Recently *Arabis mosaic virus* (ArMV) has been detected in LMoV-infected plants in this region, causing plant stunting as well as severe foliar symptoms, and likely posing a threat to lily production. Consequently, there is a need to develop simple, sensitive, and reliable detection methods for these two viruses to prevent them from spreading. Reverse transcription (RT) loop-mediated isothermal amplification (LAMP) assays have been developed to detect LMoV and ArMV using two primer pairs that match six conserved sequences of LMoV and ArMV coat proteins, respectively. RT-LAMP assay results were visually assessed in reaction tubes using green fluorescence and gel electrophoresis. Our assays successfully detected both LMoV and ArMV in lily plants without the occurrence of viral cross-reactivity from other lily viruses. Optimal conditions for LAMP reactions were 65°C and 60°C for 60 min for LMoV and ArMV, respectively. Detection sensitivity for both RT-LAMP assays was a hundredfold greater than that of our comparative RT-polymerase chain reaction assays. We have also found this relatively rapid, target specific and sensitive method can also be used for samples collected in the field and may be especially useful in regions with limited or no laboratory facilities.

**Keywords**: *Arabis mosaic virus*, lily, *Lily mottle virus*, RT-LAMP amplification, RT-PCR

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In Lanzhou, China, lilies (*Lilium spp.*) are an important economic crop of the floricultural industry. Also of economic importance are Lanzhou lily (*Lilium davidii* var. *unicolor*) bulbs, grown for their edible and medicinal properties (Zhang et al., 2018). Given that most lilies are grown by vegetative propagation, bulbs may be infected with viruses and consequently transmit from one generation to the next. Moreover, there are more than 10 different viruses that cause lily crops of quality and quantity damage throughout the world (Wang et al., 2010). The *Lily mottle virus* (LMoV; genus Potyvirus, family Potyviridae), being widespread throughout China, is a non-enveloped, rod-shaped flexuous virus with a width of 11 to 15 nm and a length of 680 to 900 nm (Zhang et al., 2017). LMoV infection is characterized by deformity, curling, mottling and necrotic of leaves along with necrotic spotting and deformation of flowers (Zhang et al., 2015b). LMoV infection can be transmitted through vegetative propagation, mechanically from plant to plant or by aphids (Asjes, 2000; Derks et al., 1994; Zhang et al., 2018).

In 2017, *Arabis mosaic virus* (ArMV; genus Nepovirus, family Comoviridae) was detected in LMoV-infected plants in a lily field within the Gaolan Research Station (Gaolan County, Lanzhou, Gansu Province, China;
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3613°N, 10347°E; Field Station, Northwest Institute of Eco-Environment and Resources, Chinese Academy of Sciences, Lanzhou, Gansu Province, China) by a lily research group from the Northwest Institute of Eco-Environment and Resources (NIEER), Chinese Academy of Sciences (CAS) (unpublished report). ArMV is isometric in shape and approximately 30 nm in diameter. In China, ArMV is a second-class quarantinable plant virus. Lilies infected only with ArMV show symptoms of leaf mosaic, chlorosis, stunting, crinkling, and necrosis (Zheng et al., 2013). Similarly, ArMV and LMoV infected plants may also show a sign of plant stunting as well as severe foliar symptoms, including leaf mosaic with necrotic leaf spotting (Fig. 1). ArMV is transmitted by nematodes such as Xiphinema diversicaudatum and Longidorus macrosoma (Brunt et al., 1996). Like LMoV, ArMV can be transmitted mechanically from plant to plant (EFSA Panel on Plant Health, 2013); or through lily bulb vegetative propagation (Zhang et al., 2015b).

Virus management is dependent upon an accurate detection procedure which is convenient, reproducible and scalable for a wide range of samples (Viswanathan et al., 2013). There are a number of approaches used in identifying LMoV and ArMV in lilies, including enzyme-linked immunosorbent assay (ELISA), reverse transcription-polymerase chain reaction (RT-PCR) and immunocapture (IC)-RT-PCR assays. While ELISA has been used to detect both LMoV and ArMV, false negative results and low sensitivity are a problem when dealing with lily bulb samples and in the detection of ArMV in hops (Adams et al., 1987; Sharma et al., 2005). RT-PCR has also been widely used in the detection of LMoV and ArMV; however, polysaccharide, polyphenol as well as other inhibitors in lily bulbs present in RNA extracts have been found to reduce RT-PCR sensitivity (Kulshrestha et al., 2005; Kwon et al., 2013; Zhang et al., 2015b). On the one hand, the IC-RT-PCR technique is not required for RNA isolation, reduces problems associated with PCR inhibitors and provides a more rapid and a less costly approach to preparing templates for amplification (Gambley et al., 2009). On the other hand, IC-RT-PCR sensitivity is likely to be inadequate in detecting low virus concentrations in lilies (Zhang et al., 2017; Zheng et al., 2013).

The RT-loop-mediated isothermal amplification (RT-LAMP) method is a rapid, specific, and sensitive diagnostic technique of relatively low cost that has been applied in the detection of a variety of plant viruses, including the Barley yellow dwarf virus (BYDV) (Zhao et al., 2010), the Piper yellow mottle virus (PYMoV), the Cucumber mosaic virus (CMV) (Bhat et al., 2013), the Turnip mosaic virus (TuMV) (Zhao et al., 2014), the Sugarcane mosaic virus (SCMV), the Sorghum mosaic virus (SrMV) (Keizerweerd et al., 2015) and the LSV (He et al., 2016). In this method, LAMP amplification reactions are typically conducted at a constant temperature between 60-70°C for approximately 1 h in a water bath or heating block. Moreover, amplicons can be readily visualized by adding SYBR Green I to the reaction mixture (Mengao et al., 2016; Parida et al., 2008).

In this study, we have developed and optimized a simple and reliable RT-LAMP method for the rapid and accurate detection of LMoV and ArMV in lily plants. Results from our RT-LAMP assays were then compared to our established RT-PCR assays.

Materials and Methods

Plant materials. Naturally infected oriental hybrid lily (Lilium oriental cv. Sorbonne) plants that exhibited typical symptoms of plant dwarfing, chlorotic (yellowing) leaf spotting or striping were collected from fields within the Gaolan Research Station. RT-PCR was used to test leaves removed from near the flower bud as described by Zhang...
Leaves from a single plant specimen that tested positive for LMoV, ArMV, LSV, or CMV, serving as a source of these viruses, were selected and stored at –70°C for use in our experiments. RT-LAMP primer design. The conserved regions of coat proteins (CP) were used as the basis for the design of RT-LAMP primers for LMoV and ArMV detection by establishing sequence alignments (multiple sequence alignment) of LMoV and ArMV CP gene sequences from lily hosts available from GenBank (Zhang et al., 2015b; Zheng et al., 2013). Primer design was conducted using PrimerExplorer V5 software, from Eiken Chemical Co., Ltd. (Tokyo, Japan) using the default settings. Two LAMP external primers (F3 and B3) and two internal primers (FIP [F1c+F2] and BIP [B1c+F2]) were designed, which are capable of detecting six LMoV and ArMV CP gene sequences, respectively (sequences shown in Table 1).

RNA extraction. Total RNA was extracted from lily leaves using the RNAprep Pure Kit (For Plant) from Tiangen Biotech (Beijing) Co., Ltd. (Beijing, China). A lily leaf sample (0.05-0.1 g) was ground in liquid nitrogen, after which 450 ml of lysate was added. Following this, further extraction steps were conducted as per the manufacturer’s instructions. The final elution step was conducted using 50 μl of RNase-free H₂O. The RNA concentration was measured using the NanoDrop ND-1000 Spectrophotometer (Nanodrop Technologies, Wilmington, DE, USA) and adjusted for RT-LAMP and RT-PCR analyses.

RT-LAMP primer design. The conserved regions of coat proteins (CP) were used as the basis for the design of RT-LAMP primers for LMoV and ArMV detection by establishing sequence alignments (multiple sequence alignment) of LMoV and ArMV CP gene sequences from lily hosts available from GenBank (Zhang et al., 2015b; Zheng et al., 2013). Primer design was conducted using PrimerExplorer V5 software, from Eiken Chemical Co., Ltd. (Tokyo, Japan) using the default settings. Two LAMP external primers (F3 and B3) and two internal primers (FIP [F1c+F2] and BIP [B1c+F2]) were designed, which are capable of detecting six LMoV and ArMV CP gene sequences, respectively (sequences shown in Table 1).

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Table 1. Sequences of primers used for RT-LAMP and RT-PCR of Lily mottle virus (LMoV) and Arabis mosaic virus (ArMV)

| Primer names   | Type                          | Position* | Sequence (5′-3′)            |
|---------------|-------------------------------|-----------|-----------------------------|
| LMoV-F3       | Forward outer primer for RT-LAMP | 9,024-9,041 | AATGGCGTGTGCGTCTAG          |
| LMoV-B3       | Backward outer primer for RT-LAMP | 9,193-9,212 | GGTGAGATTTCGCTGAAGGC        |
| LMoV-FIP      | Forward inner primer (F1c-F2) for RT-LAMP | F1c: 9,083-9,104, F2: 9,043-9,062 | CGTCGGTTTTGCGTGTTCAAGTTGGACGGA-GATCAGCAGTTT |
| LMoV-BIP      | Backward inner primer (B1c-B2) for RT-LAMP | B1c: 9,120-9,141, B2: 9,173-9,192 | GCGCAATTTCACACCACTCGGTACCTAG- GCATGTCAGG |
| LMoV-F        | For RT-PCR                     | 9,005-9,023 | TGAGCACCTCCAATGTA           |
| LMoV-B        | For RT-PCR                     | 9,399-9,419 | CACATCTCTGCTGATGCTCTCT      |
| ArMV-F3       | Forward outer primer for RT-LAMP | 2,950-2,969 | CGAGTGAGAAAACGTGCTTAAG      |
| ArMV-B3       | Backward outer primer for RT-LAMP | 3,162-3,181 | CCATGAGACGCTATCAGTACG       |
| ArMV-FIP      | Forward inner primer (F1c-F2) for RT-LAMP | F1c: 3,041-3,064, F2: 2,984-3,005 | CGAGAAAGTCCACCACCCAAAGAAAAAGCAATT-GACAGAAATGCAATTA |
| ArMV-BIP      | Backward inner primer (B1c-B2) for RT-LAMP | B1c: 3,078-3,102, B2: 3,138-3,155 | ACAGAAGAGATAGGAATAGGTCGAAAGAT- TATCTCCAGGTC |
| ArMV-F        | For RT-PCR                     | 2,241-2,262 | ACTACTGCTAGTCTGCGTATA       |
| ArMV-B        | For RT-PCR                     | 2,673-2,690 | ATAGAACCCTAGACCTAGCC         |

RT-LAMP, reverse transcription loop-mediated isothermal amplification; RT-PCR, reverse transcription polymerase chain reaction.

*Gene position based on the nucleotide sequence of the LMoV and ArMV coat proteins (CP) (accession nos. AJ564636 and AB279740.2) respectively.
RT-LAMP amplification products (5 μl) were analyzed and evaluated by direct visual inspection of the reaction tube after the addition of SYBR Green I (1:1000 TE, v/v, Invitrogen, Carlsbad, CA, USA). In addition, RT-LAMP amplification products (5 μl) were routinely detected with agarose gel electrophoresis (2.0% agarose, TAE) and by direct visual examination using SYBR Green I. As a comparison, each leaf sample was also tested using RT-PCR according to the aforementioned method.

**Results**

**Optimization of RT-LAMP reactions.** The optimum temperatures for the detection of LMoV and ArMV using RT-LAMP were confirmed by means of 100 min RT-LAMP reactions at seven different temperatures, 56°C and 70°C. For LMoV detection typical ladder-like patterns were observed using agarose gel electrophoresis at 65, 68, and 70°C; however, no amplification was seen at 56, 58, 60, and 62°C (Fig. 2A). For ArMV detection typical ladder-like patterns were clearly observed using agarose gel electrophoresis at 60, 62, 65, 68, and 70°C; however, ladder-like patterns were only weakly observed using agarose gel electrophoresis at 58°C. No amplification was observed at 56°C (Fig. 2B).

To determine the minimum reaction time for LMoV and ArMV detection, RT-LAMP was conducted at 65°C and 60°C for 20, 40, 60, 80, and 100 min for LMoV and ArMV, respectively. Typical ladder-like patterns were clearly observed at 60, 80, and 100 min; however, no amplification was observed at 20 and 40 min in either case (Fig. 2).

**Analysis of RT-LAMP specificity.** Total RNA extracts from lily leaves infected with LSV, CMV, LMoV, and ArMV were used to evaluate the cross-reactivity of the RT-LAMP assay. Water and healthy lily leaves were used as the blank and negative controls, respectively.

**Comparison between RT-LAMP and RT-PCR sensitivity.** To compare the sensitivity of RT-LAMP to RT-PCR, total RNA (100 ng/μl) from lily leaves infected with LMoV or ArMV were used for both RT reactions. The synthesized cDNA products were serially diluted 10-fold from $10^{-1}$ to $10^{-11}$ using initial cDNA. The serial dilutions were subjected to LAMP and PCR according to the aforementioned method.

**Evaluation of RT-LAMP using field-collected samples.** To validate the RT-LAMP assay, 10 Lanzhou lily and 10 oriental hybrid lily leaf samples were collected from the field within the Gaolan Research Station and were extracted as discussed above and tested for LMoV and ArMV using RT-LAMP. Amplification products (5 μl) were detected using agarose gel electrophoresis (2.0% agarose, TAE) and by direct visual examination using SYBR Green I. As a comparison, each leaf sample was also tested using RT-PCR according to the aforementioned method.

![Fig. 2](image-url) Agarose gel electrophoresis showing the effect of different amplification temperatures on the detection of *Lily mottle virus* (LMoV) (A) and *Arabis mosaic virus* (ArMV) (B) by reverse transcription loop-mediated isothermal amplification. Lane M, DL600 marker; lane 1, negative control; lanes 2-8, 56, 58, 60, 62, 65, 68, and 70°C.
Consequently, subsequent RT-LAMP analyses were conducted at 65°C and 60°C for 60 min for LMoV and ArMV, in our experiment.

Specificity of the RT-LAMP assay for LMoV and ArMV. To determine the specificity of RT-LAMP assays, we tested them using RNA extracts from lily leaves infected with LSV, CMV, LMoV, and ArMV. Following the application of RT-LAMP, only products of LMoV-infected samples exhibited a typical ladder-like pattern using agarose gel electrophoresis. No bands were observed in samples infected with LSV, CMV, or ArMV (Fig. 4A, upper). Similarly, only the products of ArMV-infected samples exhibited a typical ladder-like pattern with no bands observed in samples infected with LSV, CMV or LMoV (Fig. 4B, upper).

The reaction products of RT-LAMP were further confirmed by visualization through direct observation using green fluorescence after adding SYBR Green I. Only the products of LMoV and ArMV-infected samples exhibited a green fluorescence and were consequently visualized after adding SYBR Green I (Fig. 4A, lower and 4B, lower).

Sensitivity of RT-LAMP vs. RT-PCR. To compare sensitivities between RT-LAMP and RT-PCR, we used ten-fold serial dilutions of initial cDNA products synthesized from RT as templates. The detection limits for LMoV were determined to be $10^{-10}$ using RT-LAMP (Fig. 5A, upper) and $10^{-8}$ using RT-PCR (Fig. 6A). The detection limits for ArMV were determined to be $10^{-8}$ using RT-LAMP (Fig. 5B, upper) and $10^{-6}$ using RT-PCR (Fig. 6B). Thus, the sensitivity of IC-RT-LAMP appeared to be greater by a factor of 100 compared with the sensitivity of RT-PCR for the detection of LMoV and ArMV. The reaction products of RT-LAMP were also directly observed using SYBR Green I fluorescence. Both methods yielded similar detection limit values for both viruses (Fig. 5A, lower and 5B, lower).
Evaluation of RT-LAMP using field-collected samples.
We analyzed 20 field samples using both RT-LAMP and RT-PCR. For the lily samples, 7 were determined to be positive for LMoV and 13 were negative (Fig. 7A, upper and middle). Using the same procedure (i.e., applying both RT-LAMP and RT-PCR), of the 20 samples tested only one oriental hybrid lily sample was determined to be positive for ArMV while the remaining 19 samples were found to be negative (Fig. 7B, upper and middle). We further determined that the sequence of the ArMV CP gene from this Lanzhou isolate had a 90.2% similarity with a known ArMV lily isolate recorded in GenBank (accession no. AB279741) (data not shown).

Products from both methods analyzed using the SYBR Green I fluorescence technique with agarose gel electrophoresis provided similar results with respect to the 20 LMoV and ArMV-infected field samples to those discussed above (Fig. 7A, lower and 7B, lower).

Discussion
LMoV is one of most common viruses to infect lilies worldwide. In the Lanzhou region, such virus infections have contributed to a deterioration in bulb quality and a yield reduction greater than 50%, namely, from 23,000 to 11,000 kg per hectare (Zhang et al., 2018). Recently ArMV was detected in LMoV-infected plants in an oriental lily field in this region, there is concern that once ArMV establishes within a lily production area, and it may result in further economic losses to the lily industry.

Although different methods have been developed for LMoV and ArMV detection, many require technical expertise and specialized equipment (Kulshrestha et al., 2005; Niimi et al., 2003; Zhang et al., 2015b). In this study, we report on the successful development of a simple, sensitive and reliable method to detect LMoV and ArMV using the RT-LAMP method at temperatures validated between 65°C and 70°C and 60°C and 70°C for LMoV and ArMV,
respectively, for approximately 60 min. We found that both RT-LAMP and RT-PCR methods yielded similar results for field-collected samples. Moreover, using RT-LAMP, both detection limits were 100 times more sensitive than the RT-PCR method. These results further demonstrated that our RT-LAMP assay is a reliable method that is significantly more sensitive than RT-PCR.

Compared to the conventional RT-PCR method, RT-LAMP is not only less expensive in relation to the equipment required but also more sensitive, and it has been used not only to detect plant viruses but also various other pathogens (Arita et al., 2009; Bühlmann et al., 2013; Kubota et al., 2008). In the development of an efficient LAMP amplification, the design, ratios and concentrations of LAMP primers are key factors in determining LAMP sensitivity (Hardinge et al., 2018). In this study, we found that different primer sets yielded different amplification in LMoV and ArMV detections (data not shown), and the optimum primer sets selected in this study could be a main reason for determining the high sensitivity performance of RT-LAMP assays. The optimization of the LAMP reaction temperature and duration (time) also played an important role in the successful detection of pathogens (Bhat et al., 2013). In determining the optimum temperature, the LAMP reaction was initially conducted using a thermal-cycler to maintain a constant temperature. Subsequently, two further options were assessed to ascertain their suitability for this purpose: a water bath and a heat block. We found that all three RT-LAMP reaction incubation methods yielded similar amplification results (data not shown), which served to confirm that a device that can maintain a set temperature accurately is adequate for RT-LAMP amplification (Bhat et al., 2013).
RT-LAMP products were routinely confirmed using agarose gel electrophoresis. As reported by several other studies (Bhat et al., 2013; Mengao et al., 2016; Ravindran et al., 2012), our experimental results also confirmed that RT-LAMP products could be detected through a simple visual examination under green fluorescence using SYBR Green I. Both visual methods (gel electrophoresis and green fluorescence) yielded similar results with respect to specificity, detection limits and with field-collected samples (Figs. 4, 5, and 7). Our results also showed that visual indicators such as green fluorescence appear to be adequate for performing RT-LAMP in laboratories with minimal facilities. However, further research is required to develop a suitable rapid field-based RT-LAMP for lily crops using green fluorescence.

A previous study recently confirmed a low level of LMoV infection in the Lanzhou lily (Zhang et al., 2018). From our limited collection of field samples in this work, we neither detected LMoV nor ArMV in our Lanzhou lily samples using both RT-LAMP and RT-PCR methods. On the other hand, we did detect LMoV (70.0%) and ArMV (10.0%) in oriental hybrid lily samples using the same methods. While it is encouraging to note that LMoV and ArMV may not be well established in the Lanzhou lily in this region, further survey work needs to be undertaken to verify this conclusion. We plan to undertake further survey work in order to further clarify if either or both viruses are further establishing or spreading in Gansu Province. Based on this information monitoring and management programs of these two viruses should be strengthened in order to avoid the spread of disease and remove the threat of economic losses.

In conclusion, mixed infections of LMoV and ArMV show severe foliar symptoms, and likely bring about damage to the lily crop. Accordingly, we have developed a convenient, specific and sensitive RT-LAMP procedure for the detection of LMoV and ArMV. Through testing, we determined that this relatively simple technique should be more widely available for application owing to its low cost and minimal equipment requirements.

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