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A degenerate pair of primers for simultaneous detection of four alpha- and betanecroviruses

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Abstract

The high infection levels due to Olive latent virus 1 (OLV-1), Olive mild mosaic virus (OMMV) (alphaneoviruses) and Tobacco necrosis virus D (TNV-D) (betanecroviruses) in Portuguese olive orchards prompted us to develop a rapid PCR-based assay for the simultaneous detection of these viruses aimed at the sanitary selection and marketing of plant material in compliance with European Union regulations. A pair of degenerate oligonucleotide primers, parRdRp5 and parCoat3 was designed based on conserved regions located in the RNA-dependent RNA polymerase (RdRp) and coat protein (CP) genes of these viruses and one other alphaneovirus, Tobacco necrosis virus A. Its use in RT-PCR assays generated a product of ca. 2000 bp for the 4 viral species tested. These primers were compared with virus specific primers in multiplex RT-PCR, and identical results were obtained. Its application to dsRNA extracted from 54 olive field growing trees originated the expected ca. 2000 bp amplicon in 17 trees. The virus identity was determined by sequencing the cloned RT-PCR products. No TNV-A was found.

The RT-PCR assay using the degenerate primers described in this study were shown to be reliable in detecting any of the above-mentioned alpha- and betanecroviruses, and it is as sensitive as that which uses virus specific primers in multiplex assays. Therefore, this assay is well suited for the rapid screen of virus-free plant material in selection and improvement crop programmes. Additionally, it has the potential to reveal virus diversity and the presence of new viruses, provided the RT-PCR generated amplicon is further sequenced.

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1. Introduction

European Union regulations, namely the Council Directive 92/34/EEC and Commission Directive 92/34/EEC regarding the Conformitas Agraria Communitatis category of marketing plant material, states that olive plants must be free of all viruses as they are considered harmful. In order to examine the health status of olive orchards, appropriate and sensitive viral diagnostic tools have been developed for certification purposes, and cultivar improvement programmes have been initiated in several countries. In general, considerable levels of viral infection have been detected in many regions, including Portugal. In this country, extensive field surveys have revealed Olive latent virus 1 (OLV-1), Olive mild mosaic virus (OMMV) and Tobacco necrosis virus D (TNV-D) as predominant, reaching infection levels of 31%, with OMMV the most abundant virus (Saponari et al., 2002; Varanda et al., 2006, 2010; El Air et al., 2011). These viruses have several natural hosts. OLV-1 was first detected in olive (Gallitelli and Savino, 1985), then in citrus (Martelli et al., 1996), tulips (Kanematsu et al., 2001), and recently in tomato plants (Borodyynko et al., 2010). TNV-like viruses, that were able to be assigned to TNV-D or TNV-A, have been detected in herbaceous plants such as tobacco, tulip, bean, cucumber and potato, as well as woody crop plants such as pear and apple, citrus, grape (Kassanis, 1970; Kurstak, 1981) and olive (Félix and Clara, 2002b). OMMV was identified for the first time in olive (Cardoso et al., 2005) and genomic sequences of OMMV, published in the GenBank database, were recovered from tulips affected by the Augusta disease, leading to its identification as the causal agent of that disorder (Pham et al., 2009).

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Mixed infections of these three viruses occur naturally quite frequently (Félix and Clara, 2002a; Varanda et al., 2010). These alphapartetanecroviruses share important similarities. OMMV amino acid CP sequence shows 85.1% identity with that of TNV-D (Cardoso et al., 2009) and its RNA-dependent RNA polymerase (RdRp) has 91.2% identity with that of OLV-1 (Félix et al., 2005). In fact, differentiation between OMMV and TNV-D isolates is only possible through PCR-based assays using specific primers (Varanda et al., 2010) or genome sequencing.

The main purpose of this study was to design a single pair of degenerate primers to be used in a RT-PCR assay for a sensitive, rapid and reliable detection of either OLV-1, OMMV, TNV-D or TNV-A, applicable to plant health evaluation when the identification of the infecting species is not required, as is often the case of crop improvement programmes, which would additionally offer the opportunity for detecting the presence of new viruses or isolates by carrying out an extra sequencing step.

2. Materials and methods

OLV-1, OMMV, TNV-D and TNV-A isolates characterised previously (Félix et al., 2005; Cardoso et al., 2005, 2009; Meulenaer et al., 1998) were maintained in the experimental host plants Chenopodium murale and Nicotiana benthamiana, and used as positive controls. Inoculated plants were maintained for 2–3 weeks in a growth chamber with a 14h photoperiod at 22–24 °C. Total RNA was extracted from 100mg of symptomatic herbaceous plants using the RNeasy Plant Mini Kit (Qiagen, Hilden, Germany) in accordance with the manufacturer’s instructions.

DsRNA extracted from the fruits and stems of 54 olive field trees whose sanitary status concerning OLV-1, OMMV and TNV-D had already been determined (Varanda et al., 2010) were used in this study.

About 1μg of total RNA, or 1μg of denatured dsRNA, was used in a 20μL reaction for cDNA synthesis containing 200U of M-MLV Reverse Transcripase (Invitrogen, Carlsbad, USA) in the presence of random hexamers (Promega, Madison, USA) and 1× first strand buffer (Invitrogen, Carlsbad, USA), in accordance with the manufacturer’s instructions. For PCR amplification, two degenerative primers (parRdRp5′: 5′-CCCGCWCSCSMGGGTGKATWCAARC-3′ and parCoat3′: 5′-TATCCCCGATGWGGGAGAATK-3′) were designed based on conserved regions among published genomic sequences of OLV-1 isolates (OLV-1: NC_001721 and OLV-1*: DQ083996), OMMV wild type (NC_006939), TNV-D isolates (TNV-D: NC_00942 and TNV-D*: NC_033487) and TNV-A isolates (TNV-A: NC_001777 and TNV-A*: AY546104) (Fig. 1) in order to enable the PCR amplification of a single and similarly sized genomic fragment from any of these viral species.

The size of the RT-PCR fragment (parRdRpCoat) is predicted to be ca. 2035bp for OMMV, OLV-1 and TNV-A and ca. 2070bp for TNV-D, representing about 55% of the complete genomes.

One μl of obtained cDNA was used in PCR carried out in 20μM Tris–HCl pH 8.8, 10mM KCl, 10mM (NH₄)₂SO₄, 2mM MgSO₄, 0.1% Triton X-100, 0.1 mgml⁻¹ BSA, 0.2μM dNTPs, 0.5μM of each primer and 2.5μl of Taq DNA polymerase (Fermentas, Ontario, Canada) in a total volume of 50μl. Amplification was carried out in a Thermal Cycler (BioRad, Hercules, USA) at 94 °C for 1 min, 35 cycles at 94 °C for 1 min, 55 °C for 1 min and 72 °C for 2.5 min, and a final extension step of 72 °C for 10 min. Amplified products were analysed using agarose gel electrophoresis.

RT-PCR products were purified using GFX PCR DNA Purification kit (GE Healthcare Biosciences, Buckinghamshire, UK) and cloned into pGEM-T easy vector (Promega, Madison, USA), in accordance with the manufacturer’s instructions. Plasmid DNA was extracted from E. coli cells using GenEluteTM HP Plasmid Miniprep kit (Sigma, Missouri, USA) in accordance with the manufacturer’s instructions, after growing cells in low-salt LB medium (1% tryptone, 0.5% yeast extract, 0.5% NaCl, pH 7.5) supplemented with 100μgml⁻¹ of ampicillin and grown over night at 37 °C and shaken at 175 rpm. Ten clones from each cloned PCR product were sequenced in forward and reverse directions, by Macrogen (The Netherlands). Sequence analysis of the resulting amplicon was carried out using BioEdit (Hall, 1999).

3. Results

The expected RT-PCR amplification product (about 2000bp), using parRdRp5′ and parCoat3′ degenerate primers, was detected in herbaceous plants infected with any of the TNV-D, OLV-1, OMMV and TNV-A viruses used as positive controls (Fig. 2a). Sequencing results of the cloned amplicons confirmed the identity of each infecting virus.

The use of this test using the degenerate primers to dsRNA templates extracted from 54 olive trees RT-PCR tests demonstrated clearly that 17 of these trees were infected with the alpha- and betanecrovirus referred above (Fig. 2b). Cloning and sequencing these 17 PCR products revealed the presence of OMMV in 15 olive trees, OLV-1 in 12 trees and TNV-D in 4 trees, and mixed infections in 10 of them. This is in complete agreement with the findings of previous studies, in which single and multiplex RT-PCR assays were carried out using the same trees (Varanda et al., 2010). No TNV-A was detected in any of the sequenced clones.

4. Discussion

The present study shows that the use of the degenerate primers designed, in RT-PCR, results in an assay as sensitive and reliable in detecting the viruses under test as the assay using the three sets of primers specific for each virus. Yet it has additional advantages as it also detects TNV-A, it is more rapid, results are easier to interpret and contamination risks are lessened as only one pair of primers is used. Furthermore, since the assay developed results in a final amplicon that comprises a significant partial sequence of both RdRp and CP genes, it can be sequenced and further used for phylogenetic analysis, and enables the detection of isolate variability and a range of species with different sequences as well as other viruses.

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**Fig. 1.** Multiple sequence alignment showing the genome region of alpha- and betanecroviruses where degenerate primers were designed. The accession numbers of isolates used in the analysis are the following: NC_001721 (OLV-1), DQ083996 (OLV-1*), NC_006939 (OMMV), D00942 (TNV-D), NC_003487 (TNV-DH), NC_001777 (TNV-A), AY546104 (TNV-Ac).
previously unknown, as observed by other authors (Colinet et al., 1995). This is of particular interest for diversity and epidemiological studies of the viruses tested in this study, as it was found that such phylogenetic analysis generates the same tree topology as that which is based on the complete RdRp and CP sequences (data not shown).

The RT-PCR assay using the degenerate pair of primers is especially useful for virus free certification of O. europaea, as required by EU directive 93/48 concerning the Conformitas Agraria Communitatis plant material category, and provides support for sanitary selection and improvement programmes of olive or other crops affected by the alpha- and betanecroviruses tested in this study. In fact, this assay was found to be as sensitive, rapid and reliable as multiplex RT-PCR assays. Its application to epidemiological analysis is expected to contribute towards providing an understanding of the high level of dissemination of these viruses in nature.

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References


