

ARTIGOS

Development of a molecular method for detection and identification of *Xanthomonas campestris* pv. *viticola*

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ABSTRACT

Trindade, L.C.; Marques, E.; Lopes, D.B.; Ferreira, M.A.S.V. Development of a molecular method for detection and identification of *Xanthomonas campestris* pv. *viticola*. *Summa Phytopathologica*, v.33, n.1, p.16-23, 2007.

In order to develop a molecular method for detection and identification of *Xanthomonas campestris* pv. *viticola* (*Xcv*) the causal agent of grapevine bacterial canker, primers were designed based on the partial sequence of the *hrpB* gene. Primer pairs *Xcv*1F/*Xcv*3R and RST2/*Xcv*3R, which amplified 243- and 340-bp fragments, respectively, were tested for specificity and sensitivity in detecting DNA from *Xcv*. Amplification was positive with DNA from 44 *Xcv* strains and with DNA from four strains of *X. campestris* pv. *mangiferaeindicae* and five strains of *X. axonopodis* pv. *passiflorae*, with both primer pairs. However, the enzymatic digestion of PCR products could differentiate *Xcv* strains from the others. None of the primer pairs amplified DNA from grapevine, from 20 strains of nonpathogenic bacteria from grape leaves and 10

strains from six representative genera of plant pathogenic bacteria. Sensitivity of primers *Xcv*1F/*Xcv*3R and RST2/*Xcv*3R was 10 pg and 1 pg of purified *Xcv* DNA, respectively. Detection limit of primers RST2/*Xcv*3R was 10⁴ CFU/ml, but this limit could be lowered to 10² CFU/ml with a second round of amplification using the internal primer *Xcv*1F. Presence of *Xcv* in tissues of grapevine petioles previously inoculated with *Xcv* could not be detected by PCR using macerated extract added directly in the reaction. However, amplification was positive with the introduction of an agar plating step prior to PCR. *Xcv* could be detected in 1 µl of the plate wash and from a cell suspension obtained from a single colony. Bacterium identity was confirmed by RFLP analysis of the RST2/*Xcv*3R amplification products digested with *Hae* III.

Additional key words: PCR, grapevines, *Vitis vinifera*, bacterial canker

RESUMO

Trindade, L.C.; Marques, E.; Lopes, D.B.; Ferreira, M.A.S.V. Desenvolvimento de um método molecular para detecção e identificação de *Xanthomonas campestris* pv. *viticola*. *Summa Phytopathologica*, v.33, n.1, p.16-23, 2007.

Com o objetivo de desenvolver um método molecular para detecção e identificação de *Xanthomonas campestris* pv. *viticola* (*Xcv*), agente causal do cancro bacteriano da videira, oligonucleotídeos (*primers*) foram desenhados com base na seqüência parcial do gene *hrpB*. As combinações de *primers* *Xcv*1F/*Xcv*3R e RST2/*Xcv*3R que amplificaram fragmentos de 243 e 340 pb, respectivamente, foram testadas quanto à especificidade e sensibilidade para detecção do DNA de *Xcv*. Com os dois pares de *primers*, amplificação foi positiva com o DNA de 44 isolados de *Xcv*, mas também com quatro isolados de *X.c.* pv. *mangiferaeindicae* e cinco de *X. axonopodis* pv. *passiflorae*. Contudo, a digestão dos produtos de PCR permitiu diferenciar *Xcv* dos isolados desses patovares. Nenhum dos dois pares de *primers* amplificou o DNA de videira, nem de 20 bactérias não patogênicas isoladas da flora

da videira, ou de 10 isolados de outros seis gêneros de bactérias fitopatogênicas. A sensibilidade dos *primers* *Xcv*1F/*Xcv*3R e RST2/*Xcv*3R foi de 10 pg e 1 pg de DNA purificado de *Xcv*, respectivamente. O limite de detecção de RST2/*Xcv*3R foi de 10⁴ UFC/ml, mas empregando-se uma segunda rodada de amplificação com o *primer* interno *Xcv*1F, esse limite foi de 10² UFC/ml. Não foi possível detectar por PCR a presença de *Xcv* usando-se, diretamente na reação, o extrato do macerado de pecíolos de videira previamente inoculados. Entretanto, ampliações foram positivas quando se utilizou uma etapa de enriquecimento em meio de cultura antes da PCR. Detectou-se *Xcv* em 1 µl da suspensão obtida do lavado das placas e em uma suspensão obtida a partir de uma única colônia. A identidade da bactéria foi confirmada pela análise de RFLP dos produtos de amplificação dos *primers* RST2/*Xcv*3R com *Hae*III.

Palavras-chave adicionais: PCR, videira, *Vitis vinifera*, cancro bacteriano

Xanthomonas campestris pv. *viticola* (Nayudu) Dye (21) is the causal agent of grapevine canker disease, an important bacterial disease of grapevines in northeastern Brazil. This pathogen was first reported in the irrigated areas of the São Francisco river basin in Petrolina, state of Pernambuco (18). Later, the disease was also detected in Juazeiro, state of Bahia, and in the states of Piauí (17) and Ceará (8). *Xanthomonas campestris* pv. *viticola* is a gram-negative, non-pigmented bacterium which presents the following characteristics: aerobic metabolism, does not produce urease and oxidase but produces catalase, does not utilize asparagine as the sole source of C and N, tolerates up to 2% NaCl, and produces acid from glucose, mannose, galactose, trehalose, cellobiose and fructose (6,14,18,21). The disease is characterized by necrotic leaf spots, which sometimes coalesce, producing extensive necrosis. Symptoms also develop on leaf veins, petioles, pedicels and rachis of grape clusters. Cankers and vascular discoloration are usually observed on stems (14,18,21). Integrated management of bacterial canker can be achieved by the use of healthy propagative material, routine field inspections, drastic pruning of infected plants, roguing, management of production periods, disinfestation of vehicles, equipments and pruning tools, the use of copper and thiocarbamats as protecting fungicides, the use of windbreaks and localized irrigation to reduce bacterial dissemination (14,20).

Diagnosis of grapevine bacterial canker is based on symptoms developed on leaves, grape clusters and stems, followed by bacterial isolation and identification using biochemical and nutritional tests. Identification at the pathovar level depends on pathogenicity tests on grapevine susceptible varieties, with a period of 7-12 days for development of the first symptoms (14,18). Detecting and identifying *X. c.* pv. *viticola* in grapevine plant parts and/or propagative material, either symptomatic or asymptomatic, can be a time-consuming task with the currently available methods. Serological and molecular methods are more sensitive and specific and can improve the diagnosis of bacterial diseases. Polyclonal antibodies have already been developed for *X. c.* pv. *viticola* (3), and despite showing weak cross-reaction with *X. c.* pv. *campestris*, *X. c.* pv. *vesicatoria* and *X. c.* pv. *mangiferaeindicae*, they can potentially be used in serological tests for specific identification of *X. c.* pv. *viticola*.

Polymerase chain reaction (PCR) has been widely used and shown to have multiple applications in plant disease diagnosis, including several bacterial diseases (9,11,16,22,23). PCR-based detection methods offers several advantages such as high sensitivity to detect small amounts of DNA in infected or asymptomatic materials, without requiring organism culturing (19).

In plant pathogenic bacteria, several genomic regions have been explored in order to design primers for PCR-based detection, such as the ribosomal DNA spacer regions and the *hrp* gene cluster (16). The *hrp* genes have been discovered and characterized in several bacterial plant pathogens and they play a role in both hypersensitivity reaction and pathogenicity (15). *Hrp* genes are highly conserved in the genus *Xanthomonas* and they can be used to differentiate *Xanthomonas* strains both at pathovar and species levels. Leite Jr. *et al.* (12,13) designed specific primers based on the corresponding region of the *hrpB* gene for the detection and identification of *X. campestris* pv. *vesicatoria* by PCR. Roberts *et al.* (23) also demonstrated that the amplification and sequencing of a *hrp*-gene region allowed the selection of highly specific primers with high sensitivity for detecting *X. fragariae* in both symptomatic and asymptomatic strawberry plants. Specific primers based on *hrp* sequences are particularly useful for detection of

pathogenic xanthomonads, since nonpathogenic strains lack these genes (12).

Xanthomonas campestris pv. *viticola* is considered a pathogen of quarantine importance. The use of pathogen-free propagative material has become an important concern, considering the very restricted occurrence of this pathogen in the country and the risk of its establishment in other grapevine producing regions in south and southeast Brazil. Thus, the development of rapid and reliable procedures for detecting and identifying this pathogen is an important step for disease management in the affected areas, as well as for preventing its spread to other regions in the country. The objective of this work was to develop primers and to evaluate their potential for detection and identification of *X. c.* pv. *viticola* by PCR.

MATERIALS AND METHODS

Bacterial strains: origin and maintenance

Forty-one *Xanthomonas campestris* pv. *viticola* strains collected in grapevine producing areas in the "Submédio" of the São Francisco river valley, states of Pernambuco and Bahia, were used in this study. The strains were collected from 1998- 2003, from various grapevine cultivars. Two strains collected in Teresina, Piauí, and the reference strain of this bacterium (NCPBB 2475) were also included. Forty-two strains of *Xanthomonas* isolated from various host plants, as well as 10 strains belonging to six other genera of phytopathogenic bacteria, and 20 strains of epiphytic and/or endophytic bacteria isolated from grape leaves, were included to determine primer specificity (Table 1). All strains were cultivated on Kado's 523 medium (10). For long-term storage, strains were kept on sterile distilled water at room temperature and frozen in 30% glycerol at -80 °C.

DNA Extraction

DNA was extracted from grape leaves according to the protocol described by Doyle & Doyle (7). Bacterial DNA was extracted using a modified version of the CTAB method from Ausubel *et al.* (4). Single colonies grown on Kado's 523 medium were transferred to 1.5 ml of liquid 523 medium and cultures were grown on a shaker for 48 h at 28 °C. After this period, cultures were centrifuged at 4600x g for 5 min. The resulting pellets were resuspended in 520 µl of TE buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8.0). Fifteen microliters of 20% SDS and 3 µl of Proteinase K (20 mg/ml) were added. The mixture was incubated for 1 hour at 37°C, then 100 µl of 5 M NaCl and 80 µl of a 10% CTAB solution in 0.7 M NaCl were added and mixed. The suspension was incubated for 10 min at 65°C and kept on ice for 15 min. An equal volume of chloroform: isoamyl alcohol (24:1) was added, followed by incubation on ice for 5 min and centrifugation at 7200 x g for 20 min. The aqueous phase was transferred to a new tube, isopropanol (1: 0.6) was added and DNA was precipitated at -20 °C for 16 h. DNA was collected by centrifugation at 7200 x g for 10 min, washed with 500 µl of 70% ethanol, air-dried at room temperature for approximately three hours and finally dissolved in 50 µl of TE buffer. DNA was quantified on 0.8% agarose gels by comparison with DNA samples (phage λ DNA-*Hind* III fragments) of known concentration, diluted to working aliquots of 50 ng/µl and stored at -20 °C.

Amplification and sequencing of a *hrpB* region of *X. c.* pv. *viticola*

Primers RST2 and RST3 (12) were used to amplify a 840-bp

Table 1. Bacterial strains used in this study.

| Organism | Number of isolates | Host |
|--------------------------------------------------------------|--------------------|--------------------------------|
| <i>Xanthomonas campestris</i> pv. <i>viticola</i> | 44 | <i>Vitis vinifera</i> |
| <i>Agrobacterium tumefaciens</i> | 01 | <i>Daucus carota</i> |
| <i>Bacillus thuringiensis</i> | 01 | - |
| <i>Clavibacter michiganensis</i> subsp. <i>michiganensis</i> | 02 | <i>Lycopersicon esculentum</i> |
| <i>Curtobacterium flaccumfaciens</i> | 01 | <i>Beta vulgaris</i> |
| <i>Pseudomonas cichorii</i> | 01 | <i>Lactuca sativa</i> |
| <i>Pseudomonas corrugata</i> | 01 | <i>Lycopersicon esculentum</i> |
| <i>Ralstonia solanacearum</i> | 02 | <i>Dianthus caryophyllus</i> |
| <i>Ralstonia solanacearum</i> | 01 | <i>Solanum tuberosum</i> |
| <i>X. axonopodis</i> pv. <i>glycines</i> | 04 | <i>Glycine max</i> |
| <i>X. axonopodis</i> pv. <i>malvacearum</i> | 01 | <i>Hibiscus</i> sp. |
| <i>X. axonopodis</i> pv. <i>manihotis</i> | 07 | <i>Manihotis esculenta</i> |
| <i>X. axonopodis</i> pv. <i>passiflorae</i> | 06 | <i>Passiflora</i> sp. |
| <i>X. axonopodis</i> pv. <i>phaseoli</i> | 03 | <i>Phaseolus vulgaris</i> |
| <i>X. axonopodis</i> pv. <i>ricini</i> | 02 | <i>Ricinus communis</i> |
| <i>X. campestris</i> pv. <i>arracaciae</i> | 01 | <i>Arracacia xanthorrhiza</i> |
| <i>X. campestris</i> pv. <i>campestris</i> | 05 | <i>Brassica oleraceae</i> |
| <i>X. campestris</i> pv. <i>carotae</i> | 01 | <i>Daucus carota</i> |
| <i>X. campestris</i> pv. <i>mangiferaeindicae</i> | 01 | <i>Anacardium occidentale</i> |
| <i>X. campestris</i> pv. <i>mangiferaeindicae</i> | 03 | <i>Mangifera indica</i> |
| <i>X. campestris</i> pv. <i>vesicatoria</i> | 03 | <i>Lycopersicon esculentum</i> |
| <i>X. campestris</i> pv. <i>vesicatoria</i> | 03 | <i>Capsicum</i> spp. |
| <i>X. campestris</i> pv. <i>vitians</i> | 01 | <i>Lactuca sativa</i> |
| <i>X. melonis</i> | 01 | <i>Cucumis sativus</i> |
| Epiphytic and/or endophytic bacteria | 20 | <i>Vitis vinifera</i> |

fragment of the *hrpB* gene from genomic DNA of *X.c.* pv. *viticola* (NCPPB 2475), *X.c.* pv. *mangiferaeindicae* (UnB 764) and *X. axonopodis* pv. *manihotis* (UnB 1111). Each PCR mixture contained: 0.5 μ M of each primer; 200 μ M dNTPs; 1.5 mM MgCl₂; 1X PCR buffer (20 mM TrisHCl pH 8.4; 50 mM KCl); 1.25 U *Taq* DNA polymerase (Invitrogen, Life Technologies) and 50 ng template DNA in a final volume of 25 μ l. PCR was conducted in a PT-100 thermocycler (MJ Research, Watertown, Mass) using an initial denaturation at 95° C for 2 min, followed by 30 cycles of 95° C for 1 min, 62° C for 1 min, 72° C for 1 min, and an additional extension step at 72° C for 5 min. Specific PCR products from the three representative strains were quantified and adjusted to a final concentration of 70 ng/ μ l. After ethanol precipitation, purified PCR products were sequenced using the dideoxy chain-termination procedure (26) with fluorescent dyes (Dynamic ET terminator Cycle Sequencing Kit, Amersham Biosciences). In each sequencing reaction generating a single-strand template, total volume reaction of 10 μ l comprised 4 μ l of sequencing reagent premix, 2 μ l primer (10 μ M), 1 μ l PCR product (100 ng) and 3 μ l sterile water. Sequence reactions were carried out on a Gene Amp PCR System 9700 thermocycler (Applied Biosystems, USA) using 30 cycles of 95° C for 20 s, 50° C for 15 s, and 60° C for 1 min. Each product was purified via a standard ethanol precipitation protocol and dissolved in 10 μ l of Ddynamic ET loading solution (Pharmacia Biotech, USA) for 60 min. Automated sequencing was conducted using a MegaBace 1.000 – DNA Analyzer System (Pharmacia Biotech,

USA) with an injection voltage of 3 KV, an injection time of 60 s, and a running voltage of 9 V. Sequences were verified via BLASTN (1), and alignment conducted using the program Clustal W (29).

Primer design and selection

Potential primers were identified using program PRIMER 3 (Whitehead Institute for Biomedical Research, Cambridge, Mass.) (24). Three primers were designed and tested in five possible combinations with external primers RST2 and RST3. Primers were synthesized commercially by Invitrogen Brasil (Life Technologies, São Paulo, Brazil). The following criteria were used to choose the best primer pair combination: absence of secondary bands, reproducibility, and positive amplification with all *X.c.* pv. *viticola* strains tested.

Primer specificity

Two out of five possible primer pair combinations (Xcv1F/Xcv3R and RST2/Xcv3) were screened for specificity to *X.c.* pv. *viticola* in 12 μ l- reaction mixtures containing 10 ng of template DNA, 100 μ M each dNTPs, 1.5 mM Mg Cl₂, 0.5 μ M of each primer, 1X PCR buffer (20 mM TrisHCl pH 8.4; 50 mM KCl) and 1 U *Taq* DNA polymerase. All tests were performed in a PT 100 thermocycler (MJ Research, Watertown, Mass) with an initial denaturation step at 95° C for 2 min; 30 cycles of 95° C for 1 min, 64° C for 1 min and 72° C for 2 min, and a final extension step of 10 min at 72° C. PCR products were analyzed by loading the total reaction volume on 1% agarose gels and staining

with ethidium bromide (0.5 µg/ml).

PCR-RFLP

PCR products from *Xanthomonas* were digested with endonuclease *Hae* III (Pharmacia Biotech). Digestions were performed as follows: 6 µl PCR product; 1 µl 10X enzyme buffer and 2U of enzyme in a final volume of 10 µl, at 37°C for 16 h. Restriction fragments were separated by electrophoresis in 2.5% agarose gels or in 21% polyacrylamide gels and visualized after staining with ethidium bromide.

Primer sensitivity

The limit of detection of the *X.c. pv. viticola* specific fragment was determined by using a 10-fold serial dilutions of purified genomic DNA of strains NCPPB 2475 and UnB 1183, and a 10⁹ CFU/ml suspension culture from strain NCPPB 2475. Aliquots of 2 µl and 0.05 ml were used for PCR amplification and colony counts on 523 medium, respectively.

Detecting *X.c. pv. viticola* in symptomatic plant tissue

Grapevine plants (cv. Perlette) were inoculated by pricking the petiole surface with a sterile needle charged with a 2-day-old culture of strain UnB 1186. Three petioles were inoculated per plant and they were covered with plastic bags for 48 h after inoculation to maintain high humidity. Plants were kept under a shaded greenhouse and observed daily for symptom development. After 41 days, fragments of petioles were collected and used for PCR detection with primer pair RST2/Xcv3R. The presence of *X.c. pv. viticola* in inoculated petioles, was investigated using three different methods as follow: (1) direct detection by macerating plant tissue in 500 µl water and using 1 µl for PCR amplification; (2) using a modified BIO-PCR protocol (27), in which aliquots of macerated tissue extracts were plated on 523 medium and after 72 h the surface of the medium was washed with 1 ml of sterile water, and 1 µl of the plate wash was used for PCR, and (3) macerated tissue extracts were plated on 523 medium, after 72 h one single suspect colony was collected in 200 µl of water and 1 µl was used for PCR.

RESULTS AND DISCUSSION

The amplification of a *hrpB*-gene region from *X.c. pv. viticola* and other xanthomonads was reproducible and produced an amplicon of approximately 840 bp for all strains tested. PCR products were partially sequenced and three primers were designed (Figure 1). These primers were tested in five combinations with previously designed primers RST2 and RST3 (12): (1) RST2/Xcv3R; (2) RST2/Xcv2R; (3) Xcv1F/Xcv3R; (4) Xcv1F/Xcv2R, and (5) Xcv1F/RST3. Combinations (2) and (4) did not yield any amplified product, while combination (5) produced a fragment of expected size (~700 bp), but also several nonspecific secondary bands. Primer combinations (1) and (3) gave the expected amplicons of 340 and ~240 bp, respectively, with DNA from *X.c. pv. viticola* (Figure 1). Therefore, these two latter sets of primers were selected and tested for specificity.

Amplification was positive for all 44 *X.c. pv. viticola* strains tested with these two sets of primers, RST2/Xcv3R and Xcv1F/Xcv3R. However, amplification was also positive with DNA from four *X.c. pv. mangiferaeindicae* strains and from five strains of *X. a. pv. passiflorae*, as detected on ethidium-bromide stained agarose gels (Figure 2). Specificity of both primer pairs were also tested with genomic DNA from strains of other six genera of phytopathogenic

bacteria, grape leaves and a total of 20 epiphytic and/or endophytic bacteria isolated from grape leaves, and no detectable product was observed (Figure 2).

A differentiation of the strains of *X.c. pv. viticola* from the strains of *X.c. pv. mangiferaeindicae* and *X.a. pv. passiflorae* was performed with an additional restriction digestion step of the PCR products. Polymorphism was detected among the strains belonging to the three pathovars. *Hae*III digestion of RST2/Xcv3R-amplified products generated distinct banding patterns for each pathovar. Strains of *X.c. pv. viticola* yielded 5 bands with approximately 100, 70, 60, 55 and 25 bp, while strains of *X.c. pv. mangiferaeindicae* showed two distinct restriction profiles and *X.a. pv. passiflorae* strains showed only one profile, with 3 fragments of approximately 250, 65 and 45 bp (Figure 3).

Although they can be considered semi-specific since amplification was also positive with *X.c. pv. mangiferaeindicae* and *X.a. pv. passiflorae*, it is unlikely that strains of these pathogens would be found infecting grapevines under field conditions. Moreover, the size of the amplified RST2/Xcv3R product is suitable for restriction digestion and visualization in agarose gels, if additional confirmation is required. All *X.c. pv. viticola* tested strains produced the same restriction profile, distinct from the other two pathovars. Taxonomic and phylogenetic relationships among *X.c. pv. viticola* and these two pathovars have not been investigated. Interestingly, pathovars *viticola* and *mangiferaeindicae* share many similarities: they were both first described in India, some strains are not yellow pigmented and they have similar disease cycles and epidemiology. Their relationship should be more carefully studied.

The detection limit for primer pair Xcv1F/Xcv3R was 10 pg of purified DNA. Primer pair RST2/Xcv3R was more sensitive, and allowed the detection of 1 pg of bacterial DNA per reaction (Figure 4A). The same level of sensitivity was reported for a PCR assay developed for *Erwinia amylovora* (5), and for *Xanthomonas albilineans* (22), but for *X. oryzae pv. oryzae*, a visible band was detected with as low as 55 fg of purified DNA (25). When serial dilutions of bacterial cell suspensions were used directly in the reaction (from 10⁹ to 10 CFU/ml) a detectable product was visible up to 10⁴ CFU/ml (Figure 4B), which corresponded roughly to 20 cells per reaction tube. A 1:50 dilution of the PCR products obtained with primers RST2/Xcv3R was used in a second round of amplification with the internal primer Xcv1F and primer Xcv3R. This nested-PCR approach increased sensitivity 100-fold, and as low as 10² UFC/ml could be detected (Figure 4C). This corresponded roughly to less than one cell in the reaction. Levels of sensitivity as high as 1 CFU or less per reaction have been reported (16). High primer sensitivity is important for direct detection in plant tissue and for detecting bacteria in latent infections when pathogen population is low (below 10² CFU/ml) (5).

When primers RST2 and Xcv3R were used to detect *X.c. pv. viticola* in artificially inoculated petioles, amplification was positive only when an additional growth step was included. The bacterium could not be detected when suspensions were prepared from macerated tissue and used directly in the reaction mixture, probably due to the presence of PCR inhibitors. Lower sensitivity and inhibitors problems can be circumvented with the addition of an enrichment step, such as BIO-PCR, which has the advantage that only viable cells multiply and give a positive amplification signal (27). Moreover, sensitivity of BIO-PCR assays is extremely high, detecting as few as 1-2 CFU/ml (28).

In this study, an enrichment step was employed but bacterial

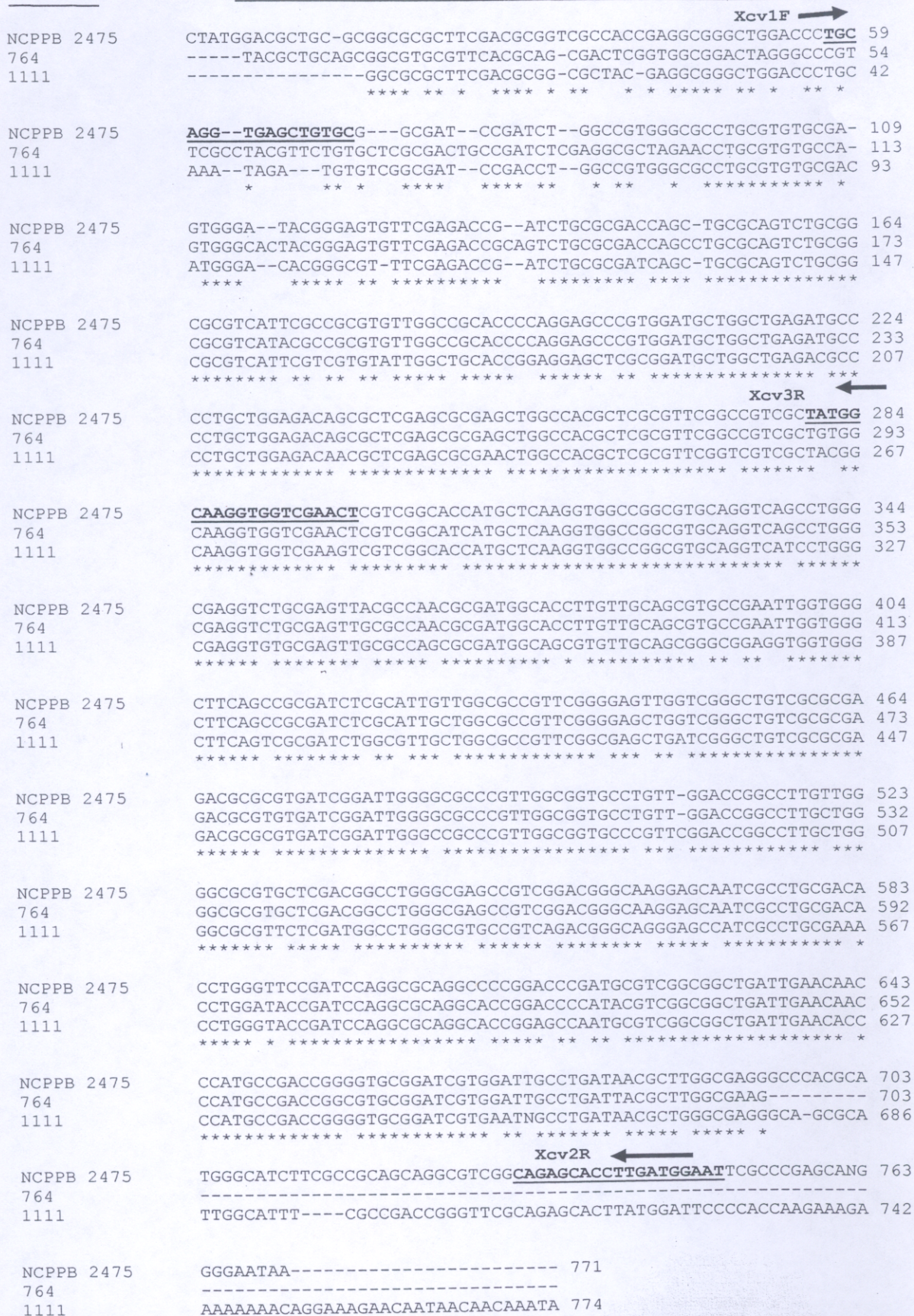


Figure 1. Partial sequence alignment of the *hrpB* gene amplified with primers RST2 and RST3 from *Xanthomonas campestris* pv. *viticola* (NCPPB 2475), *X. campestris* pv. *mangiferaeindicae* (UnB 764) and *X. axonopodis* pv. *manihotis* (UnB 1111). Nucleotide identity is indicated by *; (—) alignment gaps and deletions; underlined and bold sequences indicate the three primers designed and synthesized in this study.

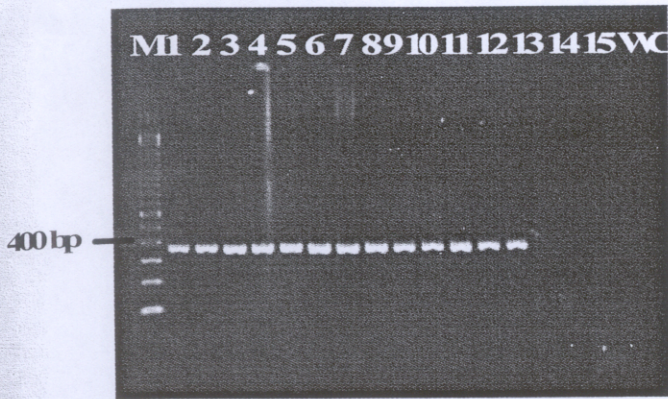


Figure 2. PCR amplification with primer pair RST2/Xcv3R, visualized on a 1% agarose gel. M - 100 bp-ladder (GIBCO/BRL); lanes (1)- NCPPB 2475; (2) - UnB - 1186; (3)- UnB 1187; (4)- IBSBF 1369; (5)- UnB 682; (6)- UnB 675; (7)- UnB 674; (8)- UnB 684; (9) - UnB 676; (10)- UnB 764; (11)- UnB 769; (12)- IBSBF 1230; (13)- IBSBF 1508; (14)- DNA from grape leaves; (15)- *Ralstonia solanacearum* and WC- negative water control. Lanes 1 - 4: *Xanthomonas campestris* pv. *viticola*; Lanes 5- 9: *X. axonopodis* pv. *passiflorae*, lanes 10-13: *X.c.* pv. *mangiferaeindicae*.

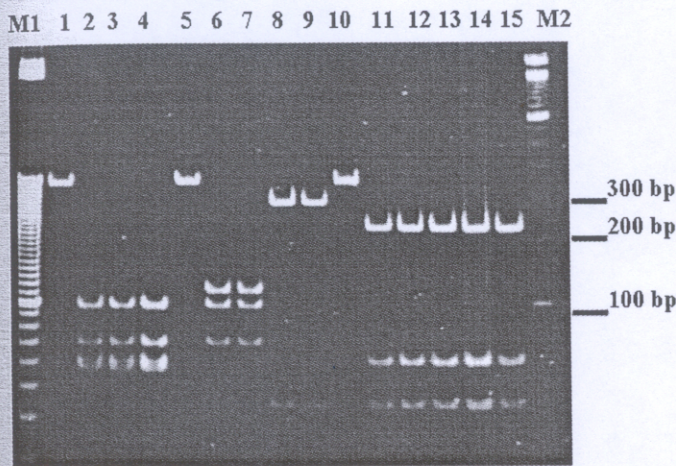


Figure 3. *Hae*III- restriction digestion of PCR products amplified with primers RST2/Xcv3R separated on 21% polyacrylamide gel. M1: 10 bp-ladder; M2: 100 bp ladder; lanes (1)-NCPPB 2475 - undigested product; (2)-NCPPB 2475; (3)- UnB 1186; (4)- UnB 1187; (5)- UnB 764 - undigested product; (6)- UnB 764; (7)- UnB 769; (8)- IBSBF 1230; (9)- IBSBF1508; (10)- UnB 682 - undigested; (11)- UnB 682; (12)- UnB 675; (13)- UnB 674; (14)- UnB 684; (15) - UnB 676. Lanes 1 - 4: *Xanthomonas campestris* pv. *viticola*; lanes 5- 9: *X.c.* pv. *mangiferaeindicae*; lanes 10-15: *X. axonopodis* pv. *passiflorae*.

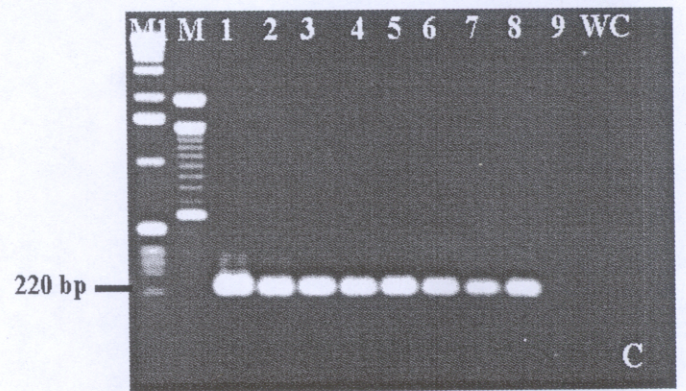
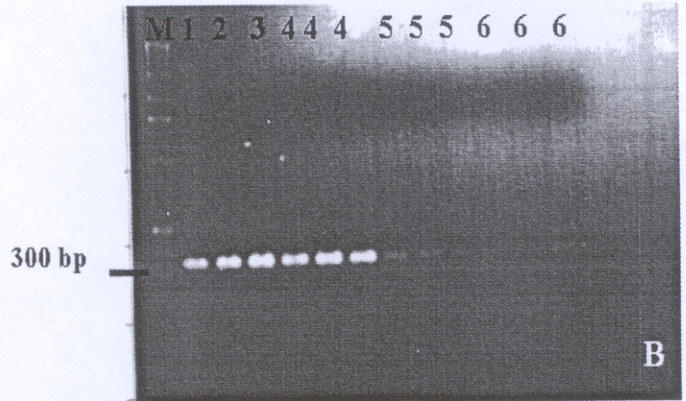
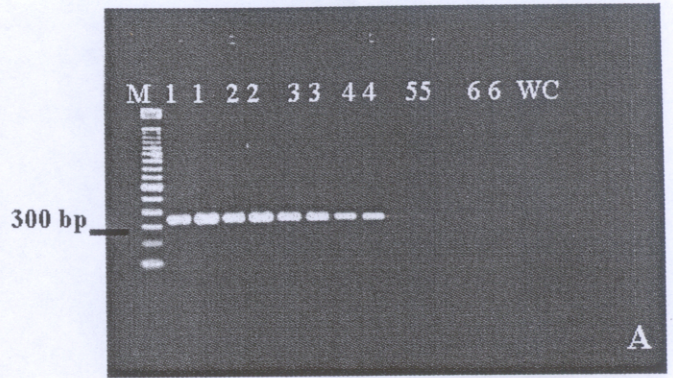


Figure 4- Detection limit of *Xanthomonas campestris* pv. *viticola* DNA amplified with primers RST2/Xcv3R. (A)- Dilution series of DNA from strain NCPPB 2475. M-100 bp ladder; Lane 1- 10 ng; 2- 1 ng; 3- 0,01 ng; 4- 0,001 ng; 5- 0,0001 ng; 6- 0,00005 ng; WC - water control. (B)- Dilution series of a cell suspension of strain NCPPB 2475. Lane 1- 10^9 CFU/ml; 2- 10^8 CFU/ml; 3- 10^7 CFU/ml; 4- 10^6 CFU/ml; 5- 10^5 CFU/ml; 6- 10^4 CFU/ml.(C)- Nested-PCR with primers Xcv1F/Xcv3R. PCR products obtained with primers RST2/ Xcv3R (Fig. 4B) were diluted (1:50) and used as templates. M1- 1 Kb ladder; M- 100 bp ladder; lane 1- 10^9 CFU/ml; 2- 10^8 CFU/ml; 3 - 10^7 CFU/ml; 4- 10^6 CFU/ml; 5- 10^5 CFU/ml, 6- 10^4 CFU/ml, 7- 10^3 CFU/ml, 8- 10^2 CFU/ml, 9- 10 CFU/ml; WC- water control.

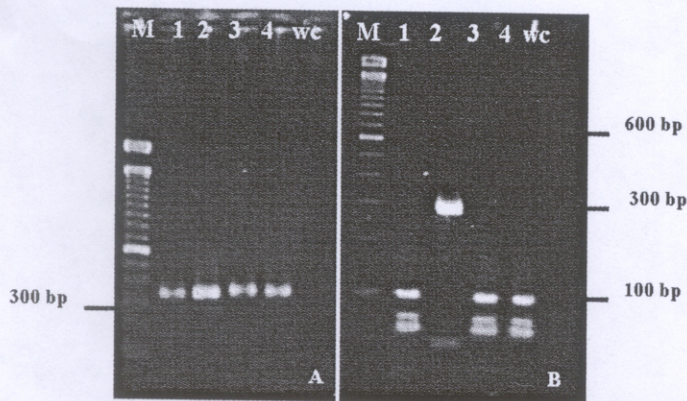


Figure 5. PCR-detection and identification of *Xanthomonas campestris* pv. *viticola* (*Xcv*) in inoculated and symptomatic grapevine petioles. (A) PCR with primers RST2/ Xcv3R: M- 100 bp DNA ladder; lane 1- *Xcv* UnB 1186 (purified DNA); lane 2- *X.c. pv. mangiferaeindicae* IBSBF 1230 (purified DNA); lane 3- agar plate-washings from macerated tissue after 72 h incubation; lane 4 - one single colony of *Xcv* grown after 72 h; and wc- water control. (B)- *Hae*III- restriction digestion of PCR products amplified with primers RST2/Xcv3R separated on a 2,5% agarose gel. Same samples as in 5A.

colonies were only recovered from symptomatic petioles. No bacterial colonies were recovered from uninoculated petioles or from inoculated asymptomatic petioles, thus PCR could not be performed with samples from asymptomatic tissue. PCR was tested with aliquots of the plate wash obtained from symptomatic tissue and using a cell suspension from a single suspect colony. In both cases amplifications were positive (Figure 5A). Bacterium identity was confirmed by *Hae*III digestion of PCR products, which produced the typical *X.c. pv. viticola* restriction profile (Figure 5B) and by observation of some cultural and biochemical characteristics of the colonies. Lack of yellow pigmentation, a positive reaction in the potassium hydroxide (3% KOH) solubility test, and lack of growth on asparagin medium were all in accordance with the PCR-based identification assay.

Total time required for *X.c. pv. viticola* PCR-based detection and identification was 3 to 4 days, which was an advantage of the method when compared to the conventional techniques (at least 10 days for isolation, culturing, and identification by traditional bacteriological and pathogenicity tests). Although PCR has proven to be a very useful method for disease diagnosis and detection of various pathogens, potential problems such as contamination, false negatives and higher cost compared to serological detection, must be considered before adapting protocols for routine or large-scale testing (19).

Plant pathogenic bacteria, represented by diverse populations in the environment, often require complementary and multiple tests for identification (2). A PCR- detection and identification method could be very useful as an additional tool for monitoring *X.c. pv. viticola*-contaminated plant material. In this study, primers targeted to a pathogenicity gene were shown to specifically amplify DNA from *X.c. pv. viticola* and did not amplify host plant DNA or bacterial DNA from grapevine microflora. Our results showed that PCR could be used to detect and identify the pathogen in symptomatic plant tissue. Evaluation of PCR to detect *X.c. pv. viticola* in different plant parts as well as in asymptomatic grapevines will be the subject of future studies.

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