Stem cankers on sunflower (Helianthus annuus) in Australia reveal a complex of pathogenic Diaporthe (Phomopsis) species

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Kev words

Diaporthe gulyae Diaporthe kochmanii Diaporthe kongii ITS phylogeny sunflower taxonomy TEF-1α

Abstract The identification of Diaporthe (anamorph Phomopsis) species associated with stem canker of sunflower (Helianthus annuus) in Australia was studied using morphology, DNA sequence analysis and pathology. Phylogenetic analysis revealed three clades that did not correspond with known taxa, and these are believed to represent novel species. Diaporthe gulyae sp. nov. is described for isolates that caused a severe stem canker, specifically pale brown to dark brown, irregularly shaped lesions centred at the stem nodes with pith deterioration and midstem lodging. This pathogenicity of D. gulyae was confirmed by satisfying Koch's Postulates. These symptoms are almost identical to those of sunflower stem canker caused by D. helianthi that can cause yield reductions of up to 40 % in Europe and the USA, although it has not been found in Australia. We show that there has been broad misapplication of the name D. helianthi to many isolates of Diaporthe (Phomopsis) found causing, or associated with, stem cankers on sunflower. In GenBank, a number of isolates had been identified as D. helianthi, which were accommodated in several clades by molecular phylogenetic analysis. Two less damaging species, D. kochmanii sp. nov. and D. kongii sp. nov., are also described from cankers on sunflower in Australia.

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INTRODUCTION

Phomopsis species are widespread and occur on a diverse range of host plants as pathogens, endophytes or saprobes (Uecker 1988). The morphological characters that define Phomopsis are dark eustromatic or pycnidial conidiomata containing elongated phialides with cylindrical, well-developed collarettes that form two types of hyaline conidia: 1-celled α -conidia that are biguttulate, fusiform, and easily germinate on artificial media, and ß-conidia that are filiform and rarely germinate (Wehmeyer 1933, Sutton 1980). Species of Phomopsis represent anamorphs of Diaporthe (Ascomycota, Diaporthales, Valsaceae) with at least 180 connections given by Uecker (1988), which represents about 80 % of named Phomopsis species. The name Diaporthe Nitschke (1870) precedes Phomopsis Sacc. & Roum. in Saccardo (1884).

Host association has often been the basis for species identification in Diaporthe and Phomopsis, as morphological and culture characteristics are inadequate or unreliable for species differentiation (van Rensburg et al. 2006). Recent studies have demonstrated that a number of Phomopsis species have wide host ranges (van Niekerk et al. 2005, Santos & Phillips 2009, Ash et al. 2010), and more than one species can occur on a single host (Mostert et al. 2001, Santos & Phillips 2009).

Molecular phylogenies, especially those derived from DNA sequence analyses of the ribosomal internal transcribed spacer (ITS) regions of the nuclear ribosomal RNA genes and translation elongation factor-1 α (TEF-1 α) have been used to identify species (Mostert et al. 2001, van Niekerk et al. 2005, van Rensburg et al. 2006, Santos & Phillips 2009, Ash et al. 2010). The polyphyletic status of D. helianthi has been recognised by Rekab et al. (2004). Hyde et al. (2010) suggested that discarding the host-based species concept was the first step in the development of a useful and reliable classification for Phomopsis and highlighted that there had been much confusion around the application of species names, drawing particular attention to the name D. helianthi.

Stem canker attributed to D. helianthi (anamorph P. helianthi) has become one of the most important diseases of sunflower (Helianthus annuus) worldwide since first described from the former Yugoslavia (Muntañola-Cvetković et al. 1981). Yield reductions of up to 40 % have been recorded in Europe (Masirevic & Gulya 1992) including the former Yugoslavia as well as France where it was considered a major pathogen of sunflower (Battilani et al. 2003, Debaeke et al. 2003). Diaporthe helianthi is also widespread in the sunflower growing regions of the USA (Gulya et al. 1997) but has not been reported from Australia.

Muntañola-Cvetković et al. (1985) found that multiple Phomopsis species were associated with cankers on sunflower in the former Yugoslavia, although only P. helianthi was responsible for the serious disease outbreaks. Gulya et al. (1997) suggested that pathogenic Phomopsis species on sunflower might consist of more than one species or biotype with apparent biological differences between the isolates from Europe and the USA. Miric et al. (2001) raised the possibility that several pathogenic Phomopsis species occurred on sunflower in Australia.

In 2009, lodging and premature senescence caused significant damage to sunflower crops in New South Wales (NSW), and to a lesser extent in Queensland (Qld), Australia, after extended periods of wet weather. The symptoms included pith damage behind elongated, brown to brown-black lesions, which weakened stems and led to mid-stem lodging as the heads filled. The aim of this study was to use morphological, molecular and pathogenicity studies to clarify the identity of the Diaporthe (Phomopsis) species occurring on sunflower in Australia.

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Table 1	Diaporthe cultures isolated from sunflower investigated in this study.	
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Species	Isolate number (BRIP) ¹	Locality	Source	Sunflower Hybrid/Wild	Virulence Rating ²	GenBank Accession numbers	
						ITS	TEF-1α
Diaporthe gulyae	53158	Goran Lake, NSW	stem	Wild H. annuus	4	JF431284	JN645799
	53166	Premer, NSW	seed	Ausigold 62	4	JF431289	JN645801
	53172	Premer, NSW	seed	Hyoleic 41	5	JF431290	JN645802
	53159	Premer, NSW	seed	Advantage	5	JF431291	JN645800
	54030	Nobby, Qld	stem	Sunbird 7	5	JF431292	JN645808
	54029	Hermitage, Qld	stem	Hyoleic 41	4	JF431293	JN645807
	54028	Hermitage, Qld	stem	Hyoleic 41	5	JF431294	JN645806
	54027	Ryeford, Qld	leaf	Sunbird 7	5	JF431297	JN645805
	54026	Ryeford, Qld	leaf	Sunbird 7	5	JF431298	JN645804
	54025	Ryeford, Qld	leaf	Sunbird 7	4	JF431299	JN645803
Diaporthe kochmanii	54033	Gatton, Qld	stem	Experimental	2	JF431295	JN645809
	54034	Gatton, Qld	stem	Experimental	3	JF431296	JN645810
Diaporthe kongii	54032	Childers, Qld	stem	Female	3	JF431300	JN645798
. 0	54031	Childers Qld	stem	Female	3	JF431301	JN645797

¹ Ex-type cultures are in **bold**.

² At 14 d after inoculation where 0 = no discolouration or very slight discolouration or scarring at site of inoculation; 1 = low level discolouration at site of inoculation; 2 = very small lesion or slight discolouration 1–2 mm diam; 3 = necrotic lesions 2–5 mm, some light stem streaking, leaf wilting and twisting; 4 = lesions 5–10 mm diam, significant necrosis and dark stem streaking, leaf and plant wilting, stunting, and some lodging; 5 = very severe necrosis and lesions, dark streaking, leaf necrosis, twisting and wilting, stunting, lodging or plant death.

MATERIAL AND METHODS

Isolates

Over 300 isolates of *Diaporthe (Phomopsis*) were obtained from stems, leaves and seed of both cultivated and wild sunflower plants exhibiting symptoms of stem canker across NSW and Qld. Small excised stem and leaf pieces with brown or brown-ish black lesions were surface-sterilised by dipping into 90 % ethanol and flaming briefly prior to placement on 1.5 % water agar amended with 100 μ g/mL streptomycin sulphate (WAS) in 9 cm diam Petri dishes. Cultures that grew from this tissue were incubated for up to 3 wk to induce pycnidial formation. For seed isolations, seeds harvested from infected crops and individual plants were incubated without surface sterilisation on WAS in Petri dishes for up to 14 d to allow pycnidia to develop.

For all isolations, conidia oozing from pycnidia were streaked onto potato-dextrose agar (Oxoid) (PDA) amended with 100 µg/mL streptomycin sulphate (PDAS). Hyphal tips were then taken from all isolates and grown on PDAS to establish pure isolates. Cultures were incubated for 7 d under ambient light at 23–25 °C. For pathogenicity experiments, 7 d old cultures were used to provide inocula. Fourteen selected isolates representing a range of virulence symptoms and morphological characteristics were deposited in the Plant Pathology Herbarium (BRIP), Brisbane, Australia as both living and dried cultures (Table 1).

Morphology

For fungal morphology, isolates were grown on PDA with pieces of sterilised wheat stems placed on the surface and incubated under 12 h near-ultraviolet light / 12 h dark (Smith 2002) at 25 °C. Fungal structures were mounted on glass slides in lactic acid (100 % v/v) for microscopic examination after 28 d of incubation. Means and standard deviations (SD) of selected structures were made from at least 20 measurements. Ranges were expressed as (min.–) mean-SD – mean+SD (–max.) with values rounded to 0.5 μ m. Images were captured with a Leica DFC 500 camera attached to a Leica DM5500B compound microscope with Nomarski differential interference contrast.

For colony morphology, 3 d old cultures on 9 cm diam plates of PDA and oatmeal agar (OA) (Oxoid) that had been grown in the dark at 23 °C were grown for a further 7 d under 12 h near-ultraviolet light / 12 h dark. Colony colours (surface and reverse) were rated according to the colour charts of Rayner (1970).

DNA isolation, amplification and analyses

Mycelia were scraped off PDA cultures and macerated with 0.5 mm glass beads (Daintree Scientific) in a Tissue Lyser (QIAGEN). Genomic DNA was then extracted with the Gentra Puregene DNA Extraction kit (QIAGEN) according to the manufacturer's instructions.

The primers ITS1 and ITS4 (White et al. 1990) were used to amplify the ITS region of the ribosome genes. To further differentiate *D. angelicae*, *D. stewartii*, *D. gulyae* and *P. dauci*, the primers EF1-728F (Carbone & Kohn 1999) and EF2 (O'Donnell et al. 1998) were used to amplify part of the translation elongation factor-1alpha (TEF-1 α) gene. Both the ITS and TEF loci were amplified with the Phusion High-Fidelity PCR Master Mix (Finnzymes). The PCR products were purified with the QIAquick PCR Purification Kit (QIAGEN) and sequenced on the 3730x/ DNA Analyzer (Applied Biosystems) using the amplifying primers.

The sequences generated in this study were assembled using Vector NTi Advance v. 11.0 (Invitrogen) and deposited in Gen-Bank (Table 2). These sequences were aligned with sequences from representative *Diaporthe/Phomopsis* species from Gen-Bank (Table 2) in MEGA v. 5.05 (Tamura et al. 2011).

The sequences of *Leucostoma persoonii* and *Valsa cerato-sperma* were used as outgroups in the ITS dataset, whilst sequences of *Leucostoma niveum* and *Valsa ambiens* were used as outgroups in the TEF-1 α dataset. Alignment gaps were treated as missing character states and all characters were unordered and of equal weight.

The ITS and TEF-1 α phylogenetic trees were inferred in MEGA v. 5.05 by Maximum Likelihood (ML). Modeltest in MEGA v. 5.05 determined that the K2+G and HKY+G models were the most suitable nucleotide substitution models for ITS and TEF-1 α , respectively. Bootstrap support values with 1 000 replications were calculated for tree branches. The sequences obtained from GenBank are listed by their taxon names followed by strain numbers in the trees (Fig. 1, 2). Nomenclatural novelties were deposited in MycoBank (www. MycoBank.org) (Crous et al. 2004).

Pathogenicity

Pathogenicity was determined by inoculating plants of the sunflower hybrid Hyoleic 41 at the V6–V8 (Schneiter & Miller 1981) growth stage and grown in a cabinet under a 25 °C 12 h light / 20 °C 12 h dark cycle using two methods, wound inoculation and mycelium contact. The wound inoculation method (adapted

Table 2 Reference isolates used in the phylogenetic analyses.

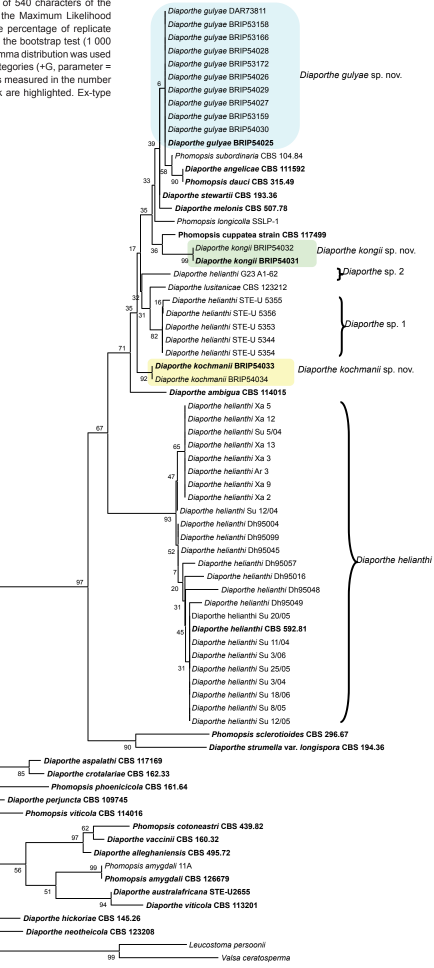
Species	Isolate no. 1, 5	Host	GenBank accession numbers		Reference	
			ITS ² TEF-1α ³			
Diaporthe alleghaniensis	CBS 495.72	Betula alleghaniensis	FJ889444	GQ250298	Santos et al. 2010	
Diaporthe ambigua	CBS 114015	Pyrus communis	AF230767	GQ250299	Mostert et al. 2001 Santos et al. 2010	
Diaporthe angelicae	CBS 111592 AR3776	Heracleum sphondylium	AY196779	GQ250302	Santos et al. 2010	
Diaporthe aspalathi	CBS 117169	Aspalathus linearis	DQ286275	DQ286249	van Rensburg et al. 2006	
Diaporthe australafricana	STE-U 2655	Vitis vinifera	AF230744		Mostert et al. 2001 van Niekerk et al. 2005	
Diaporthe crotalariae	CBS 162.33	Crotalaria spectabilis	FJ889445	GQ250307	Santos et al. 2010	
Diaporthe helianthi	Ar3 CBS 592.81 Su 5/04 Su 20/05 Su 3/04 Su 11/04 Su 3/06 Su 12/04 Su 8/05 Su 18/06 Su 12/05	Arctium lappa Helianthus annuus	FJ841859 AY705842 FJ841854 FJ841855 FJ841856 FJ841861 FJ841863 FJ841865 FJ841865 FJ841866 FJ841866 FJ841867 FJ841867	GQ250308	Vrandecic et al. 2010 Santos et al. 2010 Vrandecic et al. 2010	
	Su 25/05 Dh95016 Dh95048 Dh95057 Dh95004 Dh95045 Dh95049		FJ841868 AF358435 AF358436 AF358437 AF358438 AF358439 AF358440		Says-Lesage et al. 2002	
	Dh95099 G23 A1-62 STE-U 5355 STE-U 5353 STE-U 5354 STE-U 5356 STE-U 5354	Luehea divaricata V. vinifera	AF358441 EU878427 AY485745 AY485746 AY485747 AY485748 AY485748		Bernardi-Wenzel et al. 2010 van Niekerk et al. 2005	
	Xa 2 Xa 3 Xa 5	Xanthium italicum	FJ841860 FJ841857 FJ841852		Vrandecic et al. 2010	
	Xa 9 Xa 12 Xa 12	Xanthium strumarium	FJ841858 FJ841853 FJ841862			
Diaporthe hickoriae	Xa 13 CBS 145.26	Xanthium sp.	FJ841862 FJ889446	GQ250309		
,	CBS 143.26 CBS 123212	Carya glabra		GQ250309 GQ250310	Santos & Phillips 2000	
Diaporthe lusitanicae	Di-C001/5	Foeniculum vulgare	EU814477	60250510	Santos & Phillips 2009 Santos et al. 2010	
Diaporthe melonis	CBS 507.78	Cucumis melo	FJ889447	GQ250314	Santos et al. 2010	
Diaporthe neotheicola	CBS123208⁴ Di-C004/5	F. vulgare	EU814480	GQ250315	Santos & Phillips 2009 Santos et al. 2010	
Diaporthe perjuncta	CBS 109745	Ulmus glabra	AY485785	GQ250323	van Niekerk et al. 2005 Santos et al. 2010	
Diaporthe stewartii	CBS 193.36	Cosmos bipinnatus	FJ889448	GQ250324	Santos et al. 2010	
Diaporthe strumella var. longispora	CBS 194.36	Ribes sp.	FJ889449	GQ250325		
Diaporthe vaccinii	CBS 160.32	Oxycoccus macrocarpus	AY952141	GQ250326		
Diaporthe viticola	CBS 113201 STE-U 5683	V. vinifera	AY485750	GQ250327	van Niekerk et al. 2005 Santos et al. 2010	
Diaporthe sp.	DAR 73811	Carthamus lanatus	EU311607	FJ389003	Ash et al. 2010	
Phomopsis amygdali	CBS 126679 11A	Prunus dulcis	GQ281791 GQ281792	GQ250339	Diogo et al. 2010 Santos et al. 2010	
Phomopsis cotoneastri	CBS 439.82	Cotoneaster sp.	FJ889450	GQ250341	Santos et al. 2010	
Phomopsis cuppatea	CBS 117499	Aspalathus linearis	AY339322	AY339354	van Rensburg et al. 2006	
Phomopsis dauci	CBS 315.49	Daucus carota	FJ889451	GQ250348	Santos et al. 2010	
Phomopsis longicolla	SSLP-1	Glycine max	HQ333500	HQ333505	unpublished	
Phomopsis phoenicicola	CBS 161.64	Areca catechu	FJ889452	GQ250349	Santos et al. 2010	
Phomopsis sclerotioides	CBS 296.67	Cucumis sativus	AF439626	GQ250350	Farr et al. 2002 Santos et al. 2010	
Phomopsis subordinaria	CBS 104.84	Plantago lanceolata	GQ922519		unpublished	
Phomopsis viticola	CBS 114016	V. vinifera	AF230751	GQ250351	Mostert et al. 2001 Santos et al. 2010	

¹ CBS: Centralbureau voor Schimmelcultures, Utrecht, The Netherlands; Ph- & Di-: culture collection housed at Centro de Recursos Microbiolo' gicos, Caparica, Portugal. ² ITS: internal transcribed spacer.

³ TEF-1α: translation elongation factor-1alpha.
 ⁴ Di-C004/5 is also recorded as CBS 123208.

⁵ Ex-type cultures are in **bold**.

Fig. 1 Phylogenetic tree resulting from the alignment of 540 characters of the ITS region. The phylogenetic tree was inferred using the Maximum Likelihood method based on the Kimura 2-parameter model. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1 000 replicates) are shown next to the branches. A discrete Gamma distribution was used to model evolutionary rate differences among sites (5 categories (+G, parameter = 0.3209)). The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. Species described in this work are highlighted. Ex-type cultures are in **bold**.

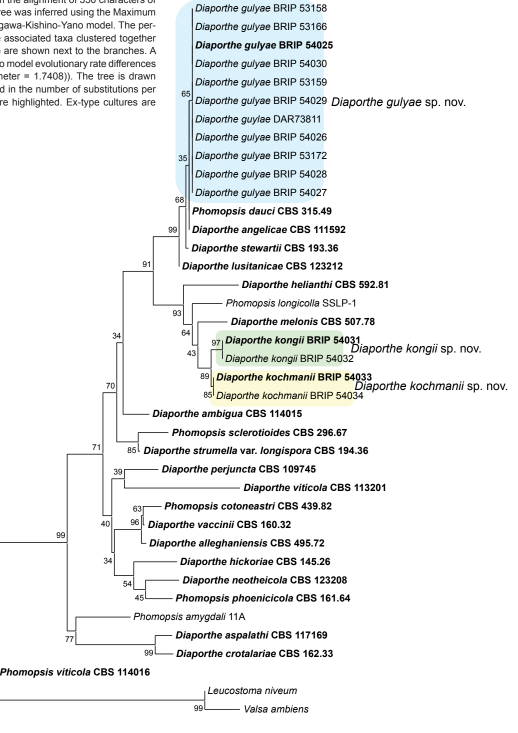




15

16

Fig. 2 Phylogenetic tree resulting from the alignment of 350 characters of the TEF-1 α region. The phylogenetic tree was inferred using the Maximum Likelihood method based on the Hasegawa-Kishino-Yano model. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1 000 replicates) are shown next to the branches. A discrete Gamma distribution was used to model evolutionary rate differences among sites (5 categories (+G, parameter = 1.7408)). The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. Species described in this work are highlighted. Ex-type cultures are in **bold**.



from Herr et al. 1983 and van Rensburg et al. 2006) required the placement of a 5 mm cube of colonised WAS into a 5–10 mm long slit made in the stem at a node. This wound was then sprayed with distilled water and wrapped with permeable film (ParafilmTM). Control plants were wounded with a 5–10 mm long slit at the nodes as for the treated plants, then wrapped with permeable film without placing an agar cube in the wound. Both inoculated and control plants were sprayed with distilled water, placed in a dew chamber and incubated at 25 °C 12 h light / 20 °C 12 h dark for 48 h then returned to a growth cabinet under the light and temperature regime described above. This test was replicated five times for each isolate.

0.1

The less invasive mycelium contact method (Miric 2002) was used as a secondary test for pathogenicity of selected isolates. A 5 mm cube of inoculated agar was placed in contact with the stem at a node, sprayed with distilled water, wrapped with permeable film and incubated as described above. Plants were assessed for lesion development at 14 d after inoculation on a scale of 1 to 5 (Table 1).

RESULTS

Phylogenetic analysis

For the ITS region, approximately 540 bases were sequenced for the isolates in this study and added to the alignment. The alignment included sequences from 58 *Diaporthe/Phomopsis* species (including two outgroups), of which 23 were from extype cultures.

For the TEF-1 α region, approximately 580 bases were sequenced for the isolates in this study. However, only 350 bases could be used to compare with the GenBank-retrieved sequences. The alignment included sequences from 24 *Diaporthe/ Phomopsis* species (including two outgroups), of which 20 were from ex-type cultures. Evolutionary relationships of these sequences were analysed using the ML method based on a K2+G model for ITS, and a HKY+G model for TEF-1 α , as determined by Modeltest in MEGA v. 5.05.

The phylogramme of the ITS region showed that the Australian isolates of *Diaporthe* from stem cankers on sunflower formed three well-supported clades, which indicate novel species (Fig. 1). One of these clades was close to ex-type strains of three species, namely *D. angelicae*, *D. stewartii* and *P. dauci*, as well as an isolate of *P. subordinaria*. Furthermore, this clade included an isolate (DAR 73811) identified by Ash et al. (2010) as *Phomopsis* sp. that was pathogenic on *Carthamus lanatus* (saffron thistle, *Asteraceae*). To improve the resolution between this clade and *D. angelicae*, *D. stewartii* and *P. dauci*, an ML analysis was conducted on the TEF-1 α dataset, which is consistent with the ITS phylogramme, but with a stronger bootstrap value (65 %) (Fig. 2).

The phylogenetic analysis of the ITS dataset included 31 isolates of *D. helianthi* sourced from five publications (Says-Lesage et al. 2002, van Niekerk et al. 2005, Bernardi-Wenzel et al. 2010, Santos et al. 2010, Vrandecic et al. 2010) and formed three distinct clades (Fig. 1). One clade included the ex-type culture of *D. helianthi* (CBS 592.81), while two other clades appeared to represent novel *Diaporthe* species (Fig. 1, *Diaporthe* sp. 1 and 2).

Pathogenicity

The 14 selected isolates inoculated onto sunflower caused a range of symptoms (Table 1), which divided them into two main groups. Ten isolates causing the most severe symptoms, rated 4 or 5 for virulence, originated from stems, seeds and leaves of infected sunflower plants from both NSW and Qld. Four isolates, causing less severe symptoms and rated 2 or 3 were collected from stems of infected plants in Queensland.

Using the wound inoculation method, tan to brown elongated lesions were evident above and below the point of inoculation

after 3–7 d for the most virulent isolates, (those rated 4 or 5) with lesions expanding rapidly upwards causing plant death after 7–14 d. Earliest symptoms at 1–3 d after inoculation for the most virulent isolates (rated 4 or 5) included brownish streaks moving upwards from the inoculation site, wilting of leaves at the node closest to the site of inoculation as well as leaves directly above the site. At times, wilting of leaves above the site of inoculation occurred without obvious stem streaking. Generally, affected leaves developed a water-soaked appearance sometimes associated with twisting.

Two to four weeks after inoculation, stem pieces above and below the site of the wound were excised from all plants with lesions, surface sterilized as previously described, and incubated on WAS at 23–25 °C for up to 3 wk. Pycnidia developed between 7–21 d. Conidia oozing from pycnidia were streaked onto PDAS and the cultures compared with those of the original isolates. Isolates were re-inoculated onto sunflower plants to confirm their pathogenicity and to complete Koch's Postulates. A comparison of wound and mycelium contact inoculation methods showed similar results for pathogenicity for individual isolates after 14 d, although wound inoculated plants displayed symptoms 1–7 d earlier than those inoculated by the mycelium contact method.

Taxonomy

Based on morphology, pathogenicity and DNA sequence analysis, three undescribed species of *Diaporthe* were recognised. Although two of the new fungi only produced an anamorphic stage, all have been described in *Diaporthe* (1870), which has priority over *Phomopsis* (1884).

Diaporthe gulyae R.G. Shivas, S.M. Thompson & A.J. Young, *sp. nov.* — MycoBank MB561569; Fig. 3

Conidiomata pycnidialia, sparsa in PDA, subglobosa, usque ad 3 mm diametro, interdum rostris ostiolatis usque ad 1 mm longis, cinctis ectostromate nigro. Conidiophora facta e strato interiore parietis locularis, interdum ramosa et septata, subhyalina, usque ad 6 µm diametro. Cellulae conidiogenae cylindraceae, hyalinae, 7–18 × 1.5–2.5 µm. Alpha conidia globosa, subglobosa, ellipsoidea, ovalia vel obovoidea, hyalina, (6–)6.5–9.0(–10) × 2.5–3.5 µm. Beta conidia haud conspecta.

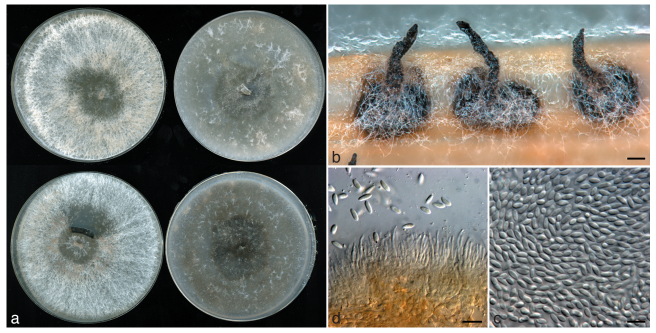


Fig. 3 Diaporthe gulyae (ex-type BRIP 54025). a. Cultures on PDA (left), OA (right) after 7 d (top) and 28 d (bottom); b. pycnidial beaks on sterilised wheat straw; c. alpha conidia; d. conidia and conidiophores. — Scale bars: b = 100 µm; c, d = 10 µm.

Etymology. In recognition of Dr Tom Gulya for his outstanding contributions to sunflower pathology research and enduring mentoring roles in the USA, Europe and Australia.

Conidiomata pycnidial, scattered on PDA, subglobose, up to 3 mm diam, occasionally with ostiolate beaks up to 1 mm long, surrounded by a black ectostroma. Conidiophores formed from the inner layer of the locular wall, sometimes branched and septate, subhyaline, up to 6 μ m diam, Conidiogenous cells cylindrical, hyaline, 7–18×1.5–2.5 μ m. Alpha conidia globose, subglobose, ellipsoidal, oval or obovoid, hyaline, (6–)6.5–9.0 (–10) × 2.5–3.5 μ m. Beta conidia not seen.

Culture characteristics — Colonies on PDA covering entire plate after 10 d, buff, ropey near the margin and adpressed in the centre, scant aerial mycelium, reverse buff with a slightly darker centre; on OA covering the entire plate after 10 d, adpressed with scattered tufts of greyish mycelium, greyish sepia, with a fuscous black central zone 3 cm diam, reverse greyish sepia with a fuscous black central zone.

Specimens examined. AustRALIA, Queensland, Ryeford near Clifton, on *Helianthus annuus* hybrid Sunbird 7, 29 Nov. 2010, *S.M. Thompson* (holotype BRIP 54025, includes ex-type culture); Ryeford near Clifton, on *Helianthus annuus* hybrid Sunbird 7, 29 Nov. 2010, *S.M. Thompson*, paratypes BRIP 54026, 54027.

Notes — Based on molecular phylogenetic inference, *D. gul*yae was placed near to the ex-type specimens of *D. angelicae*, *D. stewartii* and *P. dauci*, as well as a strain of *P. subordinaria* (Fig. 1, 2). Morphologically there is little difference between these species but unique fixed nucleotides accurately differentiate *D. gulyae*. *Diaporthe gulyae* differs from *D. stewartii* in two loci: ITS position 24 (T) and 98 (A); TEF-1 α position 19 (A), 324 (T), 30 (T), 46 (T), 47 (A) and 315 (T). *Diaporthe gulyae* differs from *D. angelicae* and *P. dauci* in two loci: ITS position 59 (C), 90 (T), 136 (A), 158 (A) and 457 (A); TEF-1 α position 30 (T) and 47 (A).

Diaporthe gulyae causes a severe stem canker on sunflower and saffron thistle. On the basis of pathology and substrate preference *D. gulyae* differs from *D. angelicae*, which is found on the decaying stems of hosts in the *Apiaceae* (Castlebury et al. 2003); *D. stewartii*, which causes stem blight of *Cosmos bipinnatus* (*Asteraceae*) (Harrison 1935); *P. dauci*, which causes inflorescence blight of *Daucus carota* (carrot, *Umbelliferae*) (von Arx 1951); and *D. adunca* (*P. subordinaria*), which attacks the scapes of *Plantago lanceolata* (*Plantaginaceae*) (Meijer et al. 1994).

Diaporthe kongii R.G. Shivas, S.M. Thompson & A.J. Young, *sp. nov.* — MycoBank MB561570; Fig. 4a, c, e

Conidiamata pycnidialia, sparsa in PDA, subglobosa, usque ad 2 mm diametro, rostris ostiolatis levibus ad apicem et saepe tectis hyphis brevibus inramosis usque ad 200 µm, cinctis ectostromate nigro. Conidiophora facta e strato interiore parietis locularis, polyangularia, interdum ramosa et septata, subhyalina ad brunneola olivacea, usque ad 6 µm diametro. Cellulae conidiogenae cylindraceae ad obclavatas, hyalinae, $6-12 \times 1.5-4$ µm. Alpha conidia ovalia ad cylindracea, biguttulata, hyalina, $5.5-7(-7.5) \times 2-2.5(-3)$ µm. Beta conidia sigmoidea vel lunata, plerumque curvata per 90–180°, hyalina, $13-23 \times 1-1.5$ µm.

Etymology. In recognition of Dr Gary Kong for his innovative contributions to sunflower pathology in Australia, specifically his investigation of the genetics of resistance to *Puccinia helianthi* and *Alternaria helianthi*.

Conidiomata pycnidial, scattered on PDA, subglobose, up to 2 mm diam, with short (less than 0.5 mm) ostiolate beaks smooth towards apex and often covered with short unbranched hyphae up to 200 μ m, surrounded by a black ectostroma. Conidiophores formed from the inner layer of the locular wall, polyangular, sometimes branched and septate, subhyaline to pale olivaceous brown, up to 6 μ m diam. Conidiogenous cells cylindrical

to obclavate, hyaline, $6-12 \times 1.5-4 \mu m$. Alpha conidia oval to cylindrical, biguttulate, hyaline, $5.5-7(-7.5) \times 2-2.5(-3) \mu m$. Beta conidia sigmoid to lunate, mostly curved through 90–180°, hyaline, $13-23 \times 1-1.5 \mu m$.

Culture characteristics — Colonies on PDA covering entire plate after 10 d, ropey with a conspicuous ring 2.5 cm in diam of tufted aerial mycelium and abundant tufts towards the margin, white to greyish white with scattered amber patches, with several scattered minute black stroma, reverse with an isabelline ring, paler towards the margin; on OA covering the entire plate after 10 d, adpressed, rosy-buff, with an irregular grey olivaceous central zone about 4.5 cm diam and smaller irregular grey olivaceous patches towards the margin containing a few minute black stroma, the central zone and patches have yellowish margins, reverse rosy buff with irregular isabelline patches.

Specimens examined. AUSTRALIA, Queensland, Childers, on Helianthus annuus hybrid PDAS, 1 Dec. 2010, S.M. Thompson (holotype BRIP 54031, includes ex-type culture); Childers, on Helianthus annuus hybrid PDAS, 1 Dec. 2010, S.M. Thompson, paratype BRIP 54032.

Notes — Based on phylogenetic inference from the ITS sequence data (Fig. 1), *D. kongii* is closely related to *P. cuppatea*, which was isolated from plants of *Aspalanthus linearis* (rooibos, *Fabaceae*) with die-back (van Rensburg et al. 2006). Morphologically *D. kongii* has smaller conidia than those of *P. cuppatea*, which measure $(10-)12-13(-14) \mu m$.

Diaporthe kochmanii R.G. Shivas, S.M. Thompson & A.J. Young, *sp. nov.* — MycoBank MB561571; Fig. 4b, d, f-h

Perithecia formata in PDA et in caulibus sterilifactis apricifloris post octo hebdomades, subglobosa, usque ad 350 µm diametro, plerumque solitaria in agaro vel aggregata in fasciculis in caulibus, cincta ectostromate nigro, uno vel pluribus collis cylindraceis nigris ostiolatis usque ad 2 mm haud distinctis ab eis in pycnidiis. Asci unitunicati, cylindracei, $33-41 \times 5-7 \ \mu m$, hyalini, octospori, biseriati, annulo conspicuo refractivo apicali. Ascosporae hyalinae, mediane septatae, ovales ad cylindraceas, haud constrictae ad septum, guttula in quaque cellula, 9–10 \times 2.5–3.5 $\mu\text{m},$ leves. Conidiomata pycnidialia, sparsa in PDA, nigra, subglobosa, usque ad 2 mm diametro, uno vel pluribus collis cylindraceis nigris ostiolatis usque ad 2 mm. Conidiophora facta e strato interiore parietis locularis, polyangularia, interdum ramosa et septata, subhyalina ad brunneola olivacea, usque ad 6 µm diametro. Cellulae conidiogenae cylindraceae ad obclavatas, hyalinae, $5-10 \times 1.5-3$ μ m. Alpha conidia ovalia ad cylindracea, (5–)5.5–7(–7.5) \times 2–3 μ m. Beta conidia flexuosa ad lunata, plerumque curvata per 45-90°, hyalina, 11-17 \times 1–1.5 µm.

Etymology. In recognition of Dr Joe Kochman who pioneered the investigation of rust races on sunflower in Australia and his widely recognised contributions to sunflower pathology.

Perithecia formed on PDA and on sterilised stems of sunflower after 8 wk, subglobose, up to 350 µm diam, usually solitary in the agar or aggregated in clusters on the stems, surrounded by a black ectostroma, with 1 or more cylindrical, black, ostiolate necks up to 2 mm, indistinguishable from those on pycnidia. Asci unitunicate, cylindrical, $33-41 \times 5-7$ (av. = $37 \times 6 \mu$ m), hyaline, 8-spored, biseriate, with conspicuous refractive apical ring. Ascospores hyaline, medially septate, oval to cylindrical, not constricted at the septum, with a guttule in each cell, $9-10 \times$ $2.5-3.5 \,\mu\text{m}$ (av. = $9.5 \times 3 \,\mu\text{m}$), smooth. Conidiomata pycnidial, scattered on PDA, black, subglobose, up to 2 mm diam, with 1 or more cylindrical black ostiolate necks up to 2 mm long. Conidiophores formed from the inner layer of the locular wall, polyangular, sometimes branched and septate, subhyaline to pale olivaceous brown, up to 6 µm diam. Conidiogenous cells cylindrical to obclavate, hyaline, 5–10 μ m × 1.5–3 μ m. Alpha conidia oval to cylindrical, $(5-)5.5-7(-7.5) \times 2-3 \mu m$. Beta conidia flexuous to lunate mostly curved through 45-90°, hyaline, $11-17 \times 1-1.5 \mu m$.

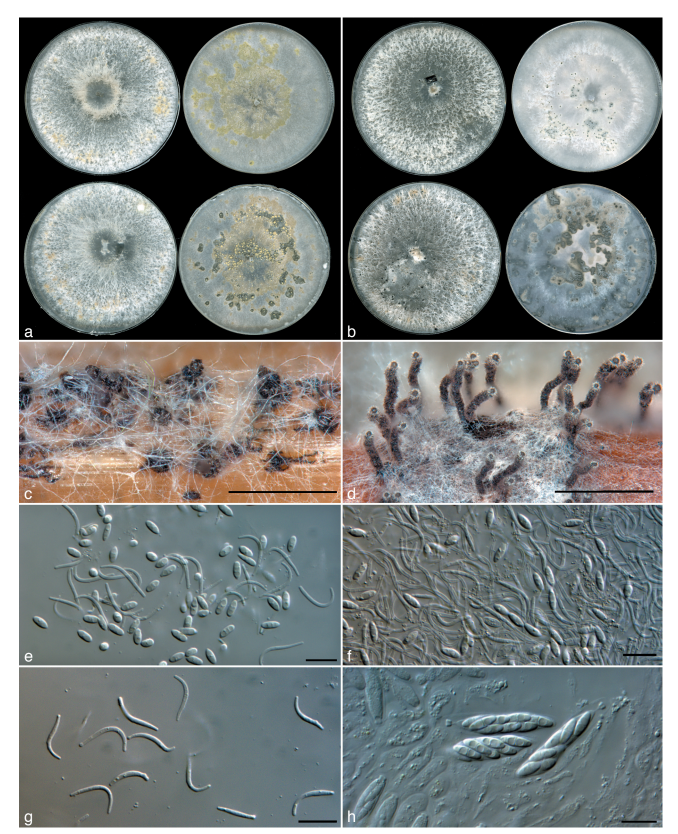


Fig. 4 Diaporthe kongii (ex-type BRIP 54031) and *D. kochmanii* (ex-type BRIP 54033). a. Diaporthe kongii cultures on PDA (left), OA (right) after 7 d (top) and 28 d (bottom); b. Diaporthe kochmanii cultures on PDA (left), OA (right) after 7 d (top) and 28 d (bottom); c. pycnidial beaks of *D. kongii* on sterilised wheat straw; d. perithecial necks of *D. kochmanii* on sterilised wheat straw; e. alpha and beta conidia of *D. kongii*; f. alpha and beta conidia of *D. kochmanii*; g. beta conidia of *D. kochmanii*; h. asci and ascospores of *D. kochmanii*. — Scale bars: c, d = 1 mm; e-h = 10 µm.

Culture characteristics — Colonies on PDA covering entire plate after 10 d, ropey with abundant tufts of mycelium, pale mouse grey, lighter towards the margin, with abundant scattered minute black stroma, reverse smoke grey with a darker central zone 5 cm diam; on OA covering the entire plate after 10 d, adpressed with scant tufted aerial mycelium, pale rosy vinaceous, with irregular pale olivaceous grey patches up to 1 cm wide containing minute black stroma, reverse pale rosy vinaceous with pale greyish areas where stroma form.

Specimens examined. AustRALIA, Queensland, Lawes, on *Helianthus annuus* Experimental Line, 25 Nov. 2010, *S.M. Thompson* (holotype BRIP 54033, includes ex-type culture); Lawes, on *Helianthus annuus* hybrid PDAS, 25 Nov. 2010, *S.M. Thompson*, paratype BRIP 54034. Notes — Based on phylogenetic inference of the TEF-1 α sequence data *D. kochmanii* is closest to *D. kongii*. Morphologically these two species cannot be reliably separated. *Diaporthe kochmanii* differs from *D. kongii* in the TEF-1 α locus position 60 (A), 83 (C), 184 (C), 219 (C), 240 (T), 260 (G), 266 (C), 268 (T), 280 (G), 284 (T), and 288 (T).

DISCUSSION

In this study, pathogenic *Diaporthe* species have been identified from wild, inbred and hybrid sunflowers grown throughout NSW and Qld. We have demonstrated that there are at least three previously unrecognised and novel species, namely *D. gulyae*, *D. kongii* and *D. kochmanii*, associated with stem cankers on sunflower in Australia. The most virulent of these species, *D. gulyae*, also contained an isolate identified by Ash et al. (2010) as pathogenic to saffron thistle. Symptoms caused by *D. gulyae* on sunflower closely resembled those of *D. helianthi*.

Unfavourable dry environmental conditions and low pathogen populations may explain the previous low frequency of sunflower stem cankers attributed to *Diaporthe* species in Australia. It is possible that severe outbreaks in Australia will remain sporadic, as has been found in Italy, despite climatic conditions appearing to be conducive to the disease (Battilani et al. 2003, Vergara et al. 2004). We consider it likely that outbreaks caused by these new *Diaporthe* species will become more widespread in the current cycles of wet summer weather, especially with the tendency towards minimum tillage practices that appear to increase pathogen inoculum in unprocessed stubble.

The molecular phylogenetic analysis showed that authentic *D. helianthi* derived from an ex-type isolate, clustered in a clade with isolates from the former Yugoslavia and France (Fig. 1). *Diaporthe helianthi* has also been recorded from hosts other than sunflower in Croatia (Vrandecic et al. 2010), which was part of the former Yugoslavia. All records of *D. helianthi* from hosts other than sunflower, without comparison to sequence data from ex-type cultures, should be treated with caution (e.g. van Niekerk et al. 2005, Bernardi-Wenzel et al. 2010). Unintentional misapplications of the name *D. helianthi* have resulted from the absence and inaccessibility of cultures derived from type material, which are needed for molecular comparison.

Based on the localities of previous *Diaporthe* collections in Australia from sunflower, soybean (*Glycine max*), Noogoora burr (*Xanthium pungens*) (Miric 2002), saffron thistle (Ash et al. 2010) plus herbarium records, we expect that future surveys will broaden the host and distribution ranges of these newly described species. We also anticipate that more species associated with stem cankers on sunflower in Australia will be identified.

The results of our study highlight the need for the re-evaluation of the identification and classification of *Diaporthe (Phomopsis*) species (Farr et al. 2002, Hyde et al. 2010, Santos et al. 2010, Udayanga et al. 2011). Accurate and reliable methods of identification for *Diaporthe* species is a major concern for biosecurity agencies in many countries, including Australia. In this regard, *D. helianthi* has not been identified from sunflower in Australia and remains a biosecurity threat.

Advances in molecular identification techniques are helping to further define species boundaries by providing more specific genetic evidence in support of taxonomic differences (Udayanga et al. 2011). The combination of pathology (host range and pathogenicity), taxonomic descriptions and molecular analyses will certainly result in the identification and description of more *Diaporthe* species from a range of host plants worldwide. Acknowledgements We thank Dr Tom Gulya, USDA-ARS Northern Crops Laboratory ND, USA, and Dr Malcolm Ryley (Department of Employment, Economic Development and Innovation QLD, Australia) for their pathology advice, Loretta Serafin (NSW Department of Primary Industries, NSW, Australia) for agronomic expertise, the Australian Oilseeds Federation (AOF), the Australian Sunflower Association (ASA), Pacific Seeds, Nuseed and HSR Seeds and Grains Research Development Corporation (GRDC) for financial support. We also acknowledge the support of the University of Queensland. Don Barrett (University of Queensland) is thanked for translating the species diagnoses into Latin.

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