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Host range of *Phytophthora parsiana*: a new high temperature pathogen of woody plants

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Summary. Among several *Phytophthora* spp. reported previously from *Pistacia vera* in Iran, a high temperature species recently identified as *P. parsiana* (formerly known as high temperature *P. cryptogea*) is becoming important in woody plants, including *P. vera*. The host range of this newly recognised species, including both annual and perennial plants, is reported here. The pathogen infected 4–5 month-old glasshouse grown seedlings of *P. vera*, *Ficus carica*, *Malus pumila* and *Prunus dulcis*, and detached stems of 23 woody plants collected during dormant and growing seasons. Nineteen field and vegetable crops and 17 weed species were not infected by *P. parsiana* in these pathogenicity assays.

Key words: Pistacia vera, Iran, Phytophthora cryptogea, gummosis, crown rot.

Introduction

Phytophthora species are major soil-borne plant pathogens in various agricultural commodity crops in Iran, causing high losses in annual and perennial plant species. Major economic losses are experienced in *Pistacia vera* (pistachio) under saline and non-saline conditions (Banihashemi, 1995).

Several species of *Phytophthora* have been reported on pistachio in Iran, including *P. citrophthora* (Banihashemi, 1983; Mirabolfathy *et al.*,1989), *P. megasperma* (Mirabolfathy and Ershad, 1987). *P. drechsleri* (Aminaee and Ershad, 1991; Banihashemi, 1995), *P. cryptogea* and *P. nicotianae* (Banihashemi, 1995). In a re-examination of *P. megasperma* and *P. drechsleri* from pistachio in Iran using RFLP and ITS sequence analyses, Mirabolfathy *et al.* (2001) showed that the *P. megasperma* like isolates from pistachio were a newly recognised species, *P. pistaciae*; while *P. drechsleri* from pistachio was considered to be *P. melonis*.

The three Phytophthora species P. melonis, P. drechsleri and P. cryptogea are morphologically identical and cannot be separated by conventional methods. Ho and Jong (1986) indicated both P. drechsleri and P. cryptogea to be identical and indicated that growth at 35° C, or this in combination with other criteria, could not separate these species. The two species were, therefore, merged into P. cryptogea, which has priority. Later (Ho and Jong ,1991) growth at 35° C was used to separate the two species.

Controversy remains concerning the morphological identity of *P. melonis*, *P.drechsleri* and *P. cryptogea*. Based on molecular analysis, some isolates of *P. drechsleri* from pistachio from Iran initially identified by morphology, were re-identified as *P. melonis* (Mirabolfathy *et al.*, 2001). The high temperature isolates of *P. cryptogea* reported from fig (Banihashemi and Ghaisi, 1993) and pistachio (MacDonald *et al.*, 1992; Banihashemi, 1995) could not be separated morphologically from common isolates of the pathogen. Mostowfizadeh-Ghalamfarsa (2005), examining the phylogeny of the taxonomically challenging species *P. drechsleri* and *P.cryptogea* from various sources, found a monophyletic group of isolates distinct from either spe-

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cies on the basis of rDNA ITS sequence, which was later reported as *Phytophthora parsiana* sp. nov. a new high temperature tolerant species (Mostowfizadeh-Ghalamfarsa *et al.*, 2008). All isolates previously identified morphologically as high temperature *P. cryptogea* isolates had been obtained from woody plants such as *Ficus carica* (Banihashemi and Ghaisi ,1993), and *P. vera* (MacDonald *et al.*,1992; Banihashemi, 1995).

The host range of *P. cryptogea* is mostly reported to be herbaceous plants, on which the pathogen causes root rot and seedling blight, but a few woody plants were included as hosts (Erwin and Ribeiro,1996).

The objective of the present study was to examine the host range of *P. parsiana* (formerly known as high temperature tolerant *P. cryptogea*) under glasshouse conditions. A summary of this study was reported earlier (Hajebrahimi and Banihashemi, 2008).

Materials and methods

Sources of isolates and host-range studies

Three representative isolates of P. parsiana from different woody hosts and locations were used in this study (Table 1).

Table 1. Sources of isolates of *Phytophthora parsiana*.

Woody plants

In a glasshouse test, six woody plant species were tested for susceptibility to *P. parsiana* (Table 2). Seeds of apple (*Malus pumila*), almond (*Prunus dulcis*) pistachio (*Pistacia vera*), walnut (*Juglans regia*) and lemon (*Citrus limon*) were surface sterilized for 5 min in 0.5% NaOCl and planted in 20 cm diameter pots containing steam sterilized soil:sand (2:1 v/v) and raised in a glasshouse at $16-35^{\circ}$ C. Stem cuttings of fig were also planted in the same substrate. Three replicate pots each containing three plant species were used in each treatment.

Herbaceous plants

Seeds of 18 species of field and vegetable crops (Table 3) were surface sterilized as described above, and planted in 9 cm diameter pots containing the same soil mix and raised in a growth chamber at $22-24^{\circ}$ C with 16:8 light-dark cycle. Three replicate pots each with five seedlings were used.

Weeds

Seeds of 16 common weed species (Table 3) were surface sterilized as described above, and planted in steam-sterilized soil mix and kept in

Isolate code	Host	Location	Year isolated	Source
SUC 7	Pistacia vera	USA	1992	Z. Banihashemi
SUC 19	Pistacia vera	Iran	1992	Z. Banihashemi
SUC 25	Ficus carica	Iran	1991	Z. Banihashemi

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Table 2	Pathogenicity	of Phytophthora	narsiana	isolates in	i some snec	ies of woody high	nts
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Scientific name	Common name —]	Pathogenicity of isolates	3 ^a
		SUC7	SUC25	SUC19
Ficus carica	Fig	+	+	+
Pistacia vera	Pistachio	+	-	+
Malus pumila	Apple	+	+	+
Prunus dulcis	Almond	+	+	+
Juglans regia	Walnut	-	-	-
Citrus lemon	Lemon	-	-	-

^a *P. parsiana* re-isolated; - *P. parsiana* not re-isolated.

glasshouse (16–35 $^{\circ}\mathrm{C}).$ Three replicate pots each with five seedlings were used.

Detached stems of woody plants

Stems $(20 \times 1 \text{ cm})$ of 21 woody plant species were collected during dormant and active growth stages (Table 4). Six replicate cuttings were inoculated for each species

Fruit, tuber and root crop plants

Fresh fruits, tuber and roots of various crops were obtained from local markets and fields and used for inoculation. Eight replicates were used for each treatment (Table 5).

Inoculum production

Inocula of the *P. parsiana* isolates were produced on vermiculite amended with hemp seed extract (Banihashemi and Fatehi 1989). Two hundred ml of vermiculite and 120 mL hemp seed extract (extract of 60 g hemp seed ⁻¹L distilled water) were autoclaved for 20 min and inoculated with three to four 6mm blocks of fresh culture grown on corn meal agar and incubated at room temperature for 4–6 weeks.

Plant	Scientific name	Common name		
Herbaceous	Cucumis sativus	Cucumber		
	C.melo var. flexuosus	Snake melon		
	C.melo	Melon		
	Cucurbita maxima	Pumpkin		
	Citrullus lanatus	Watermelon		
	Vicia sativa	Mungbean		
	Vigna unguiculata	Cowpea		
	Soja hispida	Soja		
	Phaseolus coccineus	Scarlet runner bean		
	Lens esculenta	Lentil		
	Cicer arietinum	Chick pea		
	Lycopersicum esculentum	Tomato		
	Solanum melongena	Egg plant		
	Helianthus annuus	Sunflower		
	Daucus carota	Carrot		
	Sesamum indicum	Sesame		
	Brassica napus	Rape seed		
	Spinacia oleracea	Spinach		
Weeds	Prangos uloptera	'Djashir'		
	Echinochloa sp.	Panic grass		
	Avena sativa	Oat		
	Triticum polonicum	Polish wheat		
	Hordeum spontaneum	Spontaneum barley		
	Cardaria draba	Cress		
	Chenopodium album	Pig weed		
	Rumex crispus	Curled dock		
	Solanum dulcamara	Night shade		
	Malva sylvestris	mallow		
	Launaea sp.	Launaea		
	Amaranthus sp.	Amaranth		
	Plantago major	Way bread		
	Portulaca oleraceae	Purslane		
	Melitotus alba	Sweet clover		
	Glycyrhiza glabra	Liquorice		

Table 3. Herbaceous plants and weeds inoculated with *Phytophthora parsiana* under glasshouse conditions.

Pathogenicity test

Plant inoculation

Ten to 50 ml of inoculum, depending on the size of the pot, was used in each pot containing plants. Inoculum was spread over the soil surface and, following closure of the drainage holes using melted paraffin wax, the pots were flooded with water over night with incubation at the indicated temperatures. The formation and release of zoospores were monitored for 4 weeks, as reported previously (Banihashemi, 2004). After overnight flooding, the drainage hole in each pot was re-opened by removing the paraffin wax plug. The drained water was collected separately from each pot and filtered through a double layer of cheese cloth. Fifty to 100, 6 mm citrus leaf disks were added to each collected sample and incubated at room temperature for 48 h. Subsequently, bait discs were washed gently under running tap water,

blotted dry and plated on PARP medium (Jeffers and Martin, 1986). Flooding was repeated every second week. Number of baits colonized by *P. parsiana* was recorded at each time point and percent colonization of baits counted to ensure the presence of the active pathogen in the pots.

Detached stem inoculation

Stems (20×1.5 cm) were washed, blotted dry and the surface was wiped with cotton impregnated with 95% ethanol. The two end cuts of each stem were dipped in warm melted paraffin wax to reduce desiccation during incubation. Three T-shaped cuts (2–3 cm) were made along each stem. A 6 mm corn meal agar (CMA) plug from actively growing hyphae of each isolate was inserted into each cut, the bark replaced, and the wound and inoculum wrapped with Parafilm. Stems were incubated at room tempera-

Table 4. Pathogenicity of isolates of Phytophthora parsiana on detached stems of different woody plants..

			Pathogenicity of isolates					
Scientific name	Common name	SUC19		SUC17		SUC25		
Scientific name		Winter	Growing season	Winter	Growing season	Winter	Growing season	
Morus alba	White mulberry	+	-	+	-	+	-	
Ulmus campestris	Elm	+	-	-	-	-	-	
Cupressus sempervirens	Cypress	-	-	-	-	-	-	
Magnolia grandiflora	Magnolia	-	-	-	-	-	-	
Ficus carica	Fig	+	-	+	-	+	-	
Eucalyptus globus	Eucalyptus	-	-	-	-	-	-	
Pistacia vera	Pistachio	+	+	+	+	+	+	
Fraxinus rotundifolia	Ash	-	-	-	-	+	-	
Pinus eldarica	Pine	-	-	-	-	-	-	
Platanus orientalis	Sycamore	+	-	-	-	-	-	
Acer monspessulanum	Maple	+	+	+	+	+	+	
Juglans regia	Walnut	+	-	+	-	+	-	
Ailanthus altissima	Ailanthus	+	-	+	-	+	-	
Robinia pseudoacacia	Acacia	-	-	-	-	+	-	
Citrus aurantium	Sour orange	-	-	-	-	-	-	
Punica granatum	Pomegranate	+	-	+	-	-	-	
Malus pumila	Apple	+	+	+	+	+	+	
Prunus dulcis	Almond	+	-	+	-	+	-	
Cydonia oblonga	Quince	+	-	+	-	+	-	
Persica vulgaris	Peach	-	-	-	-	-	-	
Rosa canina	Dog rose	+	-	+	-	+	-	

Scientific name	Common nome	Pathogenicity of isolates			
Scientific name	Common name	SUC25	SUC7	SUC19	
Malus pumila	Apple (cv. Golden delicious)	20	15	15	
Malus pumila	Apple (cv. Red delicious)	15	10	10	
Malus pumila	Apple (cv. Golden delicious, unripe)	10	10	10	
Citrus nobilis	Mandarine	40	40	40	
Citrus lemon	Lemon	40	40	40	
Citrus sinensis	Orange	40	40	40	
Citrus aurantium	Sour orange	40	40	40	
Musa paradisiaca	Banana	60	40	60	
Ficus carica	Fig (ripe)	40	60	50	
	Fig (unripe)	30	20	40	
Cucumis sativus	Cucumber	90	90	90	
Lycopersicon esculentum	Tomato	100	100	100	
Solanum melongena	Egg plant	20	10	50	
Daucus carota	Carrot	0	0	0	
Beta vulgaris	Sugar beet	0	0	0	
Solanum tuberosum	Potato	0	0	0	

Table 5. Percentage infection of various fruits, roots and tubers of different plants by *Phytophthora parsiana* isolates 5 days after inoculation.

ture and observed every other week for symptom development. Experimental controls comprised of stems inoculated with sterile CMA. room temperature. Colonies emerging from inoculated tissues were identified morphologically.

Fruit, tuber and root crop plants

Fruits, tubers and roots were thoroughly washed, left to dry at room temperature and wiped with 95% ethanol. A cork borer was used to make two 6 mm holes on opposing sides of each of these, and an equal diameter plug of an actively growing CMA culture of each isolate was inserted in the hole. The block of plant tissue was replaced in the wound. Wounds were covered with adhesive tape and the fruits, roots and tubers were incubated in plastic bags at room temperature. Sterile CMA plugs were used as negative experimental controls.

Re-isolation of pathogens

Roots, tubers, crowns of the seedlings, stems and fruits showing any discoloration or decay were transferred to PARP medium and incubated at

Results

Woody plants

Walnut and lime were not infected by *P. parsi*ana and the pathogen was not recovered from inoculated plants. Pistachio, fig, apple and almond were infected and the pathogen was re-isolated from infected tissues (Table 2). The first disease symptoms in pistachio appeared 5 months after inoculation. Infected plants showed mild wilting with root and crown necrosis. Apple seedlings showed disease symptoms 4 months after inoculation, including defoliation and extending necrosis in the lower stems. In almond, severe root rot and shoot necrosis occurred 4 months after inoculation. In fig, severe root rot and lower stem necrosis were observed 5 months after inoculation. The pathogen was re-isolated from infected tissues.

Herbaceous plants and weeds

Of the 18 herbaceous plants and 16 weeds (Table 3) inoculated with P. parsiana isolates, none were infected by the pathogen. No pathogen was re-isolated from inoculated plants or plant organs.

Detached stems of woody plants

Only stem samples of pistachio, *Malus pumila* and maple collected during active plant growth showed infection by *P. parsiana*. Stem samples of all 23 woody plants collected during dormant stages were infected by *P. parsiana* and the pathogen was re-isolated from infected tissues (Table 4).

Fruit, tuber and root crop plants

Phytophthora parsiana caused severe rot on all fruits but none of the roots or tubers were infected by the pathogen (Table 5).

Discussion

The present study showed that the newly described high temperature tolerant *Phytophthora parsiana* might be a serious threat to pistachio and some important woody plants especially under high temperature conditions. From a limited number of woody plants examined, both fruit and nut crops which are grown under various climatic conditions were hosts susceptible to the pathogen. None of the herbaceous plant species examined, including vegetable and field crop plants and weeds, were immune to the pathogen. At present, it is premature to finally conclude that the pathogen specifically attacks perennial woody plants.

Very few species of *Phytophthora* which are limited to woody plants grow above 35° C. *Phytophthora melonis*, *P. drechsleri* and *P. nicotianae*, which are high temperature species, infect herbaceous and woody plant species. As a result of recent climate change and global warming in many parts of the world (Garret *et al.*, 2006) the new high temperature species may become serious threats to many woody plants.

Several *Phytophthora* species have been reported to cause pistachio gummosis resulting in crown and root rot. *Phytophthora citrophthora* was the most aggressive species isolated from pistachio growing areas of southern Iran (Banihashemi, 1984), due to the low tolerance of most local Iranian rootstocks used in pistachio plantations (Banihashemi, 1998).

Other *Phytophthora* species were also reported to cause pistachio gummosis in Rafsenian a major pistachio growing area in Iran. These include P. megasperma, a high temperature P. cryptogea (McDonald et al., 1992; Banihashemi, 1995), P. drechsleri (Aminaee and Ershad, 1991; Banihashemi, 1995) and P. nicotianae (Banihashemi, 1995). Mis-identification of isolates when using conventional morphological features resulted in the introduction of hitherto unrecognised species of *Phytophthora* associated with pistachio gummosis. Mirabolfathy et al. (2001), using RFLP and ITS analyses, reported that isolates of species previously identified as P. drechsleri were actually P. melonis, and that P. megasperma was actually the new species P. pistacia, which could only be differentiated by molecular analysis. No detailed morphological features could be used to separate these species. A high temperature P. cryptogea, reported by Banihashemi (1995) as the causal agent of pistachio gummosis in Iran was also isolated from fig in southern Iran (Banihashemi and Ghaisi, 1992) and pistachio in California USA (MacDonald et al., 1992).

Based on morphological criteria, the taxonomic position of these isolates was not defined. Mostowfizadeh-Ghalamfarsa et al. (2008), using analysis of ITS sequences of various isolates of P. drechsleri and *P.cryptogea* (including high temperature isolates) found that, although the isolates were morphologically identical, the species represented distinct monophyletic groups. High temperature P. cryptogea from various hosts and geographical regions were distinct from P. drechsleri and P. cryptogea (low-maximum temperature) and all members of Phytophthora ITS clades 1-8. This lead to the proposal for the new species P. parsiana (Mostowfizadeh-Ghalamfarsa et al., 2008), which had been isolated only from woody hosts. Information on the host range of this new species is very limited.

The host range of *P. cryptogea* isolates identified on the basis of morphological criteria mostly included herbaceous plant species. Very few woody plants were considered hosts (Erwin and Ribeiro, 1996). Re-examination of the identity of the species and host range is required to determine if different species were involved.

The present study explored the host range of *P. parsiana* for disease management. None of the herbaceous plants (field and vegetable crops, weeds) were hosts for the pathogen under high inoculum po-

tential. Although maximum temperature for growth is the present criterion for separating *P. cryptogea* from *P. parsiana*, morphological or physiological features other than molecular analysis should be available for identification. Some isolates of *P. parsiana* caused mild infections in safflower (Hajebrahimi, 2008), although *P. cryptogea* is highly pathogenic to safflower cultivars (Mirtalebi and Banihashemi, 2006; Banihashemi and Mirtalebi, 2007).

Recently, several isolates of *P. parsiana* with high maximum growth temperatures have been recovered from old pistachio trees in Kerman Province (A.H. Mohammadi, personal communication); identity of the isolates was confirmed by molecular analysis (C.X. Hon g, personal communication). The presence of the pathogen from various parts of the world requires confirmation by stringent identification techniques. Distribution of the pathogen, especially under high temperature climatic conditions, also requires investigation.

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