Rapid Detection and Differentiation of Three Cucurbit-infecting Poleroviruses by Multiplex RT–PCR

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Abstract
A multiplex reverse transcription polymerase chain reaction (mRT-PCR) method was developed and optimized for the simultaneous detection and differentiation of three poleroviruses infecting cucurbits. Cucurbit
aphid-borne yellows virus (CABYV), Melon aphid-borne yellows virus (MABYV) and Suakwa aphid-borne yellows virus (SABYV) could be differentiated simultaneously using four optimized specific oligonucleotide primers, including one universal primer for detecting poleroviruses and three virus-specific primers. Amplification of three target viruses was also optimized by increasing the PCR annealing temperatures. The mRT-PCR products consisted of fragments of 700 base pairs (bp) for CABYV, 450 bp for MABYV and 950 bp for SABYV. Detection limits of the RNA quantity for mRT-PCR were 1 pg for CABYV, 0.1 pg for MABYV and 1 pg for SABYV. No specific products could be amplified from RNA of other non-target poleroviruses. The mRT-PCR was found to be a specific, sensitive and cost-effective method for detecting multiple poleroviruses in cucurbits.

Keywords: Cucurbit poleroviruses, Cucurbit aphid-borne yellows virus, Melon aphid-borne yellows virus, Suakwa aphid-borne yellows virus, Multiplex RT–PCR

1. Introduction

Cucurbits are cultivated widely in China, but viral diseases can lead to lethal syndromes, regular epidemics and considerable economic losses. Cucurbit aphid-borne yellows virus (CABYV) is the first reported polerovirus infecting cultivated cucurbits naturally and significantly reduces yield in France (Abou-Jawdah et al., 1997; Lecoq et al., 1992). Yield losses of up to 40% have been reported in melon plants, and although fruit shape and quality are not adversely impacted, infected plants characteristically produce fewer fruits per plant (Lecoq et al., 1992). Subsequently, CABYV has been found in Italy, Lebanon, Spain, Iran, Turkey, Tunisia and the USA (Abou-Jawdah et al., 1997; Bananej et al., 2006; D’Arcy and Domer, 2005; Juarez, 2004; Lecoq et al., 1992; Lemaire et al., 1993; Mnari-Hattab, 2005; Tomassoli and Meneghini, 2007; Yurdumcu and Özoğnen, 2007).

In 2006, CABYV was first detected in mainland China from nine cucurbit crops showing yellowing symptoms (Xiang et al., 2008b). Later, two new poleroviruses infecting cucurbits in China, Melon aphid-borne yellows virus (MABYV) and Suakwa aphid-borne yellows virus (SABYV), were found in our study. MABYV is accepted as one of the 13 formally described virus species, and SABYV remains a tentative designation within the Polerovirus genus by the International Committee on Taxonomy of Viruses (Knierim et al., 2010; Shang et al., 2009; Xiang et al., 2008a).

Polaroviruses are plant viruses that contain a small, single molecule of linear, positive-sense ssRNA (D’Arcy and Domer, 2005; Prüfer et al., 1995). The virus particles are approximately 25 to 30 nm in diameter, are isometric and hexagonal in outline, and have no envelope (D’Arcy and Domer, 2005). These plant viruses are phloem restricted and are not uniformly distributed in hosts (Hoffmann et. al., 2001). The routine serological detection method using polyclonal antibodies lacks the sensitivity necessary to detect poleroviruses, given that the virus occurs at very low or variable concentrations. Furthermore, current serological assays cannot differentiate the various polerovirus species that infect cucurbits, due to cross reactions (D’Arcy et al., 1989; Mnari-Hattab et al., 2009; Robertson et al., 1991). However, RT-PCR with specific primers can distinguish among these virus species (Hauser et al., 2000; Knierim et al., 2010; Lemaire et al., 1995; Mnari-Hattab et al., 2009; Shang et al., 2009; Xiang et al., 2008a; Xiang et al., 2008b).

The polerovirus genome contains six open reading frames (ORFs), has a VPg linked to the 5’ end of the genomic RNA and lacks a poly(A) tract or a tRNA-like structure on the 3’ end (Fig. 1) (D’Arcy and Domer, 2005). Poleroviruses possess an ORF0 in the 5’ end and a non-coding region of about 200 nucleotides (nt) between ORF2 and ORF3 (D’Arcy and Domer, 2005). RT-PCR with universal primers allows for the amplification of a 1.4 kb band for poleroviruses (Xiang et al., 2008a; Xiang et al., 2008b). Other primers have been used to separately detect CABYV, MABYV and SABYV, in order to describe the geographical distribution and molecular diversity of these poleroviruses in China. However, these methods only recognize one virus at a time (Shang et al., 2009; Xiang et al., 2008a). The detection of several different viruses from large numbers of cucurbit samples by running multiple simplex PCRs per sample is unnecessarily costly and time-consuming (Knierim et al., 2010; Shang et al., 2009; Xiang et al., 2008a). Multiplex RT-PCR is a popular technique that offers fast, reliable and cost-effective detection of multiple viruses simultaneously (Deb and Anderson, 2008; Viganó and Stevens, 2007; Wei et al., 2009). Here, we report a specific and sensitive multiplex RT-PCR method that can detect and differentiate three poleroviruses, CABYV, MABYV and SABYV, infecting cucurbit crops.

2. Materials and Methods

2.1 Plant material and Recombinant Plasmids

Cucurbit leaf tissues infected with individual known poleroviruses, CABYV, MABYV and SABYV, were used to standardize the multiplex RT-PCR. Each of these three viruses infecting cucurbits was identified by RT-PCR.
and sequenced in our previous research (Shang et al., 2009). Additional cucurbit samples from different fields in China showing leaf-yellowing symptoms were collected for testing.

RT-PCR products amplified with primers for polerovirus detection (PococpR/ PoconF) were purified, using a PCR DNA Purification Kit (Axygen) according to the manufacturer’s instructions, and inserted into the pMD19-T vector (TaKaRa). Plasmids pCaCAI-169, pCaMA5-85 and pTSAB-1, harboring the expected size inserts for each PCR product from viruses CABYV, MABYV and SABYV, respectively, were constructed and used to study multiplex PCR. Plasmids containing partial sequence cDNA clones of *Turnip yellows virus* (TuYV) and *Sugarcane yellow leaf virus* (ScYLV), two other poleroviruses constructed in our previous research, were used to test specificity of the multiplex PCR.

### 2.2 RNA Extraction and RT-PCR Detection

Total RNA from 0.3 g of leaf tissue from infected plants was prepared by SDS-phenol/chloroform extraction and eluted in a final volume of 10 µl of diethylpyrocarbonate-treated (DEPC) water and stored at -20ºC for the following protocols (Han et al., 2000).

Two-step RT-PCR detection for poleroviruses was performed as described earlier (Shang et al., 2009; Xiang et al., 2008a). Amplified products (5 µl each) were electrophoresed in 1% agarose gels and stained with ethidium bromide to confirm the expected size of the fragments.

### 2.3 Primer Selection and Optimization of Annealing Temperatures

The multiplex RT-PCR assay was designed to be carried out using a mixture of the universal polerovirus primer PococpR and primers specific for different viruses.

Three specific sense primers, CA3414F, MA3566F and SA3133F, had been used to detect CABYV, MABYV and SABYV separately in simplex PCR (Shang et al., 2009; Xiang et al., 2008a). However, a multiplex PCR containing each of the four primers described above could not differentiate the three viruses, even under varied PCR conditions. Under most conditions, the cDNA of MABYV could not be amplified in multiplex PCR. Accordingly, we designed other primers for the detection of MABYV based on its RNA sequence (GenBank Accession No. NC010809). We gave consideration to the size of the PCR product and to the interaction between primers, which can affect amplification efficiency. The sequences of the designed primers used in this study are listed in Table 1.

The optimization of annealing temperatures was based on 50ºC in initial protocols. Gradient PCR was performed using different temperatures that were set randomly from 45ºC to 58ºC by the PCR machine.

### 2.4 Cloning and Sequencing

Purified PCR products, amplified with primers PoT7conF and PoE5cocpR from clones pCaCAI-169, pCaMA5-85 and pTSAB-1, were inserted into pMD19-T and then transformed into competent cells of *Escherichia coli* DH5α. The sequences of the primers designed for construction of recombinant clones are listed in Table 2. Recombinant clones pMD-CAT7-9, pMD-MAT7-19 and pMD-SAT7-45, containing about 1400 nt of the cDNA fragments of CABYV, MABYV and SABYV, were constructed and included T7 polymerase and EcoRV restriction enzyme sites.

All clones were selected and identified by using colony PCR, and sequenced with M13-47 forward and M13-48 reverse primers by the dideoxynucleotide chain termination method using an automated sequencer (ABI Prism™ 3730, Applied Biosystems, USA) and the Big-Dye™ Terminator Cycle Sequencing Ready Reaction Kit.

### 2.5 In vitro RNA Transcription and Quantification

Plasmids pMD-CAT7-9, pMD-MAT7-19 and pMD-SAT7-45 were linearized by EcoRV restriction digestion and transcribed in *vitro* by T7 RNA polymerase. Reaction mixtures contained the following in a final volume of 50 µl: 1 µg linear plasmid DNA, 2 µl 20 U/µl T7 RNA polymerase and 10 µl 5X T7 polymerase reaction buffer, 5 µl 100mM DTT, 1 µl RNasin, 2.5 µl each 10 mM ATP/GTP/CTP/UTP. The reaction mixtures were incubated at 37ºC for 90 min, and the RNA polymerization reaction was terminated by the addition of 1 µl RNase-free DNasel, followed by incubation at 37ºC for 30 min according to the manufacturer’s protocol (Promega, USA). Confirmation that the RNA bands represented full-length transcripts was conducted by electrophoresis in a 1% agarose gel, and the product RNA molecules were quantified and tested by Nano analysis.

### 2.6 Sensitivity of Multiplex PCR

The sensitivity was considered as the lowest concentration of viral RNA giving a strong positive signal in mRT-PCR. To determine this threshold, ten-fold serially diluted RNA templates of three viruses were tested.
using a one-step RT-PCR kit (Qiagen, USA). Polerovirus RNA samples, ranging in quantity from 1 fg to 10 ng, were prepared in 5 μl of RNA extracted from healthy cucurbit leaf tissue and the assay was carried out as described above.

3. Results

3.1 Primer Selection and Optimization of Annealing Temperatures

Published universal polerovirus primer PococpR and species-specific primers CA3414F and SA3133F were used in our study (Shang et al., 2009; Xiang et al., 2008a). A newly designed primer MA3639F was included, to allow for mRT-PCR. We developed an mRT-PCR method using the four primers listed above that were capable of differentiating CABYV, MABYV and SABYV in infected plant tissue. Annealing temperatures between 45°C and 58°C were tested in order to optimize amplification by the Gradient PCR machine based on 50°C in our initial protocols (Wei et al., 2009). The optimal annealing temperature for mRT-PCR was determined to be 51°C. The mRT-PCR products were 700 bp for CABYV, 450 bp for MABYV and 950 bp for SABYV. A negative control containing RNA from an uninfected leaf gave no signal (Fig. 2).

3.2 Sensitivity and Specificity of Multiplex PCR

The detection limit of the multiplex PCR was determined by testing ten-fold serial dilutions of the individual transcribed RNA from CABYV, MABYV and SABYV. The quantities of in vitro-transcribed RNA with known sequence were tested and serially diluted with healthy plant RNA. The specific PCR products created using diluted individual transcribed RNA target were detected after agarose gel electrophoresis by ethidium bromide staining (Fig. 3). Detection limits of the RNA quantity for mRT-PCR were 1 pg for CABYV, 0.1 pg for MABYV and 1 pg for SABYV. Plasmids containing cDNA sequences from two other poleroviruses, TuYV and ScYLV, were used to test the specificity of the multiplex PCR, and gave no signal (Fig. 2).

3.3 Detection of the Three Poleroviruses in Field Samples

Cucurbit leaf tissue samples collected from different fields in China were used to test and standardize the mRT-PCR (Fig. 4). Samples of cushaw and squash from Fujian, cucumbers from Beijing, Suakwa vegetable sponge, and squash and cushaw from Jiangxi in China were tested. The positive control containing RNA of CABYV, MABYV and SABYV could produce three distinct fragments, 700 bp, 450 bp and 950 bp, respectively. All of the negative controls, including healthy plant samples, gave no signal as expected. In all single infections and combinations of simulated double infections, the viruses were detected and differentiated by the multiplex PCR in a single reaction using four primers together. The results were confirmed by monospecific RT-PCR. Because of the high sensitivity and specificity of the mRT-PCR method, it is easy to differentiate poleroviruses in cucurbit samples infected by CABYV, MABYV and/or SABYV.

4. Discussion

There have been considerable advances in the field of viral diagnosis, which have been brought about by rapidly advancing PCR methods that enable the diagnosis of specific viral infections. Furthermore, multiplex RT-PCR for the detection of different viruses simultaneously is popular because it provides fast, reliable and cost-effective results (Deb and Anderson, 2008; Viganó and Stevens, 2007; Wei et al., 2009).

The mRT-PCR method developed here was able to detect and differentiate simultaneously CABYV, MABYV and SABYV in one single reaction. Poleroviruses have become very important viruses infecting cucurbit crops in the world. In mainland China, CABYV is prevalent and widely occurring on nine cucurbit crops at least in 25 different provinces, while MABYV was found in Beijing, Inner Mongolia, Anhui, Jiangxi and Hainan provinces. To date, SABYV has been detected only in the southern parts of China, including the Guangxi, Guangdong and Fujian provinces (Knierim et al., 2010; Shang et al., 2009; Xiang et al., 2008b). In the past, the detection of these poleroviruses required separate, simplex PCR. Therefore, an mRT-PCR method which can rapidly identify poleroviruses will be important and helpful in studying the distribution and control of cucurbit poleroviral disease. The mRT-PCR is sensitive, specific, cost effective and useful in detection, and has the added benefits of saving time and using fewer reagents compared with simplex RT-PCR (Deb and Anderson, 2008). The application of this mRT-PCR-based assay to field samples resulted in the direct detection of RNA from CABYV, MABYV and/or SABYV infecting cucurbits.

The universal set of primers for poleroviruses were able to amplify a 1.4 kb DNA band for any of the poleroviruses, and three specific sense primers, CA3414F, MA3566F and SA3133F, were used to detect CABYV, MABYV and SABYV separately in simplex PCR as we reported (Knierim et al., 2010; Shang et al., 2009; Xiang et al., 2008b). It has been reported that amplification efficiency is inversely correlated with the amplicon size (Wei et al., 2009); shorter products could be preferentially amplified compared to larger products, although that
is not always the case. In our study, the shortest product of the cDNA of MABYV could not be amplified in multiplex PCR while other longer products from CABYV and SABYV could be amplified. So, newly designed primers were tested and then one of those primers, MA3639F, was selected to be used in mRT-PCR. Annealing temperature is an important factor for PCR specificity and amplification efficiency (Ma and Michailides, 2007; Wei et al., 2009). Therefore, the annealing temperature was optimized. The results indicate that the primers were correctly designed to avoid possible primer-dimers, and the mRT-PCR was specific and sensitive for detecting different targets in a single reaction. In our study, the newly developed multiplex RT-PCR could detect and simultaneously differentiate CABYV, MABYV and SABYV in one reaction using primers PococpR, CA3414F, MA3639F and SA3133F, which can also be used in monospecific RT-PCR.

The approach developed in this study provides a simple and convenient way to develop multiplex assays based on published primers, supplemented with newly designed primers where necessary. This specific and sensitive method for detecting multiple poleroviruses in cucurbit is could be used for large-scale sampling efforts to study the distribution of poleroviruses in China as well as other areas in the world. This detection technique could facilitate research on cucurbit-infecting polerovirus epidemiology, outbreak monitoring and investigations of interactions among viruses, hosts and vectors.

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References


Ma, Z. & Michailides, T.J. (2007). Approaches for eliminating PCR inhibitors and designing PCR primers for the


Table 1. List of primers used for developing multiplex RT-PCR

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5’ to 3’)</th>
<th>Product (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MA3817F</td>
<td>AGTATAAAATCTCAATGGTCAAGT</td>
<td>283</td>
</tr>
<tr>
<td>MA3340F</td>
<td>AGCCGGAATACGTTGTTGCAATTGC</td>
<td>760</td>
</tr>
<tr>
<td>MA3420F</td>
<td>ATTCACATCGGGCTTCCTGG</td>
<td>680</td>
</tr>
<tr>
<td>MA3603F</td>
<td>GTGGTCCAACACCACCTGGCCA</td>
<td>497</td>
</tr>
<tr>
<td>MA3639F</td>
<td>CGACGCAGCAGAATCCC</td>
<td>461</td>
</tr>
<tr>
<td>CA3488F</td>
<td>GCACACGTGCCTCAATTGTAATG</td>
<td>612</td>
</tr>
</tbody>
</table>
Table 2. Primers and designing rules used for constructing recombinant clones

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5’ to 3’)</th>
<th>Designing rule</th>
</tr>
</thead>
<tbody>
<tr>
<td>PoT7conF</td>
<td>GTAATACGACTCACTATAGTGGTTCGGTTTTGACTGG</td>
<td>19nt added to the 5’-end of primer PoconF, including promoter sequence of phage T7 polymerase (shaded) and additional G for enhancing transcription (underlined).</td>
</tr>
<tr>
<td>PoE5copol</td>
<td>GATATCGGATCCCGTCTACCTTTGCTGACCTAGG</td>
<td>12nt added to the 5’-end of primer Pocopol, including restriction enzyme cutting site of EcoRV and BamHI (shaded).</td>
</tr>
<tr>
<td>sqxT7</td>
<td>GTAATACGACTCACTATAG</td>
<td>Primer with promoter sequence of phage T7 polymerase (shaded) and additional G (underlined) designed for the detection of recombinant clones.</td>
</tr>
</tbody>
</table>

Figure 1. Diagram of the genome organization of poleroviruses

Solid lines represent RNA; boxes represent ORFs; square represents VPg. Arrowheads represent the positions of the universal primers for the detection of poleroviruses.

Figure 2. Agarose gel electrophoresis of multiplex RT-PCR products of different poleroviruses infecting cucurbits

Lane 1, DNA size marker, λ DNA digested by EcoRI and HindIII; Lane 2, CABYV; Lane 3, MABYV; Lane 4, SABYV; Lane 5, CABYV, MABYV and SABYV; Lane 6, TuYV; Lane 7, ScYLV; Lane 8, no template.
Figure 3. Evaluation of the detection limit of multiplex RT-PCR using ten-fold serial dilutions of individually transcribed RNA from CABYV, MABYV and SABYV. A for CABYV; B for MABYV; C for SABYV.

Lane 1, DNA size marker, λ DNA digested by EcoRI and HindIII; Lanes 2-9, 10 ng, 1 ng, 0.1 ng, 10 pg, 1 pg, 0.1 pg, 10 fg and 1 fg of RNA of different viruses; Lane 10, positive plasmids of CABYV, MABYV and SABYV; Lane 11, healthy control of cushaw leaf; Lane 12, no template.

Figure 4. Agarose gel electrophoresis of multiplex RT-PCR products of different poleroviruses infecting cucurbits collected from different locations.

Lane 1, DNA size marker, λ DNA digested by EcoRI and HindIII; Lanes 2-4, cushaw 1, cushaw 2 and squash from Fujian Province, China; Lanes 5-6, cucumber 1, cucumber 2 from Beijing, China; Lanes 7-9, Suakwa vegetable sponge, squash and cushaw from Jiangxi Province, China; Lane 10, positive control of CABYV, MABYV and SABYV; Lane 11, no template; Lane 12, negative control of healthy cushaw leaf.