Re-evaluation of *Phytophthora* **Species Isolated During 30 Years of Vegetation** Health Surveys in Western Australia Using Molecular Techniques

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ABSTRACT

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For 30 years, large-scale aerial photography has been used to map the extent of Phytophthora dieback disease in native forests in the southwest of Western Australia, with validation of the observations involving routine testing of soil and root samples for the presence of Phytophthora cinnamomi. In addition to P. cinnamomi, six morpho-species have been identified using this technique: P. citricola, P. megasperma, P. cryptogea, P. drechsleri, P. nicotianae, and P. boehmeriae. In recent years, many new Phytophthora species have been described worldwide, often with similar morphology to existing species; thus, as many of the isolates collected in Western Australia have been difficult to identify based on morphology, molecular identification of the morpho-species is required. Based on amplification of the internal transcribed spacer (ITS) region of the rDNA gene, sequence data of more than 230 isolates were compared with those of existing species and undescribed taxa. P. inundata, P. asparagi, P. taxon PgChlamydo, P. taxon personii, and P. taxon niederhauserii were identified based on sequence data. Phylogenetic analysis revealed that nine potentially new and undescribed taxa can be distinguished. Several of the new taxa are morphologically indistinguishable from species such as P. citricola, P. drechsleri, and P. megasperma. In some cases, the new taxa are closely related to species with similar morphology (e.g., P.sp.4 and P. citricola). However, the DNA sequences of other new taxa such as P.sp.3 and P.sp.9 show that they are not closely related to morphologically similar species P. drechsleri and P. megasperma, respectively. Most of the new taxa have been associated with dying Banksia spp., while P.sp.2 and P.sp.4 have also been isolated from dying Eucalyptus marginata (jarrah). Some taxa (P.sp.3, 6, and 7) appear to have limited distribution, while others like P.sp.4 are widespread.

The southwest of Western Australia (WA) supports one of the richest regions of floristic biodiversity in the world, with more than 5,700 described native plant species. Eucalyptus marginata (jarrah) is the dominant overstory tree species in much of the forest in this region. The introduced root and collar pathogen Phytophthora cinnamomi has had a major impact on the jarrah forest ecosystem (31,32,43). Shearer et al. (30) estimated that 2,284 of the plant species are susceptible to the pathogen, while many other species are affected indirectly through the loss of the overstory. P. cinnamomi reduces the diversity of plant species on infested sites. The altered vegetation structure in

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doi:10.1094/PDIS-93-3-0215 © 2009 The American Phytopathological Society turn affects the biodiversity of fauna (17). In WA, there has been, and continues to be, active management of Phytophthora dieback spread through extensive mapping of affected areas and then quarantining of dieback free areas (32). Infested areas of high conservation value are also treated with salts of phosphonic acid (active compound, the anion HPO_3^{2-}) to reduce the impact and rate of spread of the pathogen (18).

Extensive vegetation health survey in the WA jarrah forest has been in continuous operation since 1978 (12,34). The mapping of the extent of Phytophthora dieback disease is based on shadowless color aerial photography (34). Validation of mapping involves collecting soil and root samples from beneath dying, *Phytophthora*-sensitive native plants known as "indicator species" and the testing of them for the presence of *Phytophthora* spp. (34). Samples have also been collected from dying plants in native heathlands and other forest types, and from dying trees in *Pinus radiata* plantations. *Phytophthora* isolates

are obtained by standard baiting procedures (24) at the Vegetation Health Service (VHS) laboratory of the Department of Environment and Conservation, where isolates are identified from morphological characters where possible, and representatives are added to the VHS culture collection. P. cinnamomi is easily identified by its morphological characteristics. However, about 10 to 15% of baited samples return a positive result for Phytophthora species other than P. cinnamomi. Six other Phytophthora spp., P. citricola, P. megasperma, P. cryptogea, P. drechsleri, P. nicotianae, and P. boehmeriae, have been isolated and identified using morphological characters (10,35).

Cooke et al. (8) determined the phylogenetic relationships among 50 Phytophthora species based on internal transcribed spacer (ITS) sequence data, and identified 10 phylogenetic clades that did not correspond to previous morphological groupings (40,41). The contradiction between morphological characteristics and molecular phylogenetics indicates that the traditional morphological features used to classify Phytophthora spp. have evolved numerous times within the genus and that morphological classification alone does not result in natural assemblages. Since Cooke et al. (8), most descriptions of new Phytophthora spp. have included a molecular phylogeny where the new species is compared not only with those species with the most similar morphology, but also with those that are phylogenetically the closest (e.g., 4,6,22,26). Additional gene regions have been sequenced for most Phytophthora spp. (3,23,25). However, for an initial identification, sequencing the ITS region and conducting a BLAST search on GenBank coupled with phylogenetic analysis provides a valid identification. New taxa can also be identified in this manner, although further gene regions might be required to distinguish cryptic species.

In addition to the *Phytophthora* species identified by morphology, the VHS has acquired many isolates that could not be identified, or for which the morphological identification seemed to be misleading. Thus, the ITS gene region of more than 230 isolates, including both unidentified

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isolates and those previously identified by morphological characters, was sequenced to confirm isolate identity, and the results are reported here.

MATERIALS AND METHODS

Collection of isolates. Soil and root samples were collected from beneath dying, *Phytophthora*-sensitive indicator species in native ecosystems in the southwest of WA. Samples were baited with *Eucalyptus sieberi* cotyledons (24) that were plated after 5 days and 10 days onto NARPH (21), or P_{10} VPH (39) selective media, from which a pure culture of *Phytophthora* was

then isolated. In some cases, plant roots were surface-sterilized in 70% ethanol for 30 s, followed by four rinses in distilled water, and plated directly onto selective media. Cultures derived from earlier research and surveys have been incorporated into the present VHS collection, retaining the original abbreviations that designate their origin: MJS = M. J. C. Stukely; TCH = T. C. Hill; HSA = Hart, Simpson & Associates; DCE = E. M. Davison; and DDS = earlier prefix of the VHS collection.

Morphological identification. In general, morphological examination and identification were conducted when the isolates were initially collected. Isolates with IMI codes (Tables 1 and 2) were sent to the Imperial Mycological Institute, Kew, England for identification. Some isolates have been re-examined recently. In all cases, the same procedure is followed. Pure cultures of *Phytophthora* grown on 10% V8 juice agar were identified to species based on the morphology of their sporangia, oogonia, oospores, and related structures (15), or to the most likely Waterhouse Group (40) if all of the required structures were not formed. Sporangia were examined on squares cut from actively growing cultures 4 to 7 days old and suspended in nonsterile

Table 1. Identity (based on morphology and internal transcribed spacer [ITS] sequence data), host, location, and GenBank accession numbers for representative *Phytophthora* isolates from Western Australia, excluding isolates from Cooke's clade 6^a

Culture identification ^b	Morphology identification	Sequence identification	Host	Location	Cooke's clade ^c	GenBank accession no.
DDS3453	P. nicotianae (A2)	P. nicotianae	Chamaelaucium sp.	Perth (Marangaroo)	Clade 1b	EU301133
VHS16097	P. nicotianae (A1)	P. nicotianae	Xanthorrhoea platyphylla	Ravensthorpe	Clade 1b	EU301134
VHS16099	P. nicotianae (A1)	P. nicotianae	Dryandra cirsioides	Ravensthorpe	Clade 1b	EU301135
VHS13502	P. citricola	P.sp.2	Corymbia calophylla	Dwellingup	Clade 2	EU301119
VHS13523	P. citricola	P.sp.2	Eucalyptus marginata	Dwellingup	Clade 2	EU301120
VHS13559	P. citricola	P.sp.2	E. marginata	Dwellingup	Clade 2	EU301121
VHS13607	P. citricola	P.sp.2	E. marginata	Dwellingup	Clade 2	EU301122
VHS13615,	P. citricola	P.sp.2	E. marginata	Dwellingup	Clade 2	EF121961
WAC13038			-	N		
VHS13663	P. citricola	P.sp.2	E. marginata	Dwellingup	Clade 2	EU301123
VHS13694	P. citricola	P.sp.2	E. marginata	Dwellingup	Clade 2	EU301124
VHS13713, WAC13039	P. citricola	P.sp.2	E. marginata	Dwellingup	Clade 2	EF121962
VHS16103	P. citricola	P.sp.2	Banksia grandis	Pemberton	Clade 2	EU301125
DCE10, IMI133316	P. citricola	P.sp.4	(ex Agriculture Dept 1980)		Clade 2	EU301126
DDS1450, IMI329674	P. citricola	P.sp.4	Soil, native forest	Walpole	Clade 2	EU301127
DDS3458	P. citricola	P.sp.4	Patersonia sp.	Walpole	Clade 2	EU301128
DDS3571	P. citricola	P.sp.4	Conospermum sp.	Perth (Kings Park)	Clade 2	EU301129
HSA1210	P. citricola	P.sp.4	Gastrolobium spinosum	Coorow	Clade 2	EU301129
MJS278	P. citricola	P.sp.4	E. marginata	Jarrahdale	Clade 2	EU301130
VHS15445	P. citricola	P.sp.4	B. attenuata	Badgingarra	Clade 2	EU301131
VHS10154,	Waterhouse Group 1	1	B. littoralis	Bunbury	Clade 4	EU301112 EU301114
IMI389663	Waternouse Group I	Lisbit	D. morans	Dunbury	chude i	Lessin
VHS15453	Unidentified	P.sp.1	B. attenuata	Badgingarra	Clade 4	EU301115
VHS15485	Unidentified	P.sp.1	B. attenuata	Badgingarra	Clade 4	EU301116
VHS16282	P. citricola-like	P.sp.1	B. media	Ravensthorpe	Clade 4	EU301117
VHS9861,	Waterhouse Group 1	1	B. menziesii	Lancelin	Clade 4	EU301118
IMI389662	Waternouse Group I	1.501	Dimenziesti	Luncenn	chude i	Leouino
DDS1221	P. citricola-like	P.sp.1	Soil, native bush	Kalbarri	Clade 4	EU593265
VHS17576	Unidentified	P. taxon niederhauserii	B. prionotes	Lancelin	Clade 7b	EU301136
VHS17577	Unidentified	P. taxon niederhauserii		Lancelin	Clade 7b	EU301137
DDS3606	P. cryptogea (A2)	P. cryptogea	Daviesia sp.	Fitzgerald River NP	Clade 8	EU301138
DDS3612	P. cryptogea (A2)	P. cryptogea	D. cirsioides	Fitzgerald River NP	Clade 8	EU301139
MJS046	P. cryptogea (A2)	P. cryptogea	Pinus radiata	Nannup	Clade 8	EU301140
TCH010	Unidentified	P. cryptogea	B. ilicifolia	Mogumber	Clade 8	EU301141
VHS14620	Waterhouse Group 6		X. gracilis	Fitzgerald River NP	Clade 8	EU301142
DDS3543	P. megasperma	P.sp.9	D. falcata	Fitzgerald River NP	Clade 9	EU301143
DDS3565	P. megasperma	P.sp.9	Adenanthos cuneata	Fitzgerald River NP	Clade 9	EU301144
DDS3584	P. megasperma	P.sp.9	B. baxteri	Fitzgerald River NP	Clade 9	EU301145
DDS3601	P. megasperma	P.sp.9	D. cirsioides	Fitzgerald River NP	Clade 9	EU301146
DCE177,	P. megasperma	P.sp.9	P. radiata (H)-1980	Nannup	Clade 9	EU301147
MJS129	D	D on 0	D nadiata	Nonnun	Clade 0	EU201149
MJS186	P. megasperma	P.sp.9	P. radiata	Nannup Cana Arid	Clade 9 Clade 0	EU301148
TCH003	P. megasperma	P.sp.9	Soil, native heathland Soil	Cape Arid	Clade 9 Clade 9	EU301149
VHS16134 DDS3884	P. megasperma P. boehmeriae	P.sp.9 P. boehmeriae		Fitzgerald River NP	Clade 9 Clade 10	EU301150 EU301151
0053004	r. voenmeriae	r. ovenmeride	Persoonia longifolia	Nannup	Claue 10	EU301151

^a Isolates in bold were included in the phylogenetic study (Fig. 1).

^b Abbreviations of isolates and culture collections: IMI = CABI Bioscience (Imperial Mycological Institute), UK; WAC = Department of Agriculture and Food Western Australia Plant Pathogen Collection, Perth, Australia; VHS = Vegetation Health Service; DDS = earlier prefix of VHS collection. The following collections have been incorporated into the VHS collection: MJS = M. J. C. Stukely; TCH = T. C. Hill; HSA = Hart, Simpson & Associates; DCE = E. M. Davison.

^c Cooke et al. (8).

soil extract in daylight for ~ 24 h. Gametangia were examined in cultures 7 to 10 days old where they were produced in pure culture; for heterothallic isolates, cultures were paired with tester isolates of known mating type and examined at intervals for up to 3 months.

DNA isolation, amplification, and sequencing. The Phytophthora isolates were grown on half-strength potato dextrose agar (PDA) (Becton, Dickinson and Company, Sparks, MD, USA; 19.5 g of PDA, 7.5 g of agar, and 1 liter of distilled water) at 20°C for 2 weeks, and the mycelium was harvested and placed in a 1.5-ml sterile Eppendorf tube. Harvested mycelium was frozen in liquid nitrogen, ground to a fine powder, and genomic DNA was extracted as described previously (2). The region spanning the ITS1-5.8S-ITS2 region of the ribosomal DNA was amplified using the primers ITS-6 (5' GAA GGT GAA GTC GTA ACA AGG 3') (8) and ITS-4 (5' TCC TCC GCT TAT TGA TAT

GC 3') (42). The PCR reaction mixture, PCR conditions, the clean-up of products, and sequencing were as described previously (2).

Phylogenetic analysis. In order to compare *Phytophthora* isolates used in this study with other closely related *Phytophthora* spp., additional sequences were obtained from GenBank. Sequence data were initially assembled using Sequence Navigator version 1.01 (Applied Biosystems, Foster City, CA) and aligned in Clustal X (38). Manual adjustments were made visually by inserting gaps where necessary. All sequences derived in this study were deposited in GenBank, and accession numbers are shown in Tables 1 and 2.

Isolates representing each of the known species and undescribed taxa obtained in this study were compiled in a single dataset, and parsimony analysis was performed in PAUP version 4.0b10 (D. L. Swofford. 2003. Phylogenetic analysis using parsi-

mony (*and other methods). Sinauer Associates, Sunderland, MA). The most parsimonious trees were obtained using heuristic searches with random stepwise addition in 100 replicates, with the tree bisection-reconnection branch-swapping option on and the steepest-descent option off. Maxtrees were unlimited, branches of zero length were collapsed, and all multiple, equally parsimonious trees were saved. Estimated levels of homoplasy and phylogenetic signal (retention and consistency indices) were determined (20). Branch and branch node support was determined using 1,000 bootstrap replicates (16).

Bayesian analysis was conducted on the same individual and combined dataset as that used in the parsimony analysis. First, MrModeltest v.2.5 (J. A. A. Nylander. 2004. Evolutionary Biology Centre, Uppsala University) was used to determine the best nucleotide substitution model. Phylogenetic analyses were performed with

Table 2. Identity (based on morphology and internal transcribed spacer [ITS] sequence data), host, location, and GenBank accession numbers for representative *Phytophthora* isolates from Cooke's clade 6 from Western Australia (WA)^a

Culture identification ^b	Morphology identification	Sequence identification	Host	Location	Cooke's clade ^c	GenBank accession no.
DCE444, TCH012, IMI329666	P. drechsleri (A1)	P.sp.3	Banksia attenuata	Badgingarra	Clade 6	EU301152
DDS3760	P. drechsleri	P.sp.3	Soil - gravel pit - forest	Manjimup	Clade 6	EU301153
MJS148	Unidentified	P.sp.3	Pinus radiata	Nannup	Clade 6	EU301154
VHS13530	Unidentified	P.sp.3	Eucalyptus marginata	Dwellingup	Clade 6	EU301155
VHS13567	Unidentified	P.sp.3	E. marginata	Dwellingup	Clade 6	EU301156
VHS13761	Unidentified	P.sp.3	E. marginata	Dwellingup	Clade 6	EU301157
VHS16164	Waterhouse Group 6	P.sp.3	Banksia grandis	Pemberton	Clade 6	EU301158
DCE68, INH116468	P. megasperma var. sojae	P.sp.7	Soil, Lupin bait in 1965	Byford (Hwy)	Clade 6	EU301171
MJS235	Unidentified	P.sp.7	P. radiata	Nannup	Clade 6	EU301172
MJS238	Unidentified	P.sp.7	P. radiata	Nannup	Clade 6	EU301173
VHS9854	P. megasperma	P.sp.7	Xanthorrhoea preissii	Lancelin	Clade 6	EU301174
DDS3642	P. drechsleri (A1)	P.sp.8	Water, pear bait	Albany	Clade 6	EU301176
VHS2713	Unidentified	P.sp.8	Soil	Walpole	Clade 6	EU301175
TCH009, IMI329669	P. megasperma var. sojae	P.sp.10	B. prionotes	Badgingarra	Clade 6	EU593265
VHS17779	P. megasperma	P.sp.10	B. grandis	Perth (Bold Park)	Clade 6	EU593263
VHS17085	Unidentified	P.sp.11	Banksia sp.	Hopetoun	Clade 6	EU593262
VHS19173	Waterhouse Group 6	P.sp.11	X. preissii	Wilga	Clade 6	EU869199
DDS3599	P. megasperma	Unique	X. platyphylla	Fitzgerald River NP	Clade 6	EU593258
VHS16108	Waterhouse Group 6	Unique	Soil, native bush	Fitzgerald River NP	Clade 6	EU593259
VHS16115	Waterhouse Group 6	Unique	Soil, native bush	Fitzgerald River NP	Clade 6	EU593260
VHS17350	Waterhouse Group 6	Unique	Water, native bush	Badgingarra	Clade 6	EU593261
VHS5185	Unidentified	Unique	Soil, native bush	Pemberton	Clade 6	EU593264
VHS14801	Unidentified	P. taxon personii	Grevillea mccutcheonii	Busselton	Clade 6	EU301169
DDS3753	Unidentified	P. taxon PgChlamydo	Soil, native forest	Manjimup	Clade 6	EU301160
VHS6595	Unidentified	P. taxon PgChlamydo	Soil, native forest	Manjimup	Clade 6	EU301159
VHS17175	Waterhouse Grp. 5 or 6	P. asparagi	B. media	Esperance	Clade 6	EU301167
VHS17644	Waterhouse Grp. 5 or 6	P. asparagi	Lomandra sonderi	Perth (Murdoch)	Clade 6	EU301168
DDS1540	Unidentified	P. inundata	Adenanthos cuneata	Esperance	Clade 6	EU301161
MJS262	Unidentified	P. inundata	B. littoralis	Grimwade	Clade 6	EU301162
VHS13920, WAC12880	Unidentified	P. inundata	X. preissii	Wilga	Clade 6	EU301163
VHS16932	Unidentified	P. inundata	X. preissii	Rocky Gully	Clade 6	EU301164
VHS2910	Unidentified	P. inundata	Soil, native forest	Mount Barker	Clade 6	EU301165
VHS17183	P. megasperma	P. megasperma	X. platyphylla	Esperance	Clade 6	EU301166

^a Isolates in bold were included in the phylogenetic study (Fig. 2).

^b Abbreviations of isolates and culture collections: IMI = CABI Bioscience (Imperial Mycological Institute), UK; WAC = Department of Agriculture and Food Western Australia Plant Pathogen Collection, Perth, Australia; VHS = Vegetation Health Service; DDS = earlier prefix of VHS collection. The following collections have been incorporated into the VHS collection: MJS = M. J. C. Stukely; TCH = T. C. Hill; HSA = Hart, Simpson & Associates; DCE = E. M. Davison.

^c Cooke et al. (8).

MrBayes v. 3.1 (28) applying a general time reversible (GTR) substitution model with gamma (G) and proportion of invariable site (I) parameters to accommodate variable rates across sites. Two independent runs of Markov Chain Monte Carlo (MCMC) using 4 chains were run over 1,000,000 generations. Trees were saved each 1,000 generations, resulting in 10,001 trees. Burn-in was set at 50,001 generations (i.e., 51 trees), well after the likelihood values converged to stationary, leaving 9,950 trees from which the consensus trees and posterior probabilities were calculated.

RESULTS

Morphological identification. The morphology of many of the *Phytophthora* isolates was examined when they were first isolated and the morphological identification initially provided (Tables 1 and 2). *P. nicotianae, P. cryptogea, P. boehmeriae, P. citricola, P. megasperma,* and *P. drechsleri* were identified, and several other isolates were placed in Waterhouse groups 1, 5, and 6. However, there were also many isolates that could not be assigned to a known species or even into a Waterhouse morphological group due to the lack of useful morphological features (usually oogonia).

Phylogenetic analysis. After an initial phylogenetic analysis containing all sequenced Phytophthora isolates from WA, the dataset was split into two. The first dataset included known species and undescribed taxa from ITS clades 1, 2, 4, 7, 8, 9, and 10, and the second included known species and undescribed taxa from ITS clade 6 (sensu Cooke et al. 8). Due to the size of the first dataset, only three representative isolates were included for each of the undescribed Phytophthora taxa from WA. In addition, only those known Phytophthora spp. that were phylogenetically close to the undescribed taxa were included in the analysis. Included in the second dataset were isolates of undescribed Phytophthora taxa from ITS clade 6 with sequences similar to isolates from the current study, including representatives of all undescribed Phytophthora taxa considered by Brasier et al. (4).

The first dataset consisted of 909 characters, of which 472 were parsimony informative. The dataset contained significant phylogenetic signal compared to 1,000 random trees (P < 0.01, g1 = -0.78). Heuristic searches resulted in 8 most parsimonious trees of 1,260 steps (CI = 0.61, RI = 0.92) (Fig. 1, TreeBASE SN4084). The topology of the Bayesian tree was very similar (TreeBASE SN4084). Isolates obtained in this study resided in ITS clades 1, 2, 4, 7, 8, 9, and 10. Several isolates corresponded to known species (P. nicotianae, clade 1; P. cryptogea, clade 8; and P. boehmeriae, clade 10). P. taxon niederhauserii (clade 7) was identified solely on

The second dataset consisted of 911 characters, of which 408 were parsimony informative. The dataset contained significant phylogenetic signal compared to 1,000 random trees (P < 0.01, g1 = -1.54). Heuristic searches resulted in 1,000 most parsimonious trees of 849 steps (CI = 0.70, RI = 0.89) (Fig. 2, TreeBASE SN4084). The topology of the Bayesian tree was very similar (TreeBASE SN4084). Isolates obtained in this study resided in ITS clade 6. Only a single isolate (VHS17183), originally identified as P. megasperma on morphological characters, had 100% similarity to sequences available for P. megasperma on GenBank. Others isolates thought to be P. megasperma based on morphological characters all represented undescribed taxa in clade 6 and clade 9 (Tables 1 and 2). Several other isolates corresponded to P. inundata, both on morphology and ITS sequence data. Two isolates, VHS17175 and VHS17644, had almost identical ITS sequence to the recently described P. asparagi (29). Several more isolates have been identified only by their 100% similarity to sequences available on GenBank for the as yet undescribed taxa P. taxon personii and P. taxon Pgchlamydo.

The ITS analysis revealed three undescribed Phytophthora taxa (P.sp.3, P.sp.8, and P.sp.11) and five unique isolates (DDS3599, VHS16108, VHS16115, VHS17350, and VHS5185) (Fig. 2, Table 2). The sequences from isolates in P.sp.3 were all identical (including sequences from isolates not included in the phylogenetic study) and were a close match to Phytophthora isolate 3267 of unknown origin. P.sp.8 was represented by two isolates (VHS2713 and DDS3642) with identical sequence and was a closest sequence match to three unique isolates from WA (VHS17350, VHS5185, and VHS16108). P.sp.11 was also represented by two isolates (VHS 17085 and VHS 19173) and was closest to another unique isolate from WA (VHS16115). Together, these isolates, P.sp.3, 8, 11, and four of the unique isolates from WA and Phytophthora isolate 3267, form a clade with 74% bootstrap support and a Bayesian probability of 0.90.

Also identified in clade 6 was P.sp.7, which formed a well supported clade with *P.* taxon raspberry isolates P1049 and P896. These unidentified isolates had been collected from raspberries in Australia and had been used in previous phylogenetic studies (4,6). Two isolates were thought to represent P.sp.10; however, when sequences of other undescribed *Phytophthora* isolates were included, the clustering of these isolates was less resolved. The unknown *Phytophthora* isolates closest to P.sp.10 are from *Malus* in the United States (P462), *Prunus* in Switzerland (P532), and SCRP223 of unknown origin. A single unique WA isolate DDS3599 had identical ITS sequence to an unknown *Phytophthora* isolate 4 FFL-2008 from Hungary (Fig. 2).

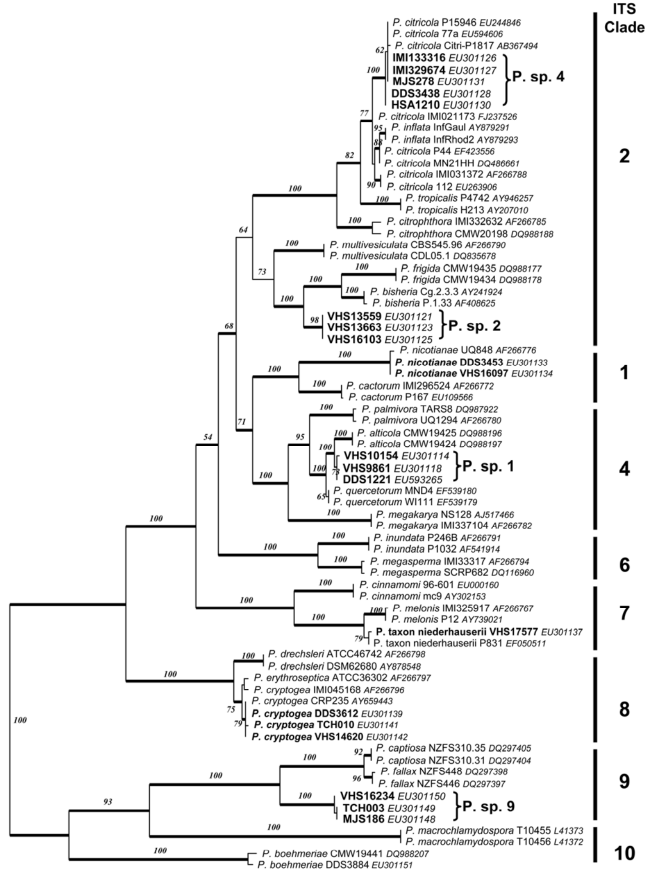
Distribution of *Phytophthora* **spp. in WA.** The known associations of *Phytophthora* species with dead and dying native plant species in WA are shown in Table 3. Besides *P. cinnamomi*, the most commonly isolated species in WA are *P. cryptogea*, *P. inundata*, P.sp.2, P.sp.3, P.sp.4, and P.sp.9. *P. cryptogea* and P.sp.4 have the largest host range and the widest distribution. The distribution of P.sp.9 and P.sp.1 is limited to the Midwest, West, and South Coast Land Management Regions of WA (Table 3, Fig. 3). Other *Phytophthora* spp. were isolated too infrequently to draw any conclusions on their distribution.

DISCUSSION

Using molecular techniques to reevaluate unidentified isolates from the VHS collection of *Phytophthora* isolates from natural ecosystems in WA, *P. inundata, P. asparagi,* and several named but as yet undescribed taxa, *P.* taxon PgChlamydo, *P.* taxon personii, and *P.* taxon niederhauserii, were identified for the first time. The presence of other species, *P. nicotianae, P. boehmeriae, P. cryptogea,* and *P. megasperma,* was confirmed. In addition, based on phylogenetic analysis, nine undescribed *Phytophthora* taxa were recognized.

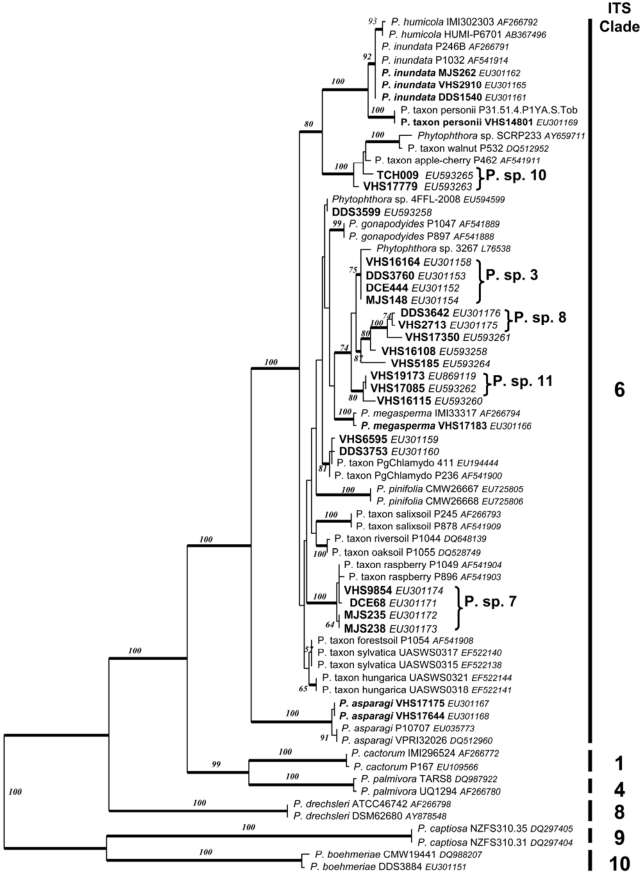
P. inundata, described in Europe in 2003 (6), was identified based on phylogenetic analysis from several locations in the southwest of WA, where it has been associated with dying native plants (36). Some of these isolates were recovered in the 1980s. P. boehmeriae has been reported from most continents except North America. It is best known as the causal agent of brown rot of citrus and boll rot of cotton (15) and has also been implicated in gummosis of Acacia mearnsii in South Africa and Brazil (14). In WA, it was isolated from soil baits at only one site, collected beneath a dead, mature Persoonia longifolia (10). P. asparagi has just been described as a pathogen of asparagus in Michigan, USA (29). P. asparagi has been recorded previously in Australia from diseased agave in the Royal Botanic Gardens in Melbourne (9), but it is intriguing that isolates with identical ITS sequences have also been isolated from natural ecosystems in WA

P. cryptogea has a cosmopolitan distribution and causes disease on many hosts (33). When sequencing of the VHS isolates



5 changes

Fig. 1. One of the eight most parsimonious trees of 1,260 steps resulting from analysis of the internal transcribed spacer (ITS) sequence data of *Phytophthora* isolates. Bootstrap values of the branch nodes are given in italics. Branches with posterior probabilities resulting from Bayesian analysis of greater than 0.85 are thickened. Isolates from this study are in bold. The ITS clade as designated by Cooke et al. (8) is indicated on the right.



5 changes

Fig. 2. One of the 1,000 most parsimonious trees of 849 steps resulting from analysis of the internal transcribed spacer (ITS) sequence data of *Phytophthora* isolates. Bootstrap values of the branch nodes are given in italics. Branches with posterior probabilities resulting from Bayesian analysis of greater than 0.85 are thickened. Isolates from this study are in bold. The ITS clade as designated by Cooke et al. (8) is indicated on the right.

commenced, a large group of isolates with identical sequences were observed which had the closest sequence match to P. cryptogea. Initially, it was thought to be a distinct species, and was named P.sp.5. However, as additional sequence information appeared on GenBank, the WA isolates clustered within P. cryptogea sensu lato. This complex is currently undergoing extensive revision, which will result in the description of several new species (M. D. Coffey, personal communication). The isolates from WA may reside in one of these new species. P. cryptogea is commonly isolated in WA, but its role in the natural environment is currently unclear.

P. megasperma has previously been reported from dying native vegetation in WA (35). However, after molecular reevaluation, many isolates thought to be P. megasperma (ITS clade 6) based on morphology actually represent a new taxa belonging to ITS clade 9, and have been provisionally named P.sp.9. Other isolates represent the new taxa P.sp.7 and P.sp.10 (ITS clade 6). Phylogenetic analysis resolved only a single, recent, WA isolate (VHS17183) as a "true" P. megasperma. The earlier identification of P. megasperma on morphological characters alone was thus not reliable, and the ITS sequence data confirm that the P. megasperma morpho-species in WA represent a complex of several distinct taxa.

As with *P. megasperma*, so few isolates of *P.* taxon Pgchlamydo have been recorded among the VHS collection, that their role in the landscape is considered minor. In Europe, *P.* taxon Pgchlamydo is closely related to and inhabits a similar niche to *P. gonapodyides* (5). *P. gonapodyides* is common in North America and Europe and has been reported from Australia, New Zealand, and Chile, where it is predominantly known as a minor root and seedling pathogen with a restricted host range (15). Brasier et al. (4) suggests *P. gonapodyides* and some other taxa in ITS clade 6 may play a significant role in the breakdown of green plant litter, especially in waterways.

Hardy and Sivasithamparam (19) reported P. cactorum, P. citricola, P. crypto-

gea, P. drechsleri, P. megasperma, and P. nicotianae from container-grown nursery plants in WA. Davison et al. (11) reported several *Phytophthora* species on nursery plants imported into WA: P. nicotianae, P. palmivora, P. drechsleri, P. citrophthora, P. taxon parvispora, and P. taxon nieder-hauserii. These were intercepted prior to placement in WA nurseries. Among the VHS collection were isolates of P. crypto-gea, P. nicotianae, and P. taxon nieder-hauserii and an isolate of P. megasperma, indicating that these species are estab-

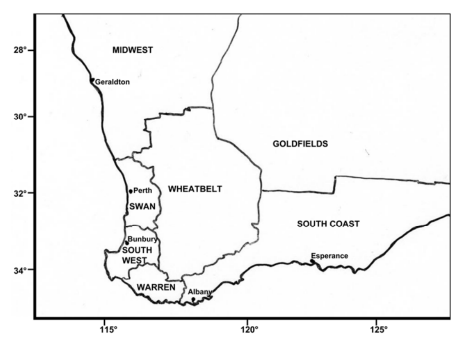


Fig. 3. Map of the southwest of Western Australia showing Department of Environment and Conservation Land Management Regions.

Table 3. Numbers of isolates of each *Phytophthora* species identified from the Vegetation Health Service (VHS) collection (excluding *P. cinnamomi*), including host and distribution across the southwest of Western Australia based on Land Management Regions (Fig. 3)

Phytophthora species	No. isolates	Host species	Distribution (region)
P. nicotianae	3	Chamaelaucium sp., Xanthorrhoea platyphylla, Dryandra cirsioides	Swan, South Coast
P. taxon personii	1	Grevillea mccutcheonii	South West
P. asparagi	2	Banksia media, Lomandra sonderi	Swan, South Coast
P. taxon PgChlamydo	2	Soil, native forest	Warren
P. inundata	15	Banksia littoralis, Xanthorrhoea preissii, Adenanthos cuneata	South West, Warren, Wheatbelt, South Coast
P. megasperma	1	X. platyphylla	South Coast
P. taxon niederhauserii	2	Banksia prionotes	Midwest
P. cryptogea	28	Banksia grandis, B. ilicifolia, Dryandra cirsiodes, Xanthorrhoea gracilis, X. platyphylla, Patersonia sp., Crowea angustifolia, Isopo- gon buxifolius, Daviesia sp.; Pinus radiata (plantation)	Midwest, Swan, South West, Warren, Wheatbelt, South Coast
P. boehmeriae	1	Persoonia longifolia	South West
P.sp.1	10	Banksia menziesii, B. littoralis, B. attenuata, B media	Midwest, South West, South Coast
P.sp.2	21	Eucalyptus marginata, Corymbia calophylla, B. grandis, Dryandra squarrosa, Patersonia xanthina	Swan, South West, Warren
P.sp.3	15	<i>E. marginata, B. attenuata, B. grandis; Pinus radiata</i> (plantation)	Midwest, Swan, South West, Warren
P.sp.4	64	E. marginata, B. attenuata, B. grandis, B. littoralis, B. menziesii, B. prionotes, Conospermum sp., Leucopogon verticillatus, X. gracilis, Podocarpus drouyniana, Patersonia sp., Bossiaea sp., Gastrolobium spinosum; Pinus radiata (plantation)	Midwest, Swan, South West, Warren, Wheatbelt, South Coast
P.sp.7	4	X. preissii; Pinus radiata (plantation)	Midwest, Swan, South West
P.sp.8	3	Soil and water baits, native forest	Midwest, Warren, South Coast
P.sp.9	23	B. attenuata, B. baxteri, D. cirsioides, D. falcata, A. cuneata, Isopo- gon sp.; Pinus radiata (plantation)	Midwest, South West, South Coast
P.sp.10	2	B. prionotes, B. grandis	Midwest, Swan
P.sp.11	2	Banksia sp., X. preissii	South West, South Coast

lished in natural ecosystems. The other species reported in nursery plants have not been collected from natural ecosystems in WA.

P.sp.4 is morphologically very similar to P. citricola, and it resides within the P. citricola complex in clade 2. However, there is strong bootstrap support for the separation of these isolates into a new taxa. P.sp.4 is the most commonly isolated Phytophthora taxa in WA after P. cinnamomi. It has been found on 14 native host plant species representing seven families, from a wide range of locations, as well as on Pinus radiata, an introduced tree species in WA. Until recently, P.sp.4 was thought to be P. citricola, but after sequencing 73 isolates, not a single isolate was found with ITS sequence corresponding exactly to P. citricola sensu stricto. It has long been recognized that P. citricola is most likely a species complex (27). The widespread nature of this undescribed taxa in WA and the lack of consistent association with "high impact" disease suggests this species may be indigenous to the region. Interestingly, three sequences that are identical to P.sp.4 have been recently submitted to GenBank: EU244846 from Spain, EU594606 from Hungary, and AB367494 from Japan.

P.sp.2 has been isolated from the lower stem and roots of dying 1- to 2-year-old jarrah (Eucalyptus marginata) seedlings in rehabilitated open-cut bauxite mine pits in the WA jarrah forest since 2003, with unusually large numbers of isolations in the summer of 2004-05 (37). It has also been isolated from similar aged Corymbia calophylla in the same sites (M. J. C. Stukely, unpublished data) and from undisturbed jarrah forest on several occasions since the mid-1980s (37). P.sp.2 is morphologically similar to P. citricola, but most closely related to the recently described P. bisheria and P. frigida, the former from plants in the Rosaceae on three continents (1) and the latter from Eucalyptus in South Africa (26). The pathogenicity of P.sp.2 to jarrah is currently under investigation.

P.sp.1 has been isolated only from dead and dying Banksia spp. in coastal areas of the southwest of WA. It has a morphology similar to P. cactorum, but has the closest ITS sequence match to P. alticola and P. quercetorum, the former isolated and described from Eucalyptus in South Africa (26) and the latter from Quercus in the United States. P.sp.9 has been recovered from six WA native hosts, but not from Eucalyptus spp. Morphologically, P.sp.9 resembles *P. megasperma*, and for several years, the death of young Pinus radiata in plantations in the Donnybrook Sunklands (Blackwood Plateau) in WA was attributed to P. megasperma (7). However, molecular re-evaluation places these isolates in ITS clade 9, most closely related to the newly described P. captiosa and P. fallax from

Many new undescribed taxa and unique isolates were recognized within ITS clade 6. Among the isolates in the VHS collection was a single isolate, originally designated P.sp.6, which corresponds to P. taxon personii (Z. G. Abad, personal communication). The most commonly encountered taxa in WA from ITS clade 6 is P.sp.3. This taxa is morphologically very similar to P. drechsleri and P. cryptogea, and has been isolated from soil associated with dying E. marginata and Banksia spp., as well as P. radiata. There is very little sequence variation among P.sp.3 isolates, and it is widely distributed in WA. In the phylogenetic analysis, there were eight other isolates from WA which formed a strongly supported clade together with P.sp.3. Four of these isolates were unique and did not correspond to any sequence currently on GenBank. The other four isolates fell into two subclades designated P.sp.8 and P.sp.11. The role of these ITS clade 6 isolates in WA ecosystems is unclear. P.sp.3 is commonly isolated from stream baiting within natural ecosystems (unpublished data), and perhaps the other undescribed species in this clade will also be found in water bodies once more baiting is conducted.

In conclusion, based on phylogenetic analysis, nine potentially new and undescribed taxa of Phytophthora can be distinguished among isolates from natural ecosystems in the southwest of WA. Several of these undescribed taxa are morphologically indistinguishable from known species such as P. citricola, P. megasperma, and P. drechsleri. In some cases, the new taxa are closely related to known species with similar morphology (e.g., P.sp.4 and P. citricola). However, the DNA sequences of other new taxa show that they are not closely related to morphologically similar species (e.g., P.sp.3 and P. drechsleri, P.sp.9 and P. megasperma). Further work is planned to describe the new taxa and their relationships, and to test their pathogenicity, so that an estimate of the level of threat they pose to native vegetation can be made. It is not known if these new Phytophthora taxa are endemic or introduced, and thus their quarantine status within Australia and elsewhere remains unknown.

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