Rapid PCR-based Detection of Phytoplasmas from Infected Plants

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Abstract. Five simplified DNA preparation procedures for polymerase chain reaction (PCR) amplification were tested for detection of phytoplasmas from infected herbaceous and woody plants. Thin freehand cross-sections made from infected plant tissues and stored in acetone were used as sources for DNA preparation. The tissue sections were treated by: 1) grinding in sodium hydroxide; 2) sonication in water; 3) microwaving in water; 4) boiling in sodium hydroxide; or 5) placing directly in PCR tube. PCR amplification was performed with a universal phytoplasma-specific primer pair in a reaction buffer containing 0.5% (v/v) Triton X-100, 1.5 mM magnesium chloride, and 10 mM Tris-HCl. All five procedures provided phytoplasma template DNA for successful PCR amplification from infected herbaceous plants (periwinkle [Catharanthus roseus], green ash (Fraxinus pennsylvanica)), carrot (Daucus carota L.), maize (Zea mays L.), while the grinding, microwaving, and boiling procedures also allowed positive amplification from a woody plant [green ash (Fraxinus pennsylvanica Marsh.)]. The quality of the resulting DNA was adequate for subsequent identification of the aster yellows and ash yellow phytoplasmas through nested-PCR using phytoplasma group-specific primer pairs. These methods provide remarkable savings in labor and materials, making disease testing and indexing of plant materials much more attractive.

Phytoplasmas [formerly known as mycoplasma-like organisms (MLOs)] are a group of cell wall-less, nonculturable prokaryotes. These insect-transmitted, phloem-restricted microorganisms are associated with diseases in more than 300 species of higher plants worldwide, from cool temperate to tropical regions (Kirkpatrick, 1989). Conventional diagnostic procedures for detection of phytoplasma infection requires pre-development of an antibody or DNA probe specific to phytoplasmas and is only moderately sensitive. If the diagnosis is negative, it is unknown whether it is due to a low titer of phytoplasmas in the plant tissue or due to a lack of infection. Polymerase chain reaction (PCR) amplification is a very sensitive diagnostic method, but the normal DNA extraction processes are tedious, especially when handling a large number of samples. Traditional DNA preparation from one plant sample takes 8–10 h, with an added 20 min for each additional sample, up to 10 samples. Beyond 10 samples, another set must be run on a different day. Although many protocols for phytoplasma DNA extraction have been developed for PCR amplification (Gibb and Padovan, 1994; Green et al., 1999; Lee et al., 1993b; Levy et al., 1994), all require multiple steps or commercial kits. Several researchers have reported simplified DNA preparation procedures for detection or diagnosis of other organisms (Liu et al., 1995; Wang et al., 1993; Zhang and Goodwin, 1997). We report here adaptations of these simplified DNA preparation methods for PCR-based detection and diagnosis of phytoplasmas. They can significantly speed up the overall PCR process and reduce chemical and labor costs.

Materials and Methods

The plant materials, the associated phytoplasmas, and their sources for this research are listed in Table 1. Various methods were used to confirm the existence of the respective phytoplasmas (Table 1). Infected periwinkle plants containing various phytoplasmal strains were kept in the greenhouse and the phytoplasmas were maintained in them by grafting a branch from the diseased plant to a healthy one. Samples of green ash, carrot, and maize were from the various sources. Both phytoplasma-infected and noninfected plant samples were used. Very thin freehand cross-sections of leaf midribs, young shoots, or roots from the various plant materials were fixed in acetone for subsequent DNA extraction (Guo et al., 2000). Five methods were used to prepare the phytoplasma template DNA for testing.

For method 1, 10–20 fixed sections were ground in 20 µL of 0.3 N NaOH with a tissue grinder (Kontes, NJ) in a 1.5-mL microtube. After clarification by full-speed centrifugation (Eppendorf centrifuge, model 5415D) for 30s, 2 µL of extract was mixed with 48 µL of 100 mM Tris (pH 8.0). For the sonication method, 5–10 fixed sections were sonicated in 10 µL water for 1 min (model #2210, Bransonic Ultrasound Cleaner; Branson, Danbury, Conn.). For the microwaving method, 5–10 fixed sections were microwaved in 10 µL water for 5 min (model #564.8844881; Kenmore Sensor Cook Microwave Oven; Sears, Roebuck & Co., Mountain View, CA). For all the procedures for PCR amplification, 0.5 µL PCR product was electrophoresed on a 1.5% agarose gel.

| Plant     | Associated phytoplasmas | Source 1 | Confirmation methods 1  |
|-----------|--------------------------|----------|------------------------|
| Periwinkle | [Catharanthus roseus (L.) G. Don] | T.A. Chen | Symptoms, +IF          |
|           | Eastern X-disease phytoplasmas | T.A. Chen | Symptoms, +IF          |
|           | Aster yellows phytoplasmas | W. Sinclair | Symptoms, +IF          |
|           | Ash yellows phytoplasmas | W. Sinclair | Symptoms, +IF          |
|           | Elm yellows phytoplasmas | J.A. Walla | No symptoms, –IF       |
|           | Noninfected plants |          |                        |
| Green ash  | (Fraxinus pennsylvania Marsh.) |          |                        |
|           | Ash yellows phytoplasmas | Walla et al., 2000 | Symptoms, +IF          |
|           | Noninfected plants | J.A. Walla | No symptoms, –IF       |
| Carrot     | (Daucus carota L.) |          |                        |
|           | Aster yellows phytoplasmas | C.W. Lee | Symptoms, +IF          |
|           | Noninfected | C.W. Lee | No symptoms, –IF       |
| Maize      | (Zea mays L.) |          |                        |
|           | Aster yellows phytoplasmas | T.A. Chen | Symptoms, +IF          |
|           | Noninfected | T.A. Chen | No symptoms, –IF       |

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and Co., Chicago). For the boiling method, 5–10 fixed sections were boiled in 10 µL 0.3 N NaOH for 5 min and then neutralized with 10 µL 0.3 N HCl. For the direct tissue method, 3–5 fixed sections were placed directly in the PCR reaction tube for amplification. A 1-µL aliquot of solution from methods 1–4 was used in each PCR reaction. DNA prepared by a more conventional method (Lee et al., 1993a) was used as the control. A universal phytoplasma-specific primer pair (forward primer P16S3F: CGGGTTTGTACACACCGCCCTCA and reverse primer P235R:TCTTAGTGC-CAAGGCATCCACTGTC), which amplifies the 16S/23S ribosomal RNA spacer region, (Guo et al., 2000) was used in PCR amplification. The program for PCR amplification was denaturing for 1 min at 94 °C, annealing for 2 min at 60 °C and extension for 3 min at 72 °C, followed by a 10 min final extension at 72 °C. The PCR reaction was performed in a final volume of 25 µL containing 0.5% (v/v) Triton X-100, 1.5 mM MgCl₂, 10 mM Tris-HCl (pH 8.0), 200 µM each of dNTPs, 0.4 µmol primers, and 0.625 units of Taq DNA polymerase (Promega Corp., Madison, Wis.) for 35 cycles in a thermocycler (RoboCycler Gradient 96, Stratagene, La Jolla, Calif.).

For the periwinkle plants infected by aster yellows phytoplasmas and the green ash plants infected by ash yellows phytoplasmas and their respective control plants, the DNA prepared by these five methods was also used in a standard two-step nested-PCR (Lee et al., 1993b). In the nested-PCR, the universal phytoplasma-specific primer pair was used first to determine whether the plant was infected with any phytoplasma, and then a second, phytoplasmal group-specific primer pair was used to determine the specific phytoplasmal group.

All amplified products (10 µL each) were electrophoresed in a 1% agarose gel and visualized by UV light transillumination after staining with ethidium bromide.

Results and Discussion

All five methods provided phytoplasma DNA for successful PCR amplification from periwinkles infected with X-disease (shown in Fig. 1), aster yellows, ash yellows, and elm yellows phytoplasmas, carrot with aster yellows phytoplasmas, and maize with maize bushy stunt phytoplasmas, but not from any noninfected controls. However, the amount of amplified DNA was substantially lower using the direct tissue method. The grinding, microwaving and boiling, but not the sonication and direct tissue methods, provided phytoplasmal DNA for successful PCR amplification from green ash infected with ash yellows phytoplasmas (Fig. 2). The microwaving method resulted in a substantially weaker band than the grinding and boiling methods.

Results using nested-PCR consistently showed positive amplifications of DNA from infected, but not from noninfected, periwinkle and green ash plant materials (data not shown). Thus, the prepared DNA can be successfully used with various primer pairs to detect and diagnose various phytoplasmas.
Compared to published protocols of conventional preparation of phytoplasma DNA, the methods described here are much simpler. These processes apparently disrupt phytoplasma cells in the plant tissue sections and release phytoplasma DNA to the solution in less than 10 min. The extracted/exposed DNA can be used directly as the template for the single-step PCR or two-step nested-PCR amplification. Because all five methods allowed positive amplifications of phytoplasma DNA from periwinkles, we recommend the direct tissue method for diagnosis of phytoplasmas in herbaceous plants because it is the simplest method. Among the methods that worked with the woody plant samples, we prefer the boiling method for phytoplasmal diagnosis with woody plants because grinding woody plant sections, which is difficult, and the microwaving method resulted in weaker amplification. These methods significantly simplify the laborious process of DNA preparation for PCR amplification, yet maintain the high sensitivity of traditional PCR methods, and therefore, can dramatically reduce the cost of the PCR-based diagnosis of phytoplasma diseases.

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