

RESEARCH ARTICLE

Development of a new probe for specific and sensitive detection of '*Candidatus Phytoplasma mali*' in inoculated apple trees

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Keywords

'*Candidatus Phytoplasma mali*'; detection; real-time PCR; sensitivity.

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Received: 7 December 2006; revised version accepted: 11 June 2007.

doi:10.1111/j.1744-7348.2007.00171.x

Abstract

A new TaqMan minor groove binding (MGB) probe and new PCR conditions were designed for quick, specific and sensitive detection of '*Candidatus Phytoplasma mali*'. The new probe can distinguish a single mismatch between '*Ca. P. mali*' and '*Candidatus Phytoplasma prunorum*', this constituting an improvement over a previously published method. In our study, the relative position of the mismatch in the MGB probe influenced greatly the specificity of detection. Our new real-time PCR protocol was able to detect one plasmid copy in water and 100 copies in healthy plant DNA extracts. The sensitivity of this new real-time PCR method, three other real-time PCR protocols and a conventional PCR with fU5/rU3 primers was compared. Periwinkles and MM106 rootstocks were grafted with infected material and surveyed over time by symptom observation, conventional PCR and real-time PCR. Phytoplasma infection was detected by symptom observation in all periwinkles within 4 months and in 75% of the MM106 rootstocks by the end of 7 months. Conventional PCR detected phytoplasma infection in all periwinkles within 4 months and in all MM106 rootstocks within 7 months. Best results were obtained by our real-time PCR, which detected phytoplasma infection in all grafted plants within 3 months after inoculation.

Introduction

Phytoplasmas are prokaryotes inhabiting phloem sieve elements of infected plants (McCoy *et al.*, 1989; Lee *et al.*, 1995). '*Candidatus Phytoplasma mali*' (Seemüller & Schneider, 2004), classified as an A2 pest by European and Mediterranean Plant Protection Organization (EPPO), is a quarantine organism (EPPO/CABI, 1996) associated with one of the most serious diseases of apple trees, apple proliferation disease, in most countries of western Europe (Frisinghelli *et al.*, 2000). Typical symptoms are witches' broom, early leaf colouration in autumn and small size and incomplete colouration of fruit. This phytoplasma belongs to the 16SrX group, which also includes '*Candidatus Phytoplasma pyri*' and

'*Candidatus Phytoplasma prunorum*' (Seemüller *et al.*, 1998). Phylogenetic analyses have revealed that these pathogens have nearly identical 16S rDNA sequences (Seemüller & Schneider, 2004). Restriction fragment length polymorphism analysis of chromosomal DNA has demonstrated their relationship (Lee *et al.*, 1998; Aldaghi *et al.*, 2005).

Candidatus Phytoplasma mali being a quarantine organism, its accurate and rapid detection is a major prerequisite to controlling the disease and fulfilling the certification requirements. Yet the relatively low titres and uneven distribution of phytoplasmas in plant hosts (Andersen *et al.*, 1998) and their inability to be cultured *in vitro* make it hard to develop an accurate and reliable diagnostic procedure.

Real-time PCR provides a straightforward method for sensitive and specific detection of phytoplasmas (Jarusch *et al.*, 2004). Recently, Torres *et al.* (2005) developed a polyvalent real-time PCR protocol based on SYBR Green technology for simultaneous detection and quantification of the three phytoplasmas of the 16SrX group. SYBR Green technology has also been used to quantify *Ca. P. mali* in plants and insect vectors (Jarusch *et al.*, 2004). Baric & Dalla-Via (2004) have developed a new approach to specific detection of *Ca. P. mali* in plant material: a real-time PCR assay using a TaqMan minor groove binding (MGB) probe. The MGB moiety stabilise probe hybridisation with single-stranded DNA targets, raising the melting temperature of the oligodeoxynucleotide. This property allows the use of shorter oligodeoxynucleotide (Kutyavin *et al.*, 2000). TaqMan MGB probes display higher specificity than classical TaqMan probes (Yao *et al.*, 2006). Successful discrimination of a single mismatch has been achieved by various authors (Marbot *et al.*, 2002; Salmon *et al.*, 2002a; Yoshitomi *et al.*, 2003; Van Hoeyveld *et al.*, 2004; Massart *et al.*, 2005). Yet, recent studies (Yao *et al.*, 2006) have shown that a TaqMan MGB probe may tolerate one or two mismatches with its template and cause a nonspecific fluorescence emission.

The aim of the present study was to develop a new real-time PCR methodology for specifically detecting *Ca. P. mali* infection in apple trees. The specificity of the methodology described by Baric & Dalla-Via (2004) was improved by designing a new MGB probe and new PCR conditions. The initial and improved methods were compared with other existing conventional and real-time PCR methods regarding their sensitivity.

Materials and methods

Phytoplasma isolates, biological materials and indexing

The phytoplasma isolates, their host plants and their origins are listed in Table 1. Isolates of *Ca. P. mali*, '*Ca. P. pyri*', and '*Ca. P. prunorum*' received in infected periwinkles [*Catharanthus roseus* (L.) G. Don] were maintained in periwinkles by periodic grafting, and obtained apple plant materials containing the isolates of *Ca. P. mali* were grafted on apple trees (variety: MM106) by whip or bark grafting in an insect-proof greenhouse. After grafting, suitable conditions for growth were provided: 14 h of photoperiod, high relative humidity and temperature of 20–25°C). For biological indexing, the appearance of symptoms was timed by monitoring inoculated plants (three periwinkles and four apple trees per isolate).

Table 1 Phytoplasmas and host plants examined in this study

Strain	Sample Form	Provider
AT1	Periwinkle infected	Seemüller (Germany)
AT	Periwinkle infected	Schneider (Germany, accession no. X68375)
AP15	Periwinkle infected	Schneider (Germany, accession no. AJ542541)
AT1/93	Periwinkle infected	Schneider (Germany, accession no. AJ542542)
AT5/93	Periwinkle infected	Schneider (Germany)
AT12/93	Periwinkle infected	Schneider (Germany)
AT1-proliferation	Apple infected	Pradier (Lempdes, France)
AT1-No. 2	Apple infected	Pradier (Lempdes, France)
AP-N17	Apple infected	Pradier (Lempdes, France)
AT1-IDARED	Apple infected	Pradier (Lempdes, France)
AT2-SO8D	Apple infected	Pradier (Lempdes, France)
LB/AP1	Extracted DNA	Baric (Bolzano, Italy)
LB/AP2	Extracted DNA	Baric (Bolzano, Italy)
AP2174	Liophilised DNA	Fialova (Olomouc, Czech Republic)
AP2371	Lyophilised DNA	Fialova (Olomouc, Czech Republic)
AP-Melo1	Lyophilised DNA	Firrao (Udine, Italy)
AP-Melo2	Lyophilised DNA	Firrao (Udine, Italy)
ESFY	Periwinkle infected	Seemüller (Germany)
ESFY1	Extracted DNA	Bertaccini (Italy, accession no. AJ542544)
ESFY2	Extracted DNA	Bertaccini (Italy, accession no. AJ542545)
PD	Extracted DNA	Bertaccini (Italy, accession no. AJ542543)
PD1	Periwinkle infected	Seemüller (Germany)
EY (16SrV)	Extracted DNA	Bertaccini (Italy, accession no. X68376)
ASLO (16SrXII-A)	Extracted DNA	Bertaccini (Italy)
KVM (16SrI-C)	Extracted DNA	Bertaccini (Italy)

Phytoplasma DNA extraction

Fresh apple tree leaf midribs (0.5 g) were ground in 4-mL extraction buffer (2% cetyltrimethylammonium bromide, 1.4 M NaCl, 20 mM ethylenediaminetetraacetic acid, 100 mM Tris-HCl, pH 8.0) with a mortar and pestle. Samples were incubated at 60°C for 30 min and shaken every 5 min. Three millilitres of chloroform/isoamyl alcohol (24:1) were added on ice, and the mixture was vortexed vigorously before centrifugation at 16 100 *g* for 5 min at 4°C. The supernatant was transferred to a clean tube, and after adding an equal volume of ice-cold (–20°C) isopropanol, the solution was centrifuged at 16 100 *g* for 8 min at 4°C. The pellet was rinsed with 70% ethanol and dried. An optional ethanol precipitation was also carried out. If required, the total nucleic acid concentration was estimated with a spectrophotometer (LKB Biochrom Ultraspec II, Cambridge, UK).

Detection of phytoplasma by conventional PCR

The fU5/rU3 primer pairs and the protocol described by Lorenz *et al.* (1995) were used to detect phytoplasma by classical PCR. Two adaptations were made: an additional step at 95°C for 10 min (polymerase activation) and amplification by 45 PCR cycles. Primers were obtained from Eurogentec (Liège, Belgium). The 1× PCR buffer and the HotGoldStar DNA polymerase (Eurogentec) were used. PCR products were electrophoresed through 1.5% agarose gel and revealed using ethidium bromide under UV illumination.

Cloning and sequencing of PCR products

The PCR products obtained with the qAP-16S-F/R primers (Baric & Dalla-Via, 2004; Table 2) were purified from the agarose gel using the QIAEX II purification kit (Qiagen, Hilden, Germany) as recommended by the manufacturer. These fragments were further cloned with the TA Cloning[®] Kit (Invitrogen, Carlsbad, CA, USA). Plasmids containing the PCR product were extracted according to the ULTRAPrep[®] PLASMID DNA kit protocol (AHN Biotechnologie, Nordhausen, Germany). The amplified plasmids were either sequenced (by GATC Biotech, Konstanz, Germany) or quantified by optical density at 260 nm. After quantification, 10-fold serial dilutions of the purified plasmid, ranging from 1 to 10⁸ copies per PCR, were prepared to evaluate the limit of detection of the method.

Oligonucleotide design

A new TaqMan MGB probe [with the carboxyfluorescein (FAM) reporter dye] was designed using the 'Primer Express' software version 5.0 (PE Applied Biosystems, Foster City, CA, USA). The design of this new probe was based on our own sequences, corresponding to isolates AP15 (DQ661859) and AT1 (DQ661860) of *Ca. P. mali*, and based on at least 20 sequences from the NCBI database [six *Ca. P. mali* isolates: X76426, AJ542542, AJ54254, X68375, AY598319 and AF248958; six *Ca. P. prunorum* isolates: AY029540, AJ542544, AJ542545, AJ575105, AJ575106 and AJ575107; three isolates of *Ca. P. pyri*: Y16392, AJ542543 and AJ964959 and some other phyto-

plasmas more distinct from AP group including '*Candidatus Phytoplasma ulmi*' (X68376), '*Candidatus Phytoplasma asteris*' (AF503568), '*Candidatus Phytoplasma solani*' (AF248959), '*Candidatus Phytoplasma vitis*' (X76560), and strain CPh of '*Ca. P. asteris*' (X77482)].

Detection of phytoplasma by real-time PCR using TaqMan MGB probe

The reactions were carried out in a total volume of 25 µL containing 10–100 ng template DNA, 1× qPCR Master Mix (Eurogentec) containing dUTP, 400 nM of each primer and 200 nM probe. Amplification and detection were performed with the GeneAmp 5700 Sequence Detection System (SDS; PE Applied Biosystems). The thermal cycle consisted of 2 min at 50°C (carry-over inactivation) and 10 min at 95°C (HotGoldStar DNA activation), followed by 45 cycles of 15 s at 95°C and 60 s at 60°C, 62°C, or 64°C. Each reaction included at least one blank without template and one negative control corresponding to healthy plants. All samples and controls were amplified in triplicate. Threshold levels were calculated according to two recommended methods: (a) threshold = 10 × standard deviation (SD) of the fluorescence values during 5th to 15th cycle and (b) point of inflexion of the fluorescence curve. The threshold cycle (*C_t*) value of each PCR reaction was calculated by GeneAmp 5700 SDS software.

Specificity and limit of detection

The specificity of the real-time PCR using the newly designed probe for *Ca. P. mali* was assessed using DNA extracts containing other European fruit tree phytoplasma isolates (two *Ca. P. pyri* and four *Ca. P. prunorum*) or other more different phytoplasmas (Table 1). The limit of detection of our real-time PCR method was compared with that of four other PCR protocols: one conventional PCR (Lorenz *et al.*, 1995) and three real-time PCRs using TaqMan MGB (Baric and Dalla-Via, 2004) or SYBR Green (Jarusch *et al.*, 2004; Torres *et al.*, 2005) fluorescence chemistry. Forty-five PCR cycles were performed with all protocols. The SYBR Green Mastermix used in the original protocols (Jarusch *et al.*, 2004; Torres *et al.*,

Table 2 List of primers and probes used for phytoplasma detection

Name	Sequence 5'–3'	Reference
qAP-16S-F	CGAACGGGTGAGTAACACGTAA	Baric & Dalla-Via (2004)
qAP-16S-R	CCAGTCTTAGCAGTCGTTTCCA	Baric & Dalla-Via (2004)
qAP-16S	FAM-TAACCTGCCTCTTAGACG-MGB	Baric & Dalla-Via (2004)
AP-MGB	FAM-CTGCCTCTTAGACGAGG-MGB	Designed in this study

MGB, minor groove binding.

2005) was employed. Three samples, from three different apple trees, were extracted independently from leaves (A and C) or vascular tissue (B). Ten-fold serial dilutions (10^{-1} to 10^{-8}) were prepared from each sample. Each dilution was amplified twice independently in triplicate.

Results

Real-time PCR and new probe design

In our experiments, target DNA from all *Ca. P. mali* isolates (Table 1) was successfully detected with the qAP-16S probe (Table 2) and a hybridisation temperature of 60°C (Baric and Dalla-Via, 2004). No signal was obtained from healthy plants (apple or periwinkle), but significant fluorescence signals were also detected for *Ca. P. prunorum*, and *Ca. P. pyri* isolates. The amplification curves of the *Ca. P. prunorum* samples were sigmoid like those of the *Ca. P. mali* samples and crossed the threshold line set by either the $10 \times$ SD or the inflexion point method. A positive signal was still observed for *Ca. P. prunorum* samples with the hybridisation temperature of 62°C or

64°C, but the amplification curves were not completely sigmoid and the final fluorescence was much lower (Fig. 1).

Therefore, a sequencing effort and a sequence alignment (Fig. 2) were carried out to select a new TaqMan MGB probe (AP-MGB). Sequence alignments showed that the fragments amplified from our *Ca. P. mali* strains were 100% identical to the previously published *Ca. P. mali* sequences. They also revealed that the *Ca. P. prunorum* isolates used in this study had only one mismatch with the qAP-16S probe (nucleotide at position 36 in Fig. 2). Our new probe was designed to locate the mismatch in the middle part of the probe (Fig. 2). This probe displayed no mismatch with any of the six *Ca. P. mali* sequences. It did display mismatches with the *Ca. P. prunorum* sequences (one or two mismatches), *Ca. P. pyri* sequences (three mismatches), other more distant phytoplasmas (at least two mismatches) and also with unspecific amplicons obtained from healthy plants (seven mismatches, Fig. 2). With the AP-MGB probe and a hybridisation temperature of 64°C, specific detection of *Ca. P. mali* was achieved (Fig. 1) without cross-amplification of DNA from any other phytoplasma

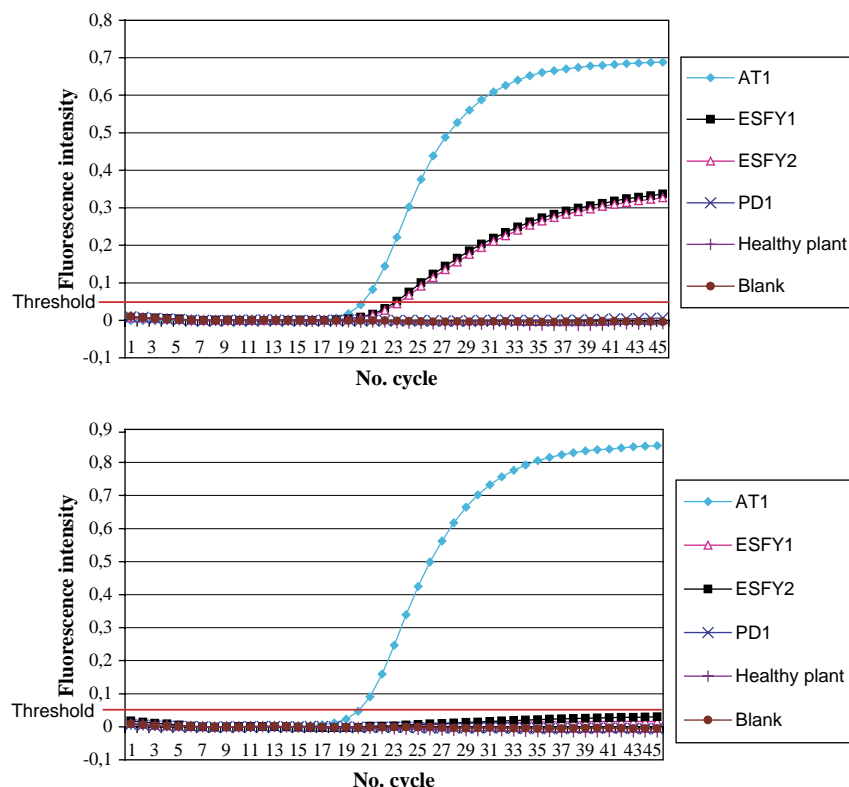


Figure 1 Fluorescent intensity (delta Rn) of amplification of different phytoplasma isolates, '*Ca. P. mali*' (AT1), '*Ca. P. prunorum*' (ESFY1 and ESFY2) and '*Ca. P. pyri*' (PD1) obtained by real-time PCR using qAP-16S-F/R primers and qAP-16S (upper diagram) or AP-MGB (lower diagram) probes at 64°C as hybridisation temperature.

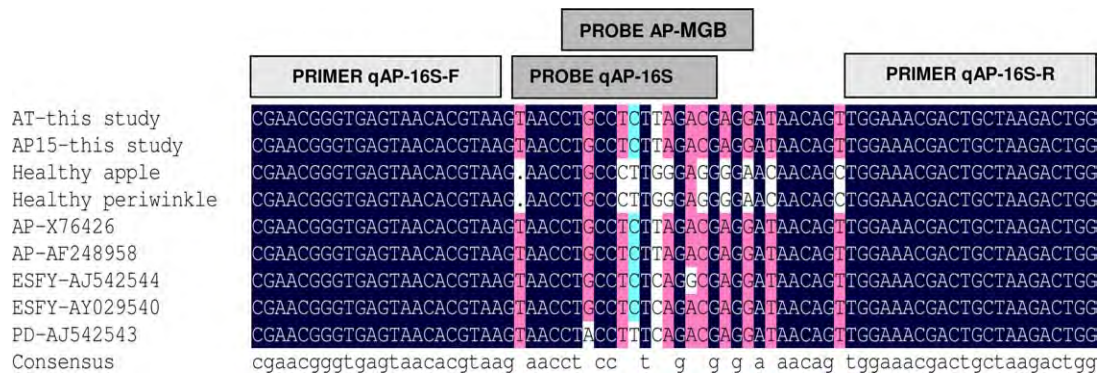


Figure 2 Alignment comparison of a sequenced region of 16S rRNA gene in two strains of ‘*Ca. P. mali*’ (AT and AP15) and two healthy samples (apple and periwinkle) with reference strains of ‘*Ca. P. mali*’, ‘*Ca. P. prunorum*’ (ESFY) and ‘*Ca. P. pyri*’ (PD). In sequence alignments, the dots mean absence of nucleotide.

species detailed in Table 1, whatever the methodology used to select the threshold (Fig. 1).

Limits of detection of conventional and real-time PCR protocols

Our new real-time PCR assay successfully detected a single copy of plasmid in water (Table 3). When the plasmid was diluted in DNA extract from healthy plants, 100 plasmid copies were still detected reproducibly.

We then compared the limits of detection of the five PCR protocols. The classical PCR protocol successfully detected the phytoplasma down to dilution 10⁻⁵ of samples A and B and down to dilution 10⁻⁴ of sample C (data not shown). Both real-time PCR protocols using SYBR Green (Jarausch *et al.*, 2004; Torres *et al.*, 2005) displayed a reproducible limit of detection of 10⁻⁵ whatever the sample. Both real-time PCR protocols using TaqMan MGB probes successfully detected the target in each sample and each PCR run down to dilution 10⁻⁶ (data not shown).

Table 3 C_t values of serial dilution of standard (plasmid) prepared in water or DNA extract of healthy plants (pure or 100-fold diluted in water)

Standard Copy Number in PCR Reaction ^a	Standard Prepared in Water	Standard Prepared in DNA Extract of Healthy Apples
One copy	40.37	ND
10 copies	36.18	ND
10 ² copies	33.31	44.72
10 ³ copies	30.53	34.42
10 ⁴ copies	27.77	28.66
10 ⁵ copies	24.25	24.92

ND, not-detected.

^aThere was no significant difference between C_t values of different treatment of standard from 10⁵ to 10⁸ copies (LSD test available as SAS software, P < 0.05).

Detection of phytoplasma after grafting by biological indexing and PCR

The appearance of symptoms was periodically surveyed (Table 4). Three months after grafting of infected materials on periwinkles, the symptoms of phytoplasma disease (yellowing and reduced leaf size) were conspicuous on 71% of the plants. Only after 4 months, did all the plants display proliferation symptoms. On grafted MM106 rootstocks, 10% of the plants showed symptoms after 4 months. After 7 months, 25% of the plants still showed no symptoms.

Phytoplasma infection was also evaluated by conventional PCR using the universal primer pair fU5/rU3 (Table 4). It was detected in all periwinkle and apple plants after 4 and 7 months, respectively. Symptom indexing and conventional PCR detected 100% of the infected periwinkles at the same moment (at 4 months), but a higher percentage of infected plants was detected by conventional PCR than by symptom indexing during the second and third months for periwinkle and from the second to seventh month for apple. Conventional PCR thus has an important advantage: the ability to detect *Ca. P. mali* in all infected apple trees within 7 months.

The third method used to detect phytoplasma was our new real-time PCR protocol (Table 4). It proved even more advantageous than the conventional PCR protocol, detecting *Ca. P. mali* infection in all grafted plants, both periwinkle and apple, only 3 months after grafting, that is earlier than either conventional PCR or biological indexing.

Discussion

In this study, we have designed a new MGB probe and new PCR conditions to improve the specificity of *Ca. P.*

Table 4 Results of indexing and detection of phytoplasma by different PCR methods in different periods after grafting the infected samples on healthy plants in greenhouse

Period	Symptom Indexing		Conventional PCR ^a		Real-time PCR ^b	
	Periwinkle (%)	Apple (%)	Periwinkle (%)	Apple (%)	Periwinkle (%)	Apple (%)
First month (15 days)	0 ^d	0	NT	0	NT	5
Second month (40 days)	26	0	50	35	71	70
Third month (70 days) ^c	71	0	96	80	100	100
Fourth month ^c	100	10	100	90	100	100
Seventh month ^c	100	75	100	100	100	100

NT, not-tested.

^aAmplification using universal primers fU5/rU3 (Lorenz *et al.*, 1995).

^bOur real-time PCR protocol by AP-minor groove binding probe.

^cDetection of phytoplasma in one leaf midrib.

^dPercentage of positive samples versus total number of tested plants (24 periwinkles and 20 apples).

mali detection. Our new real-time PCR protocol detects in water a single copy of DNA target per reaction and is ten times more sensitive than the conventional PCR protocol and two SYBR Green real-time PCR protocols.

We first tested the qAP-16S probe designed by Baric & Dalla-Via (2004). In our hands, this probe was not specific for *Ca. P. mali* detection because *Ca. P. prunorum* isolates were also detected. This result may be because of two factors: (a) the *Ca. P. prunorum* isolates used in our study only have a single mismatch with the sequence of the qAP-16S probe, whereas those of Baric & Dalla-Via (2004) had two mismatches and (b) the threshold line was set by inflexion point in the previous publication (Baric & Dalla-Via, 2004).

The position of a mismatch is crucial to the specificity of an MGB probe. MGB probes have been successfully employed to detect single-nucleotide mutations in the central region (Marbot *et al.*, 2002; Salmon *et al.*, 2002b) or the 3' terminal region (Kutyavin *et al.*, 2000; Salmon *et al.*, 2002a; Massart *et al.*, 2005) of the probe. Yao *et al.* (2006) observed that two mismatches in the centre of the probe-annealing region generated a signal in their MGB probe assay. Here, a single mismatch within the annealing region of the qAP-16S probe also generated a fluorescence signal. In both studies, similar fluorescence curves were obtained, with low fluorescence increases and low final delta Rn values. This low level of fluorescence for *Ca. P. prunorum* isolates was below the threshold line set by the inflexion point method but not below the line set by the 10 × SD method. Besides, an MGB probe may require careful optimisation to allow specific detection (Yao *et al.*, 2006). Under our optimised PCR conditions, the newly designed TaqMan MGB probe successfully detected all three described subtypes (Jarausch *et al.*, 2000) of *Ca. P. mali* (AP, AT1 and AT2) of different geographical origins, while no signal was observed with strains belonging to closely related phyto-

plasmas. This new TaqMan MGB probe thus showed higher specificity. Under these PCR conditions, the single-nucleotide polymorphism was detected more readily when located in the central region of the probe (position 9 out of 17) than in the 3' region (position 13 out of 18).

This new *Ca. P. mali*-specific probe can be helpful in ascertaining the presence of *Ca. P. mali* specifically. The finding that a single host plant can be doubly or multiply infected by different phytoplasmas (Bianco *et al.*, 1993; Lee *et al.*, 1994, 1995) suggests that the aetiologies of diseases associated with perennial fruit crops may be particularly complex. Fruit trees might harbour other phytoplasmas, and as several apple-growing areas worldwide have not yet been surveyed, we cannot just assume that *Ca. P. mali* is the only phytoplasma affecting apple.

The new protocol and that of Baric and Dalla-Via (2004) display similar sensitivity. Both are 10–100 times more sensitive than classical PCR protocol and 10 times more sensitive than two other real-time PCR protocols. This difference between real-time PCR protocols in sensitivity may be because of fluorescence chemistry, primer efficiency or the real-time PCR cycler. Clearly, this difference should be confirmed and better investigated through, for example, an interlaboratory comparison of these methods.

In our grafting experiments, the new real-time PCR protocol detected 100% of the infected material earlier than classical PCR or symptom observation. This confirms the higher sensitivity of real-time PCR and should be useful in the framework of a certification programme, where early detection of the infected status of a suspicious sample after grafting is of the essence. Importantly, in the case of a quarantine organism, high sensitivity should also make it possible to improve disease control through earlier detection *Ca. P. mali* in an infected tree. The earlier eradication measures can be taken, the lower the risk of contaminating other trees in the orchard. As mentioned

previously (Mumford *et al.*, 2006), real-time PCR is a reliable, quick and simple molecular method, well suited for high-throughput analysis of numerous samples.

In summary, we have designed a new real-time PCR protocol with improved specificity and high sensitivity to detect *Ca. P. mali*. This technique has great potential as a routine diagnostic tool in a procedure of plant nursery certification.

Acknowledgements

We are grateful to A. Bertaccini for critical reading of the manuscript. We thank A. Bertaccini, E. Seemüller, B. Schneider, B. Pradier, S. Baric, G. Firrao and R. Fialova for providing phytoplasma strains. J. Kummert, C. Bragard, R. Kettmann, O. Dutrecq and S. Steyer offered useful advice and suggestions, which we appreciate sincerely.

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