Phylogeny and morphology of four new species of Lasiodiplodia from Iran

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Key words
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EF-1α
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Abstract
Four new species of Lasiodiplodia, L. citicola, L. gilanensis, L. hormozganensis and L. iranensis from various tree species in Iran are described and illustrated. The ITS and partial translation elongation factor-1α sequence data were analysed to investigate their phylogenetic relationships with other closely related species and genera. The four new species formed well-supported clades within Lasiodiplodia and were morphologically distinct from all other known species.

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INTRODUCTION
Members of the Botryosphaeriaceae (Botryosphaeriales, Dothideomycetes, Ascomycota) are cosmopolitan and occur on a wide range of monocotyledonous, dicotyledonous and gymnosperm hosts (von Arx & Müller 1954, Barr 1987). They are associated with various symptoms such as shoot blights, stem cankers, fruit rots, dieback and gummosis (von Arx 1987) and are also known as endophytes (Slippers & Wingfield 2007). Based on 28S rDNA sequence data Crous et al. (2006) showed that Botryosphaeria is polyphyletic and they divided it into several genera distinguishable by conidial morphology and phylogenetic data. Botryosphaeria was thus restricted to species with Fusiccoccom anamorphs. However, the clade containing Diplodia/Lasiodiplodia could not be fully resolved. In a multigene genealogy Phillips et al. (2008) resolved and separated this clade into six genera including Diplodia, Lasiodiplodia, Neodeeitonia, Barriopsis, Phaeobotryon and Phaeobotryosphaeria. Morphological characters of the anamorphic and teleomorphic states also supported the separation of these genera.

Lasiodiplodia species are common, especially in tropical and subtropical regions where they cause a variety of diseases (Punithalingam 1980). According to Sutton (1980) the genus is based on Lasiodiplodia theobromae. The main features that distinguish this genus from other closely related genera are the presence of pycnidal paraphyses and longitudinal striations on mature conidia. Thus far 20 species have been described and they are differentiated on the basis of conidial and paraphyses morphology. The more recently described species (described since 2004) have been separated not only on morphology, but also on the basis of ITS and EF-1α sequence data. Punithalingam (1976) included several of the species known at that time as synonyms of L. theobromae since he could not separate them on morphological characters. However, on account of its morphological variability and wide host range it seems likely that L. theobromae is a species complex. Recent studies based on sequence data have confirmed this and eight new species have been described since 2004 (Pavlic et al. 2004, 2008, Burgess et al. 2006, Damm et al. 2007, Alves et al. 2008).

There have been no studies on the Lasiodiplodia species in Iran apart from a few reports of L. theobromae. In a survey of Botryosphaeriaceae in Iran some Lasiodiplodia isolates that differed from L. theobromae in terms of morphology and ISSR fingerprinting profile were found. The aim of this study was to characterise these isolates in terms of anamorph morphology and phylogenetic analysis.

MATERIALS AND METHODS

Fungal isolation
During a survey of Botryosphaeriaceae in different regions of Iran in 2005–2007 some 30 Lasiodiplodia-like isolates were collected from various tree species showing symptoms of branch dieback, cankers and fruit rot. Isolations were made from single conidia or by directly plating out pieces of diseased tissue after surface sterilization (1–4 min in 70 % ethanol). Representative isolates were deposited in the culture collection of the Iranian Research Institute of Plant Protection (IRAN, Tehran, Iran) and the Centraalbureau voor Schimmelcultures (CBS, Utrecht, The Netherlands). Isolates included in the morphological and phylogenetic analyses are listed in Table 1.

Morphology and culture characteristics
To induce sporulation, isolates were transferred to 2 % water agar with sterilised pine needles on the agar surface and incubated under mixed near-UV and cool-white fluorescent light in a 12 h light-dark regime for 2–6 wk at 25 °C. Vertical sections through conidiomata were made for some isolates with a Leica CM1100 cryostat microtome. Structures were mounted in 100 % lactic acid and digital images were recorded with a Leica DFC 320 camera on a Leica DMR HC microscope. Measurements were made with the Leica IM500 measurement module. From measurements of 50 conidia the mean, standard deviation

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### Table 1
Isolates included in the phylogenetic study.

| Species | Culture no. | Substrate | Locality | Collector | ITS | GenBank |
|---------|-------------|-----------|----------|-----------|-----|---------|
| *Botryosphaeria tsugae* | CBS 418.64 | Tsuga heterophylla | Canada | A. Funk | DQ458898 | DQ458897 |
| *Diplodia corticola* | CBS 112549 | Quercus suber | Portugal | A. Alves | AR259100 | AY573227 |
| *D. cupressi* | CBS 261.85 | Cupressus sempervirens | Bet Dagan, Israel | Z. Solel | DQ458894 | DQ458879 |
| *D. muta* | CBS 112553 | Vitis vinifera | Portugal | A.J.L. Phillips | AR259093 | AY573220 |
| *D. pinea* | CBS 230.30 | Pinus nigra | Netherlands | H.A. van der Aa | DQ458895 | DQ458892 |
| *D. rosulata* | CBS 112546 | Pinus radiata | South Africa | M.J. Wingfield | DQ458896 | DQ458881 |
| *D. scrobiculata* | CBS 112555 | Pinus africana | Ethiopia | A. Gure | AR210345 | EU430268 |
| *D. sericata* | CBS 119049 | Pinus africana | Ethiopia | A. Gure | AR210345 | EU430268 |
| *Lasiodiplodia* | CBS 15038 | Malus pumila | Netherlands | A.J.L. Phillips | AR573206 | AY573223 |
| *Lasiodiplodia cincta* | IRAN 1521C | Citrus sp. | Iran | A. Shekarli | GU945353 | GU945339 |
| *Lasiodiplodia cincta* | IRAN 1523C | Citrus sp. | Iran | J. Abdollahzadeh/A. Javadi | GU945354 | GU945340 |
| *Lasiodiplodia cincta* | CMV 13488 | Eucalyptus urophylla | Venezuela | S. Mohali | DQ103552 | DQ103559 |
| *Lasiodiplodia cincta* | CBS 118741 | Santalum album | Australia | T.I. Burgess/B. Dell | DQ103550 | DQ103557 |
| *Lasiodiplodia cincta* | IRAN 1501C | Unknown | Iran | J. Abdollahzadeh/A. Javadi | GU945352 | GU945341 |
| *Lasiodiplodia cincta* | CBS 118512 | Syzygium cordatum | South Africa | D. Pavlic | DQ458892 | DQ458877 |
| *Lasiodiplodia cincta* | CBS 116355 | Syzygium cordatum | South Africa | D. Pavlic | AR639594 | QD103567 |
| *Lasiodiplodia cincta* | IRAN 15444 | Mangifera indica | Iran | J. Abdollahzadeh/A. Javadi | GU945356 | GU945344 |
| *Lasiodiplodia cincta* | IRAN 1500C | Olea sp. | Iran | J. Abdollahzadeh/A. Javadi | GU945355 | GU945343 |
| *Lasiodiplodia cincta* | IRAN 921C | Mangifera indica | Iran | N. Khezrinejad | GU945346 | GU945334 |
| *Lasiodiplodia cincta* | IRAN 15035 | Juglans regia | Iran | A. Javadi | GU945347 | GU945335 |
| *Lasiodiplodia cincta* | IRAN 1517C | Juglans regia | Iran | J. Abdollahzadeh/A. Javadi | GU945349 | GU945337 |
| *Lasiodiplodia cincta* | IRAN 1519C | Mangifera indica | Iran | J. Abdollahzadeh/A. Javadi | GU945350 | GU945338 |
| *Lasiodiplodia cincta* | IRAN 1520C | Selvadaria pericha | Iran | J. Abdollahzadeh/A. Javadi | GU945348 | GU945336 |
| *Lasiodiplodia cincta* | CBS 122519 | Adansonia gibbosa | Western Australia | T.I. Burgess | EU144050 | EU144065 |
| *Lasiodiplodia cincta* | CBS 122065 | Adansonia gibbosa | Western Australia | T.I. Burgess | EU144050 | EU144066 |
| *Lasiodiplodia cincta* | CBS 494.78 | Cassava-field soil | Colombia | O. Rangel | EF622094 | EF622064 |
| *Lasiodiplodia cincta* | CBS 456.78 | Cassava-field soil | Colombia | O. Rangel | EF622093 | EF622063 |
| *Lasiodiplodia cincta* | CBS 121103 | Vitis vinifera | South Africa | F. Hallesen | AR343842 | EF45396 |
| *Lasiodiplodia cincta* | CBS 120832 | Prunus salicina | South Africa | U. Damm | EF445362 | EF445395 |
| *Lasiodiplodia cincta* | CBS 116499 | Gmelina arborea | Costa Rica | J. Carranza-Velasquez | EF622077 | EF622057 |
| *Lasiodiplodia cincta* | CBS 374.54 | Coffea sp. | Zaire | Unknown | EF622090 | EF622059 |
| *Lasiodiplodia cincta* | IRAN 1518C | Citrus sp. | Iran | J. Abdollahzadeh/A. Javadi | GU973874 | GU973866 |
| *Lasiodiplodia cincta* | CJA36 | Citrus sp. | Iran | J. Abdollahzadeh/A. Javadi | GU973875 | GU973867 |
| *Lasiodiplodia cincta* | CBS 164.96 | Fruit on coral reef coast | New Guinea | A. Apte | AR641026 | AY641058 |
| *Lasiodiplodia cincta* | CBS 1125C | Unknown | Unknown | Unknown | AR622074 | AR622064 |
| *Lasiodiplodia cincta* | IRAN 1233C | Unknown | Iran | J. Abdollahzadeh/A. Javadi | GU973868 | GU973860 |
| *Lasiodiplodia cincta* | IRAN 1496C | Mangifera indica | Iran | J. Abdollahzadeh/A. Javadi | GU973869 | GU973861 |
| *Lasiodiplodia cincta* | IRAN 1496C | Mangifera indica | Iran | J. Abdollahzadeh/A. Javadi | GU973870 | GU973862 |
| *Lasiodiplodia cincta* | CJA198 | Unknown | Iran | Unknown | GU973871 | GU973863 |
| *Lasiodiplodia cincta* | CJA199 | Unknown | Iran | Unknown | GU973872 | GU973864 |
| *Lasiodiplodia cincta* | CJA279 | Coccos sp. | Unknown | J. Abdollahzadeh | GU973873 | GU973865 |
| *Lasiodiplodia cincta* | WAC 12539 | Acacia mangium | Venezuela | S. Mohali | DO120348 | DO120368 |
| *Lasiodiplodia cincta* | WAC 12540 | Acacia mangium | Venezuela | S. Mohali | DO120348 | DO120369 |
| *Spencermartinsia* | CBS 117006 | Vitis vinifera | Spain | J. Lurque & S. Martos | AR905555 | AY905562 |
Fig. 1 One of six most parsimonious trees obtained from combined ITS and EF-1α sequence data. MP and NJ bootstrap values are given based on 1000 pseudoreplicates above and below the branches respectively. The tree is rooted to Dothiorella sarmentorum (CBS115038) and Spencermartinsia sp. (CBS117006).

and 95% confidence intervals were calculated. Dimensions are given as the range of measurements with extremes in parentheses followed by 95% confidence limits and mean ± standard deviation. Dimensions of other structures are given as the range of at least 20 measurements. Colony morphology, colour (Rayner 1970), and growth rates between 5 and 35 °C in 5 °C intervals, were determined on 2% malt extract agar (MEA, Difco laboratories) in the dark. Nomenclatural novelties and descriptions were deposited in MycoBank (www.MycoBank.org; Crous et al. 2004).

Phylogenetic analysis
Isolates were grown in 2% malt extract broth (MEB) incubated at room temperature for 4–7 d. Mycelial mats were collected by filtration and washed with sterile distilled water and freeze dried with an Edward MicroModulyo 1.5K System (England) freeze drier. DNA was extracted using the method of Raeder & Broda (1985) with modifications as described by Abdollahzadeh et al. (2009). PCR reaction mixtures were prepared according to Alves et al. (2004), with the addition of 5% DMSO to improve the amplification of some difficult DNA templates. The ITS1-5.8S-ITS2 plus D1/D2 domain of the 28S rDNA gene, and the translation elongation factor-1alpha (EF-1α) were amplified with the primer pairs ITS1 (White et al. 1990) /NL4 (O’Donnell 1993) and EF1-688F/EF1-1251R (Alves et al. 2008), respectively. PCR conditions, purification and sequencing were as described in Abdollahzadeh et al. (2009). The nucleotide sequences were read and edited with Bioedit Sequence Alignment Editor v7.0.9.0 (© 1997–2007, Tom Hall). Sequences of both DNA regions of additional isolates were retrieved from GenBank (Table 1).
Phylogenetic analysis

The nucleotide sequences were aligned with ClustalX v1.83 (Thompson et al. 1997) and manually adjusted when necessary. Phylogenetic information contained in indels (insertions/deletions) was incorporated into the phylogenetic analyses using simple indel coding as implemented by GapCoder (Young & Healy 2003). Trees were rooted to Dothiorella sarmentorum and Spencermartinsia sp. Phylogenetic analyses were performed using PAUP v4.0b10 (Swofford 2003) for neighbour-joining (NJ) and maximum-parsimony (MP) analyses. The neighbour-joining analysis was performed using Kimura-2-parameter nucleotide substitution model (Kimura 1980). All characters were unordered and of equal weight. Bootstrap values were obtained from 1000 NJ bootstrap replicates. Maximum-parsimony analysis was performed using the heuristic search option with 100 random taxon additions and tree bisection and reconnection (TBR) as the branch-swapping algorithm. All characters were unordered and of equal weight and gaps were treated as missing data. Branches of zero length were collapsed and all multiple, equally parsimonious trees were collapsed and all multiple, equally parsimonious trees were saved. The robustness of the most parsimonious trees was evaluated by 1000 bootstrap replicates (Hillis & Bull 1993). Other measures used were consistency index (CI), retention index (RI) and homoplasy index (HI). A partition homogeneity test was done to determine the possibility of combining the ITS and EF-1α datasets (Farris et al. 1995, Hulsenbeck et al. 1996). New sequences were deposited in GenBank (Table 1) and the alignment in TreeBASE (S10302).

RESULTS

Phylogenetic analysis

The partition homogeneity test in PAUP was not significant (P = 0.08) indicating that the ITS (566 characters) and EF-1α (330 characters) datasets were congruent. Therefore the two datasets were combined in a single analysis. ITS and EF-1α sequences for the 20 isolates studied were combined and aligned with 37 sequences of 19 taxa, including the outgroup, retrieved from GenBank. Incomplete portions at the ends of the sequences were excluded from the analyses. The combined dataset after alignment contained 987 characters including alignment gaps, of which 74 were excluded, 552 were constant, 62 were variable and parsimony-uninformative and 299 were parsimony-informative. A heuristic search of the remaining 299 parsimony-informative characters resulted in six most parsimoni-
dark slate blue (39'"k) at the reverse after 2 wk in the dark at 25 °C. Colonies reaching 85 mm on MEA after 2 d in the dark at 25 °C. Cardinal temperatures for growth; min ≤ 10 °C, max ≥ 35 °C, opt 25–30 °C.

Substrate — Citrus sp.
Distribution — Chaboksar (Gilan Province), Sari (Mazandaran Province), Northern Iran.

Specimens examined. Iran, Gilan Province, Chaboksar, on twigs of Citrus sp., June 2007, J. Abdollahzadeh and A. Javadi, holotype IRAN 14270F, culture ex-type IRAN 1522C = CBS 124707; Mazandaran Province, Sari, on twigs of Citrus sp., June 2007, A. Shekari, IRAN 1521C = CBS 124706.

Notes — Phylogenetically Lasiodiplodia citricola is closely related to L. parva, but conidia of L. citricola, (20–)22–27(–31) × (10.9–)12–17(–19) μm, are longer and wider than those of L. parva, (15.5–)16–23.5(–24.5) × (10–)10.5–13(–14.5) μm. This species produces a pink pigment in PDA cultures at 35 °C.

Lasiodiplodia gilanensis Abdollahzadeh, Javadi & A.J.L. Phillips, sp. nov. — MycoBank MB516778; Fig. 3

Teleomorph. Unknown.

Lasiodiplodia plurivora similis sed paraphyses brevioribus et angustioribus.

Etymology. Named after Gilan Province in Iran where it was first found.

Conidiomata stromatic, pycnidial, produced on pine needles on WA within 2–4 wk, superficial, dark brown to black, covered with dense mycelium, mostly uniloculate, up to 940 μm, solitary, globose, thick-walled, non-papillate with a central ostiole. Paraphyses hyaline, cylindrical, thin-walled, initially aseptate, becoming up to 1–3 septate when mature, rarely branched, rounded at apex, up to 95 μm long, 2–4 μm wide. Conidiophores absent. Conidiogenous cells holoblastic, discrete, hyaline, smooth, thin-walled, cylindrical, 11–18 × 3–5 μm. Conidia initially hyaline, aseptate, ellipsoid to ovoid, with granular content, rounded at apex, base mostly truncate, wall < 2 μm, becoming pigmented, verruculose, ellipsoid to ovoid, 1-septate with longitudinal striations, (25.2–)28–35(–38.8) × (14.4–)15–18(–19) μm, 95 % confidence limits = 30.6–31.4 ×
16.5–16.7 μm (av. ± S.D. = 31 ± 2.4 × 16.6 ± 1 μm, l/w ratio = 1.9 ± 0.2).

Culture characteristics — Colonies with abundant aerial mycelia reaching to the lid of Petri plate, aerial mycelia becoming smoke-grey (21''''f) to olivaceous-grey (21'''''i) at the surface and greenish grey (33''''i) to dark slate blue (39''''k) at the reverse after 2 wk in the dark at 25 °C. Colonies reaching 80 mm on MEA after 2 d in the dark at 25 °C. Cardinal temperatures for growth; min ≤ 10 °C, max ≥ 35 °C, opt 25–30 °C.

Substrate — Unknown.

Distribution — Rahimabad-Garmabdost (Gilan Province), Northern Iran.

Specimens examined. Iran, Gilan Province, Rahimabad-Garmabdost, on twigs of unknown woody plant, June 2007, J. Abdollahzadeh and A. Javadi, holotype IRAN 14272F, culture ex-type IRAN 1523C = CBS 124704; Gilan Province, Rahimabad-Garmabdost, on twigs of unknown woody plant, June 2007, J. Abdollahzadeh and A. Javadi, IRAN 1501C = CBS 124705.

Notes — Phylogenetically L. gilanensis is closely related to L. plurivora, but can be distinguished on average conidial dimensions. Moreover, the paraphyses of L. gilanensis are up to 95 μm long and 4 μm wide, whereas paraphyses of L. plurivora are up to 130 μm long and 10 μm wide (Damm et al. 2007). Also, the 1–3 basal cells of L. plurivora paraphyses are often broader than the apical cells whereas, in L. gilanensis, they are the same as the apical cells. This species produces a pink pigment in PDA cultures at 35 °C.

Lasiodiplodia hormozganensis Abdollahzadeh, Zare & A.J.L. Phillips, sp. nov. — MycoBank MB516779; Fig. 4

Teleomorph. Unknown.

Lasiodiplodia citricola et L. parva phylogenetically similar. Differs from L. parva conidia majoribus (20.2 ± 1.9 × 11.5 ± 0.8 μm) et L. citricola minoribus (24.5 ± 0.2 × 15.4 ± 1.8 μm), et paraphyses minoribus.

Etymology. Named after Hormozgan Province in Iran where it was first found.

Conidiomata stromatic, pycnidial, produced on pine needles on WA within 2–4 wk, superficial, dark-brown to black, covered

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Fig. 3 Lasiodiplodia gilanensis holotype. a. Conidiomata on pine needles in culture; b–d. conidia developing on conidiogenous cells between paraphyses; e. paraphyses; f. g. hyaline, immature conidia; h, i. mature conidia in two different focal planes to show the longitudinal striations. — Scale bars: a = 1 000 μm; b, c = 5 μm; d–i = 10 μm.
with dense mycelium, mostly uniloculate, up to 950 μm, solitary, globose, thick-walled, non-papillate with a central ostiole. Paraphyses, hyaline, cylindrical, thin-walled, initially aseptate, becoming up to 1–7 septate when mature, rarely branched, occasionally basal, middle or apical cells swollen, rounded at apex, up to 83 μm long, 2–4 μm wide. Conidiogenous cells holoblastic, discrete, hyaline, smooth, thin-walled, cylindrical, 9–15 × 3–5 μm. Conidia initially hyaline, aseptate, ellipsoid to cylindrical, with granular contents, rounded at apex, base round or truncate, wall < 2 μm, becoming pigmented, verruculose, ellipsoid to ovoid, 1-septate with longitudinal striations, (15.3–)18–24(–25.2) × 11–14 μm, 95% confidence limits = 21.2–21.7 × 12.4–12.6 μm (av. ± S.D. = 21.5 ± 1.9 × 12.5 ± 0.8 μm, l/w ratio = 1.7 ± 0.2).

Culture characteristics — Colonies with abundant aerial mycelium reaching to the lid of Petri plate, aerial mycelium becoming smoke-grey (33"''i) to olivaceous-grey (31"''i) at the reverse after 2 wk in the dark at 25 °C. Colonies reaching 83 mm on MEA after 2 d in the dark at 25 °C. Cardinal temperatures for growth; min ≤ 10 °C, max ≥ 35 °C, opt 25–30 °C.

Substrates — Mangifera indica, Olea sp.

Distribution — Rudan-Kheirabad (Hormozgan Province), Southern Iran.

Notes — Phylogenetically this species is closely related to Lasiodiplodia citricola and Lasiodiplodia parva but can be distinguished on average conidial dimensions and length of its paraphyses. Conidia of

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**Fig. 4** Lasiodiplodia hormozganensis holotype. a. Conidiomata on pine needles in culture; b, c. conidia developing on conidiogenous cells between paraphyses; d, e. paraphyses; f. hyaline immature conidia; g, h. mature conidia in two different focal planes to show the longitudinal striations. — Scale bars: a = 1 000 μm; b, c = 5 μm; d–h = 10 μm.
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*L. hormozganensis* are larger (21.5 ± 1.9 × 12.5 ± 0.8 μm) than those of *L. parva* (20.2 ± 1.9 × 11.5 ± 0.8 μm), but smaller than those of *L. citricola* (24.5 ± 0.2 × 15.4 ± 1.8 μm). Paraphyses of *L. hormozganensis* are shorter (up to 83 μm) than those of *L. parva* (up to 105 μm), and *L. citricola* (up to 125 μm). This species did not produce a pink pigment in PDA cultures at 35 °C.

**Lasiodiplodia iraniensis** Abdollahzadeh, Zare & A.J.L. Phillips, *sp. nov.* — MycoBank MB516780; Fig. 5

*Teleomorph.* Unknown.

*Lasiodiplodia theobromae* phylogenetic simile, sed conidiis minoribus.

**Etymology.** Named after Iran where it was first found.

*Conidiomata* stromatic, pycnidial, produced on pine needles on WA within 2–4 wk, superficial, dark brown to black, covered with dense mycelium, mostly uniloculate, up to 980 μm, solitary, globose, thick-walled, non-papillate with a central ostiole.

*Paraphyses*, hyaline, cylindrical, thin-walled, initially aseptate, becoming up to 1–6 septate when mature, rarely branched, occasionally basal, middle or apical cells swollen, rounded at apex, up to 127 μm long, 2–4 μm wide. *Conidiophores* absent. *Conidiogenous cells* holoblastic, discrete, hyaline, smooth, thin-walled, cylindrical, 9–16 × 3–5 μm. Conidia initially hyaline, aseptate, subglobose to subcylindrical, with granular content, both ends rounded, wall < 2 μm, becoming pigmented, verrucose, ellipsoid to ovoid, 1-septate with longitudinal striations, (15.3–)17–23 (−29.7) × 11–14 μm. 95 % confidence limits = 20.6–20.8 × 13–13.1 μm (av. ± S.D. = 20.7 ± 2 × 13 ± 0.9 μm, l/w ratio = 1.6 ± 0.2).

*Culture characteristics* — Colonies with abundant aerial mycelium reaching to the lid of Petri plate, aerial mycelium becoming smoke-grey (21′′′f) to olivaceous-grey (21′′′i) at the surface and greenish grey (33′′′i) to dark slate blue (39′′′k) at the reverse after 2 wk in the dark at 25 °C. Colonies reaching 80 mm on MEA after 2 d in the dark at 25 °C. Cardinal temperatures for growth; min ≤ 10 °C, max ≥ 35 °C, opt 25–30 °C.

**Fig. 5** *Lasiodiplodia iraniensis* holotype. a. Conidiomata on pine needles in culture; b, c. conidia developing on conidiogenous cells between paraphyses; d, e. paraphyses; f. hyaline, immature conidia; g, h. mature conidia in two different focal planes to show the longitudinal striations. — Scale bars: a = 500 μm; b, c = 5 μm; d–h = 10 μm.
Substrates — Mangifera indica, Eucalyptus sp., Citrus sp., Salvadora persica, Juglans sp., Terminalia catapæ.

Distribution — Hormozgan & Golestan Provinces, Southern and Northern Iran.

Specimens examined. Iran, Hormozgan Province, Bandar Abbas, Geno mountain, on twigs of Salvadora persica, Mar. 2007. J. Abdollahzadeh and A. Javadi, holotype IRAN 14268; culture ex-type IRAN 1520C = CBS 124710; Golestan Province, Gorgan-Tooshan, on twigs of Juglans sp., June 2007, A. Javadi, IRAN 1502C = CBS 124711; Additional isolates are listed in Table 1.

Notes — Phylogenetically L. iraniensis is clearly distinct from other species, but is most closely related to L. theobromae. Conidia of L. iraniensis are smaller (15.3–17–23–(29.7)×11–14 μm) than L. theobromae ((19–)21–31–(32.5)×(12–)13–15.5 (–18.5) μm). Conidial dimensions of L. iraniensis are similar to those of L. parva, but the subglobose to subcylindrical conidia with both ends rounded distinguish this species from L. parva, in which the conidia are ovoid with apex broadly rounded and the base rounded or truncate. This species produces a pink pigment in PDA cultures at 35 °C.

DISCUSSION

In this study six species of Lasiodiplodia were associated with a variety of symptoms on a range of woody hosts in Iran. Four of these (L. citricula, L. gilanensis, L. hormozganensis and L. iraniensis) are recognised as new. All four species can be distinguished morphologically and phylogenetically from one another and from previously described species. Although 24 species are currently known in Lasiodiplodia (including those described here), cultures of only 12 are available, and all of these are of species described since 2004. For this reason it was possible to include only the more recently described species in the phylogenetic analysis. Thus, it is possible that some of the species described before 2004 are the same as those included in the phylogenetic tree in this paper. To complicate matters, none of the currently extant isolates of L. theobromae can be linked to the type. Pavlic et al. (2004) were unable to locate the holotype of L. theobromae and relied on the original description of this species, and its various synonyms, to differentiate L. gonubiensis from L. theobromae. Burgess et al. (2006), Damm et al. (2007) and Alves et al. (2008) followed the example of Pavlic et al. (2004) and included strains that have previously been recognised as representative of L. theobromae in their phylogenies. This is not wholly satisfactory, but until the species is recollected and an epitype is proposed there is no alternative. However, that does not resolve the possibility that new species names are applied to existing species. To differentiate species in the absence of cultures or sequence data it is necessary to rely on morphological characters and the original descriptions of the older species.

Species in Lasiodiplodia have been distinguished based on their DNA phylogeny in association with conidial morphology and dimensions, and morphology and size of paraphyses. Burgess et al. (2006) used septation of pycnidial paraphyses to differentiate Lasiodiplodia species including L. crassisspora, L. gonubiensis, L. rubropurpurea, L. theobromae and L. venezuelensis. However, this character needs to be interpreted carefully since paraphyses are asceptate when they are young but later they become septate. For example, Burgess et al. (2006) could not find septate paraphyses in the isolates of L. theobromae that they studied. Nevertheless, septa have been reported in this species by Punthalingam (1976) and Alves et al. (2008). Damm et al. (2007) distinguished L. plurivora from L. crassisspora and L. venezuelensis on the length and shape of the paraphyses. In a similar way in the present study maximum length of paraphyses differentiated L. gilanensis from L. plurivora, and L. hormozganensis from L. parva and L. citricula. Burgess et al. (2006) used conidial dimensions to differentiate L. crassisspora, L. rubropurpurea and L. venezuelensis from L. gonubiensis and L. theobromae. Furthermore, Alves et al. (2008) distinguished L. parva, and Pavlic et al. (2008) distinguished L. margaritacea from all other species on account of their small conidia.

Culture morphology has rarely been used as a character for species separation in Lasiodiplodia. However, Alves et al. (2008) distinguished L. parva and L. pseudotheobromae from L. theobromae based on the ability of the first two species to produce a pink pigment in PDA cultures at 35 °C. However, in this study all species except L. hormozganensis produced a pink pigment in PDA cultures at 35 °C and the theobromae isolates produced a very weak pigment. Furthermore, all isolates studied in the present work could grow at 10 °C, which is in contrast to the report of Alves et al. (2008) who found that only L. pseudotheobromae was capable of growing at this temperature. Thus, cultural characters can vary widely between isolates of any given species, and thus are of limited value in species determination.

Punthalingam (1976) regarded L. nigra, L. triflora and L. tuberculca as synonyms of L. theobromae and this was confirmed from the morphological data presented by Pavlic et al. (2004) for these four species. According to descriptions of L. abnormis, L. florii and L. thomasianae given by Saccardo (1913), these are also likely to be synonyms of L. theobromae, but this would have to be confirmed from a study of type material. From Saccardo’s (1899) description of L. paraphysaria (under Diplodia paraphysaria) this species is similar to L. gonubiensis except that the conidia are smaller (30–32 × 15–16 μm) and the paraphyses are longer (90–100 μm). Nevertheless, conidia of L. paraphysaria are substantially longer than any other known species of Lasiodiplodia, apart from L. gonubiensis. On the other hand, conidia of L. ricinii have similar dimensions to L. parva (16–19 × 10–11 μm), but the paraphyses are much shorter (25–35 μm). Little information is available on L. undulata. Abbas et al. (2004) regarded this as a synonym of L. theobromae and report the conidia as 20–32 × 13.5–19.2 μm. In the original description, Berkeley (1868) gives the conidia as 33 μm long, and this was confirmed by Saccardo (1884) who reported them as 30–33 μm long. Since conidia of L. theobromae rarely exceed 30 μm (Punthalingam 1976, Pavlic et al. 2004, Alves et al. 2008) it seems unlikely that L. undulata is a synonym of L. theobromae.

Since 2004, 12 new species have been described in Lasiodiplodia, while in the preceding 108 years only 13 species were introduced. The recent increase in the number of species recognised is largely due to the use of phylogenetic data, but is also due to sampling in relatively unexplored regions including Venezuela (Burgess et al. 2006), Western Australia (Pavlic et al. 2008) and Iran (this paper).

Since 2004 phylogenetics has played a significant role in distinguishing species in Lasiodiplodia. Pavlic et al. (2004) used ITS sequence data to distinguish L. gonubiensis from L. theobromae. Burgess et al. (2006) described a further three new Lasiodiplodia species clearly separated from L. theobromae based on ITS sequences. Inclusion of EF-1α sequences in the phylogenetic analysis gave stronger support for these species (Burgess et al. 2006). In a study of Botryosphaeriaceae on Prunus species in South Africa, Damm et al. (2007) described L. purivora as a new species. This species is closely related to L. theobromae and the two species could not be distinguished solely on the basis of ITS sequence data but they were clearly separated when EF-1α data was included. Alves et al. (2008)
used ITS and EF-1α together with morphological data to characterise a collection of isolates originally identified as *L. theobromae*. In this way they showed that *L. theobromae* is a complex of cryptic species and described *L. pseudotheo-

bromae* and *L. parva* as new. In the present paper we reveal a further four species in this complex. The eight species currently recognised in the complex cannot be distinguished solely on their ITS sequences, and phylogenetic separation is effectively based on a single gene region, namely EF-1α. However, the differences in EF-1α are fixed within the isolates studied thus far and the species can be separated on morphological features. Nevertheless, if further species appear in this complex in the future it would seem prudent to include further gene regions in the analyses to strengthen the support for them and to separate the existing ones.

All the new species described in this study were isolated from dead twigs of various hosts, but it is not known if they are primary pathogens or saprobes that developed on diseased wood. While *L. citrullina* was isolated only from *Citrus* sp., it is not possible to determine any degree of host specificity. Indeed, the other three new species were each isolated from several different hosts thus suggesting a plurivorous nature. Although *L. theobromae* has been reported from more than 500 hosts (Punithalingam 1976), host ranges of species described in recent years have been reportedly restricted (Pavic et al. 2004, Burgess et al. 2006, Damm et al. 2007). However, it is not clear if the narrow host range of the more recently described species is a reflection of sampling rather than a real representation of host range. Thus it is highly possible that there is a variation in the breadth of host range between species as seen in other genera in the Botryosphaeriaceae. For example, *D. seriata* has a very broad host range while *D. pinea* is restricted to pines and *D. corticola* is restricted to Quercus species.

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