INTRODUCTION

Almond (Prunus dulcis) is a common crop cultivated in many Mediterranean countries as well as in California (USA), South Africa, and some countries in South America and Australasia. According to the Food and Agriculture Organization (FAO 2010), Spain is the second largest almond producer after California, accounting for 11.9 % of the world’s almond production, which yielded 2.31 million tons in 2009. In Spain, almonds are grown in the south-eastern regions and on the Balearic Islands (western Mediterranean Sea), an important region with 23 432 ha of this crop cultivated in 2007 (4.7 % of the Spanish almond cultivation) (INE 2011).

In summer 2008, severe decline of almond trees was noticed in several orchards on the island of Mallorca (Balearic Islands). Disease symptoms included rapid collapse of branches during mid-summer, chlorosis of leaves, which suddenly wilted and died, as well as bud and shoot dieback. Internal wood symptoms ranged from brown to black vascular streaking, visible in cross sections as spots or circular discolouration of the xylem tissue. Additionally, wedge-shaped necroses were frequently observed. Some affected trees in the orchard died within a few weeks of showing first symptoms.

Different studies have shown that Prunus represents a rich catch-crop for many fungal trunk pathogens. Species of Botryosphaeriaceae (Aplosporella, Botryosphaeria, Diplodia, Dothiorella, Lasiodiplodia, Macrophomina, Neofusicoccum, Spenicermartina and Sphaeropsis), Calosphaeriaceae (Calosphaeria and Jattaea), Coniochaetaeae (genus Coniochaeta), Diaportheaeae (Phomopsis), Diatrypaceae (Cryptovalsa, Diatrype, Eutypa and Eutypella), Herpotrichiellaceae (Pheomoniella), Montagulaceae (Paraconiothyrium), Togniniaaeae (Pheaeoacremonium) and species of the genus Collophora have been reported on Prunus trees. A list of fungal trunk pathogens isolated from Prunus spp. and their worldwide distribution is shown in Table 1. While several fungal species belonging to a number of genera are well-recognized pathogens of Prunus trees, the causal agent of the severe decline of almond trees on the island of Mallorca is still unknown. Therefore, the objective of this study was to determine the aetiology of trunk diseases associated with wood necroses of almond trees in this region of Spain.
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**Table 1** List of fungal trunk pathogens isolated from *Prunus* spp. and their worldwide distribution.
Morphological identification and characterisation

Species of Botryosphaeriaceae were identified based on colony and conidial morphology as described by Phillips (2006). In order to enhance sporulation, cultures were amended with sterilised pine needles on 2 % water agar (WA; Biokar-Diagnostics) and incubated at 25 °C under near UV light with a 12 h photoperiod (Phillips TDL18W/33) (Slippers et al. 2004). Isolates were examined weekly for formation of pycnidia and conidia in order to record their morphology (size, shape, colour, presence or absence of septa and cell wall structure).

Since it is difficult to distinguish species or even genera within the Diatrypaceae based on morphological characters of their Libertella anamorph (Glawe & Rogers 1984), the morphological identification of Eutypa lata was only tentative using characters such as conidial size and shape, and colony characters on PDA.

Morphological characters used to distinguish Phaeoacremonium species included conidiophore morphology, phialide type and size, size and shape of hyphal warts and conidial size and shape. Colony characters and pigment production on MEA, PDA and oatmeal agar (OA; 60 g oatmeal; 12.5 g agar; Difco, France) (Crous et al. 2009) incubated at 25 °C were noted after 8 and 16 d.

Collophora species were characterised based on the presence of conidiomata, microcyclic conidiation or endoconidia additional to conidia formed on hyphae, as well as size and shape of conidia and conidiophores (Damm et al. 2010). To enhance sporulation, double-autoclaved pine needles were placed onto the surface of synthetic nutrient-poor agar medium (SNA; Nirenberg 1976). The cultures were incubated at 24 °C in the dark, microscopically examined after 2 wk and additionally inspected after 4 wk. Colony characters and pigment production were noted after 2 wk of growth on MEA, PDA and OA (Crous et al. 2009) incubated at 24 °C.

Species of Phomopsis were identified based on morphology of conidia formed in pycnidia (van Niekerk et al. 2005). The sporulation was enhanced by amending 2 % WA cultures with sterilised pine needles and incubating them at 25 °C under near UV light with a 12 h photoperiod. Isolates were examined weekly for formation of pycnidia and conidia.

Microscopic observations for all fungi were made from mycelium of colonies cultivated on the respective medium or by using slide culture technique, as explained by Arzanlou et al. (2007) when studying the genus Mycosphaerella. Photos were captured with a Nikon camera system (Digital Sight DXM 1200, Nikon Corp., Japan), with a Nikon SMZ1000 dissecting microscope (DM) or with a Nikon Eclipse 80i microscope using differential interference contrast illumination (DIC). Structures were mounted in lactic acid, and 30 measurements (1 000× magnification) were determined. The 5th and 95th percentiles were defined for all measurements with the extremes given in parentheses. Colony colours were determined using the colour charts of Rayner (1970). Cardinal growth temperatures were determined by incubating MEA plates in the dark at 6–40 °C with 3 °C intervals (Collophora) or 5–40 °C with 5 °C intervals (Phaeoacremonium), also including 37 °C. Radial growth was measured after 8 d at 25 °C (Phaeoacremonium) or 2 wk at 24 °C (Collophora). Isolates of novel species were deposited in the culture collection of the CBS-KNAW Fungal Biodiversity Centre (CBS), Utrecht, The Netherlands.
DNA isolation

Fungal mycelium and conidia from pure cultures grown on PDA for 2 wk at 25 °C in the dark were scraped and mechanically disrupted by grinding to a fine powder in liquid nitrogen with a mortar and pestle. Total genomic DNA was extracted with the E.Z.N.A. Plant Miniprep Kit (Omega Bio-tek, Norcross, GA) following the manufacturer’s instructions. DNA was visualised on 0.7 % agarose gels stained with ethidium bromide and the DNA aliquots were stored at -20 °C.

Molecular identification and phylogenetic analysis

Morphological identifications of Botryosphaeriaceae spp., dityraeous fungi and Phomopsis spp. were confirmed by sequence analysis of the internal transcribed spacer (ITS) nrDNA region using the primers ITS1 and ITS4 (White et al. 1990). Species in the Botryosphaeriaceae were also confirmed by analysis of partial β-tubulin gene (BT) sequences amplified using primers Bt2a and Bt2b (Glass & Donaldson 1995). For species of Phaeoacremonium, ± 600 bp of the 5’ end of the BT and ± 300 bp of the 5’ end of the actin (ACT) genes were amplified as described by Mostert et al. (2006) using primer sets T1 (O’Donnell & Cigelnik 1997) and Bt2b, and ACT-512F and ACT-783R (Carbone & Kohn 1999), respectively. For Collophora spp., the ITS region was amplified using the primer pairs ITS1-F (Gardes & Bruns 1993) and ITS4. Additionally, a 200-bp intron of the glyceraldehyde-3-phosphate dehydrogenase (GAPDH) and a partial sequence of the translation elongation factor 1α (EF-1α) were amplified and sequenced using the primer pairs GDF1 and GDR1 (Guerber et al. 2003) and EF1-986R (Carbone & Kohn 1999). PCR products were purified with the High Pure PCR Product Purification Kit (Roche Diagnostics, Germany) and sequenced.
in both directions by Macrogen Inc., Sequencing Center (Seoul, South Korea). Sequences were edited using the Sequencher software v. 4.7 (Gene Codes Corporation, Ann Arbor, MI).

The Collophora sequences (ITS, GAPDH, EF-1α) were added to reference sequences (Damm et al. 2010) and the outgroup (Caldophora luteo-olivacea CBS 141.41, ITS: GU128588, GAPDH: JN808849, EF-1α: JN808856). The multi-locus alignment was manually adjusted using Sequence Alignment Editor v. 2.0a11 (Rambaut 2002). To determine whether the three sequence datasets were congruent and combinable, tree topologies of 70 % reciprocal Neighbour-Joining bootstrap with Maximum Likelihood distances (10 000 replicates) with substitution models determined separately for each partition using Modeltest v. 3.5 (Posada & Crandall 1998) were compared visually (Mason-Gamer & Kellogg 1996). The Phaeacremonium sequences (BT and ACT) together with the reference sequences (Mostert et al. 2006, Damm et al. 2008b, Essaki et al. 2008, Graham et al. 2009, Gramaje et al. 2009b) and the outgroup taxa, Pleurostomophora richardsiae (ACT: AY579271, BT: AY579334) and Wuestneia molokaiensis (ACT: AY579272, BT: AY579335) obtained from GenBank were aligned using MAFFT sequence alignment program v. 6 (Katoh & Toh 2010) followed by manual adjustments of the alignments in Sequence Alignment Editor v. 2.0a11. The BT and ACT alignments were concatenated. A partition homogeneity test was conducted in PAUP (Phylogenetic Analysis Using Parsimony) v. 4.0b10 (Swofford 2003). The congruence between the ACT and TUB datasets were tested at 1 000 replicates. Phylogenetic analyses of all aligned sequence data were performed with PAUP. Alignment gaps were treated as missing data and all characters

Fig. 1 Disease symptoms on almond trees on the island of Mallorca associated with fungal trunk pathogens. a, b. Dieback and wilting of branches; c–h. internal symptoms visible when transversal and longitudinal cuts were made in branches used for fungal isolation: black spots and dark brown to black streaking of the xylem tissue (d, h), circular (c, g) or sectorial necrosis (f), and wood discoloration (e).
were unordered and of equal weight. Any ties were broken randomly when encountered. All characters were unordered and of equal weight.

Maximum parsimony analysis was performed for the combined Phaeoacremonium dataset using the heuristic search option with 10 random simple taxon additions and tree bisection and reconstruction (TBR) as the branch-swapping algorithm with the option of saving no more than 10 trees with a score greater than or equal to 5 (Harrison & Langdale 2006). The maximum parsimony analysis for the combined Collophora dataset was performed using the heuristic search option with 100 random sequence additions and TBR without further restrictions for tree saving. Branches of zero length were collapsed and all multiple, equally parsimonious trees were saved. The robustness of the trees obtained was evaluated by 1 000 bootstrap replications (Hillis & Bull 1993). Tree length (TL), consistency of the trees obtained was evaluated by 1 000 bootstrap multiple, equally parsimonious trees were saved. The robustness of the trees obtained was evaluated by 1 000 bootstrap replications (Hillis & Bull 1993). Tree length (TL), consistency (CI), retention index (RI) and rescaled consistency index (RC) were calculated.

Sequences derived in this study were lodged at GenBank, the alignments in TreeBASE (www.treebase.org/), and taxonomic novelties in MycoBank (www.Mycobank.org; Crous et al. 2004). GenBank accession numbers of the strains collected during this study are listed in Table 2. Additional GAPDH and EF-1α sequences were generated for strains CBS 141.41 (see above), CBS 120878 (JN808847, JN808854) and CBS 120873 (JN808848, JN808855).

RESULTS

Morphological identification and characterisation

Based on their appearance in culture, the isolates obtained in this study, could be assigned to five main fungal groups (Table 2). The first group was characterised by dark green or grey to dark grey fast-growing mycelium on PDA. Some isolates produced a yellow pigment after 3 days that diffuses into the agar. With age, most of these cultures developed single or grouped, black, globose fruiting bodies (pycnidia) on the surface of pine needles on WA releasing either pigmented or hyaline conidia. Based on descriptions of species of Botryosphaeriaceae (van Niekerk et al. 2004, Phillips 2006) and comparison with previously identified isolates from Spain, fungal cultures with pigmented conidia were assigned to two species: Diplodia seriata and Diplodia sp. Fungal cultures with hyaline conidia were assigned to three different species: Botryosphaeria dothidea, Neofusicoccum parvum and Neofusicoccum sp.

The second group of isolates was characterised by white to reddish cream, slow growing mycelium, turning red to blood colour with age. Isolates formed a red pigment that coloured the colony and surrounding medium. Conidiophores on hyphae were reduced to conidiogenous cells. Sporulation was abundant, with hyaline, 1-celled conidia aggregated in masses around the hyphae. All morphological characters observed were consistent with the description of Collophora spp. (Damm et al. 2010). However, the isolates could not be assigned to one of the known species.

The third fungal group was characterised by having white to white-cream cottony slow-growing mycelium on PDA lacking fruiting structures after an incubation time of 3 wk in the dark. With age, some cultures developed a grey pigment. After 3–4 wk under continuous fluorescent light, small black pycnidia were formed on the pine needles. Conidia developing in the fruiting bodies were filiform and mostly curved in shape, which corresponds to descriptions of species in the Diatrypaceae family (Glawe & Rogers 1984).

The fourth fungal group was characterised by pale to medium brown flat slow-growing cultures on MEA. Different types of phialides that were variable in size and shape were observed in the aerial mycelium, and either discrete or integrated in conidiophores. Sporulation was abundant and conidia hyaline and aseptate. All morphological characters corresponded to the genus Phaeoacremonium (Mostert et al. 2006).

The last fungal group was characterised by white, cottony, slow-growing, raised mycelium, with margins becoming pale brown with age. Dark brown, euromromatic pycnidia released a mucilaginous light-cream drop containing only one characteristic spore type, usually ovoid-ellipsoidal with one obtuse and one acute end. These morphological characteristics resembled those of Phomopsis species (van Niekerk et al. 2005). Botryosphaeriaceae spp. were the most common fungi isolated from symptomatic almond wood from Mallorca Island, followed by Eutypa lata, Phaeoacremonium spp., Collophora hispanica and Phomopsis amygdali (Table 2). Of the Botryosphaeriaceae species isolated, N. parvum was the most abundant species (13 strains). In contrast, B. dothidea was the least abundant, with only one strain. Other species of Botryosphaeriaceae were also frequently isolated, and included D. olivarum (7 strains), D. seriata (1 strain) and D. pilipes (4 strains).

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Fig. 2 One of two most parsimonious trees obtained from heuristic searches of ITS, GAPDH and EF-1α gene sequences of Collophora species. Bootstrap support (1 000 replicates) above 70% are shown at the nodes. Cadophora luteo-olivacea CBS 141.41 was used as outgroup. Ex-type strains for each species are indicated with a ‘T’ after the strain number.
N. australis (6 strains) and D. seriata (4 strains), Diatrypaceae, represented by E. lata, were also frequently isolated in this study (7 strains), while only one strain of Ps. amygdali was collected. While Ps. iranianum was infrequently isolated (one strain), the novel species Ps. amygdalinum and Co. hispanica were collected several times (4 strains each).

Species of Botryosphaeriaceae, Eutypa lata and Ps. amygdali isolates were mostly isolated from circular (Fig. 1c, g) or sectorial necrosis (Fig. 1f), and wood discoloration (Fig. 1e). Phaeoacremonium and Collophora spp. were isolated from black spots and dark brown to black streaking of the xylem tissue (Fig. 1d, h).

**Molecular identification and phylogenetic analyses**

To confirm the identification based on morphology, BLASTn searches in GenBank showed that ITS sequences of Botryosphaeriaceae isolates had 99–100 % identity with isolates of D. seriata CBS 121485 (GenBank EU650671), B. dothidea CBS 121484 (GenBank EU650670) and N. parvum CBS 110301 (GenBank AY259098). Diplodia sp. isolates showed 100 % identity with isolates previously described as D. olivarum (GenBank GQ923873, GQ923874; Lazzizera et al. 2008a), while Neofusisococcum sp. isolates showed 99 % identity with isolates previously identified as Neofusisococcum australae CBS 115185 (GenBank FJ150696) and CBS 110301 (GenBank DQ299244). The ITS sequences of the second group of isolates were 96 % identical to those of Co. africana STE-U 6113 (GenBank GQ154570) and Co. capensis STE-U6341 (GenBank GQ154574), while the EF sequences of these isolates were 91 % identical to that of Co. rubra STE-U 6198 (GenBank GQ154642). The ITS sequences of the Diatrypaceae isolates from this study had 99–100 % identity with isolates previously identified as Eutypa lata (GenBank AY462541, AY662394). Regarding Phaeoacremonium isolate Pir-1 from Mallorca Island, BT sequences had 99–100 % identity with Ps. iranianum isolates (GenBank EU128077, FJ872406). The ACT sequences of the remaining...
Phaeoacremonium isolates were 87 % identical to those of *Pm. pallidum* STE-U 6104 (GenBank EU128103) and *Pm. viticola* STE-U 6180 (GenBank EU128094), while the BT sequences were 81 % identical to those of *Pm. theobromatis* CBS 111586 (GenBank DQ173132) and *Pm. viticola* CBS 100947 (GenBank DQ173134). A BLASTn search showed that the ITS sequence of isolate Pam-1 had 100 % identity with isolates previously identified as *Ps. amygdali* (GenBank AF102996, AF102997). Sequences of three representative isolates of each species derived in this study were lodged in GenBank (Table 2).

Phylogenetic analysis was performed only with genera of unknown species, *Collophora* and *Phaeoacremonium* (Fig. 2, 3). The ITS, GAPDH and EF-1α sequence datasets of the genus *Collophora* did not show any conflicts in tree topology for the 70 % reciprocal bootstrap trees, which allowed us to combine them. The combined sequence dataset consisted of 11 isolates including the outgroup and had 1 097 included characters, of which 133 characters were parsimony-informative, 130 parsimony-uninformative and 834 constant. After a heuristic search, two equally most parsimonious trees with identical topologies were retained (length = 318 steps, CI = 0.921, RI = 0.907, RC = 0.836) of which one is shown in Fig. 2. Isolates of Col-1, Col-3, Col-4 and Col-5 from almond trees on the island of Mallorca form a distinct clade (100 % bootstrap support) sister to *Co. rubra*. Sequences of the type strains of *Co. africana* (CBS 120872) and *Co. capensis* (CBS 120879) as well as *Co. paarla* (CBS 120877) and *Co. pallida* (CBS 120878) are each present in well-supported clades (100 % bootstrap support) with no or little variability.

The partition homogeneity test of the BT and ACT alignments of *Phaeoacremonium* gave a *P*-value of 0.05 indicating that the datasets were congruent and could be combined. The combined sequence dataset consisted of 78 isolates including the outgroup and had 918 characters, of which 510 characters were parsimony-informative, 126 parsimony-uninformative and 282 constant. Ninety equally most parsimonious trees were retained (length = 2 690 steps, CI = 0.465, RI = 0.823, RC = 0.383). A tree that closely resembled the strict consensus tree was chosen and is presented in Fig. 3. The strain Pir-1, grouped inside the *Pm. iranianum* clade with 100 % bootstrap support. Four strains

![Fig 4](https://example.com/figure4.jpg)

*Collophora hispanica*. a. Colony on MEA that is stained red by the pigment exuded by the fungus; b. conidioma on pine needle; c–h. conidiogenous cells and conidia on hyphal cells; i, j. conidiophores formed in conidiomata on pine needles; k, l. microcyclic conidiation (indicated by arrows) in conidia from conidiomata (k) and from hyphal cells (l); m. conidia formed in conidiomata after 4 wk; n, o. endoconidia; p, q. conidia formed on hyphal cells after 4 wk; r. conidia formed on hyphal cells after 2 wk. a, b: DIC, c–r: DM. — Scale bars: a = 1 mm; b = 100 µm; c = 5 µm; scale bar for c applies to c–r.
(Psp 1–4) grouped together in a monophyletic clade with 100% bootstrap support, basal to the clades containing T. minima and Pm. novae-zeelandiae, with no other closely related species.

**Taxonomy**

Based on the DNA sequence analyses and morphological characters, two species, one species each of Collophora and Phaeoacremonium, proved distinct from all known species, and are newly described below.

*Collophora hispanica* D. Gramaje, J. Armengol & Damm, *sp. nov.* — MycoBank MB561926; Fig. 4

**Etymology.** Named after Spain where this fungus was first collected.

Vegetative hyphae hyaline, smooth-walled, septate, branched, 1–3.5 µm wide, lacking chlamydospores. Sporulation abundant, conidia formed on hyphal cells, occasionally in hyphae (endoconidia) and in conidiomata or due to microcyclic conidiation. **Conidiophores on hyphae** reduced to conidiogenous cells. **Conidiogenous cells** enteroblastic, hyaline, mainly intercalary, reduced to collarettes formed directly on hyphal cells or on short necks; necks cylindrical, 0.5–2 µm long; discrete phialides often observed, cylindrical to ampulliform, 4.5–8 × 1–2 µm; collarettes cylindrical to narrowly funnel shaped, very thin-walled, 0.5 µm long, opening 0.5 µm wide, inconspicuous, periclinal thickening sometimes visible. **Conidia** aggregated in masses around the hyphae, hyaline, 1-celled, cylindrical, sometimes obovate, often slightly bent, with both ends obtuse or with a papillate apex, smooth-walled, (2.5–)3.5–5(–6.5) × (1–)1.5–2 µm, av. ± SD = 4.3 ± 0.7 × 1.5 ± 0.2 µm, L/W ratio = 2.8, after 4 wk many clavate, limoniform, subapical, or irregularly inflated conidia observed that are often > 2 µm wide. **Microcyclic conidiation** occurs occasionally, with conidia developing into mother cells, becoming > 7 µm long, 2–3 µm wide, and sometimes septate, with a short neck or with a mere opening with a minute collarette at one end. **Endoconidia** uniseriate within hyphae, hyaline, 1-celled, cylindrical, slightly bent, with both ends obtuse, same size as conidia formed on hyphal cells. **Conidiomata** occasionally formed on pine needles in 2–4 wk. **Conidiophores** hyaline to slightly reddish, smooth-walled, septate, branched. **Conidiogenous cells** enteroblastic, hyaline to slightly reddish, smooth-walled, cylindrical to ampulliform, conidiogenous loci formed terminal and intercalary, immediately below the septum (acropyleurogenously), 3–6 × 1–3 µm, collarettes sometimes visible, ≤ 0.5 µm long, opening minute ≤ 0.5 µm wide, periclinal thickening not observed. **Conidia** hyaline or reddish, 1-celled, cylindrical, sometimes obovate, often slightly bent, with both ends obtuse or with a papillate apex, smooth-walled, (2.5–)3–5(–7) × 1–2(–3.5) µm, av. ± SD = 4.2 ± 1.0 × 1.7 ± 0.5 µm, L/W ratio = 2.5.

**Aerial structures in vitro on MEA:** Mycelium consisting of branched, septate hyphae that occurs singly or in bundles of up to 10; hyphae tuberculate with warts up to 2 µm diameter, verrucose to smooth, medium brown to pale brown and 2–3.5 µm wide. **Conidiophores** mostly short, usually unbranched, arising from aerial or submerged hyphae, erect to flexuous, up to 5-septate, medium brown to pale brown, verruculose on the lower part, (12–)15.5–40(–55) (av. = 29) µm long and 1.5–3 (av. = 2.1) µm wide. **Phialides** terminal or lateral, often polyphialidic, smooth to verrucose, hyaline, collarettes, 1.5–2.5 µm long, 1–1.5 µm wide; type I phialides mostly cylindrical, occasionally widened at the base, (3–)3.5–7.5(–10) × 1–2 (av. = 6 × 1.5) µm; type II phialides most predominant, either subcyllindrical or navicular, some elongate-ampulliform and attenuated at the base, (9–)10–16.5–17) × 1.5–2.5 (av. = 13.5 × 2) µm; type III phialides cylindrical to subcylindrical, 17–27 × 1.5–2.5(–3) (av. = 21 × 2) µm. **Conidia** hyaline, oblong ellipsoidal or obovoid, (3–)4–5 × 1.5–3 (av. = 4.5 × 2) µm, L/W ratio = 2.1.

On surface or submerged in the agar — **Phialides** hyaline, mostly cylindrical, 4–12 × 1–2 (av. = 7.5 ± 1.5) µm. **Conidia** hyaline, mostly allantoid, few reniform, (5–)5.5–7(–10) × 1–2 (av. = 6.5 ± 1.5) µm, L/W ratio = 4.4.

**Specimens examined.** SPAIN, Mallorca, Sant Llorenç del Cardassar, isolated from branches of *Prunus dulcis* trees, June 2010, J. Armengol, CBS H-20518 holotype, culture ex-type CBS 128568 = Col-4; Mallorca, Sant Llorenç del Cardassar, isolated from branches of *Prunus dulcis* trees, June 2010, J. Armengol, Col-1 herb, CBS H-20516, culture Col-1 = CBS 128566; Mallorca, Sant Llorenç del Cardassar, isolated from branches of *Prunus dulcis* trees, June 2010, J. Armengol, Col-3 herb, CBS H-20517, culture Col-3, CBS 128567; Mallorca, Sant Llorenç del Cardassar, isolated from branches of *Prunus dulcis* trees, June 2010, J. Armengol, Col-5 herb, CBS H-20519, culture Col-5, CBS 128569.

**Notes.** The phylogeny of the combined sequence dataset showed that *Co. hispanica* does not group with any of the known species. Colonies of *Co. hispanica* resemble those of *Co. africana* and *Co. rubra* in forming red pigments that stain the colony and surrounding medium. Conidia formed in the mycelium are often slightly curved like those of *Co. rubra*. Unlike both of those species, discrete phialides are common and endoconidia are formed similar to *Co. palida* and *Co. paarla*. Few conidiomata were formed on pine needles, but not on agar medium. The conidia in these conidiomata formed on conidio- phores similar to those of other *Collophora* species such as *Co. africana*, however no wall structures were observed.

**Collophora africana** Damm & Crous, Persoonia 24: 65. 2010


**Collophora paarla** Damm & Crous, Persoonia 24: 67. 2010.

**Notes.** *Collophora africana* and *Co. capensis* were regarded as distinct taxa based on differences in conidial morphology (on hyphae and in conidiomata) and differences in cardinal temperature requirements for growth. Furthermore, *Co. palida* and *Co. paarla* were differentiated based on their conidial morphology, width of vegetative hyphae, and exudates formed in culture (Damm et al. 2010). However, the phylogeny of the multigene sequence dataset of *Collophora* spp. generated in this study (Fig. 2), only supports two species, namely *Co. africana* and *Co. paarla*, suggesting that the observed variation (Damm et al. 2010) was not informative at species level.

**Phaeoacremonium amygdalinum** D. Gramaje, J. Armengol & L. Mostert, *sp. nov.* — MycoBank MB561925; Fig. 5

**Etymology.** Named after the host it was isolated from, almond (*Prunus dulcis*), which is in Greek amygdali (*αμυγδαλί*).

Aerial structures in vitro on MEA: Mycelium consisting of branched, septate hyphae that occurs singly or in bundles of up to 10; hyphae tuberculate with warts up to 2 µm diameter, verrucose to smooth, medium brown to pale brown and 2–3.5 µm wide. **Conidiophores** mostly short, usually unbranched, arising from aerial or submerged hyphae, erect to flexuous, up to 5-septate, medium brown to pale brown, verruculose on the lower part, (12–)15.5–40(–55) (av. = 29) µm long and 1.5–3 (av. = 2.1) µm wide. **Phialides** terminal or lateral, often polyphialidic, smooth to verrucose, hyaline, collarettes, 1.5–2.5 µm long, 1–1.5 µm wide; type I phialides mostly cylindrical, occasionally widened at the base, (3–)3.5–7.5(–10) × 1–2 (av. = 6 × 1.5) µm; type II phialides most predominant, either subcyllindrical or navicular, some elongate-ampulliform and attenuated at the base, (9–)10–16.5–17) × 1.5–2.5 (av. = 13.5 × 2) µm; type III phialides cylindrical to subcylindrical, 17–27 × 1.5–2.5(–3) (av. = 21 × 2) µm. **Conidia** hyaline, oblong ellipsoidal or obovoid, (3–)4–5 × 1.5–3 (av. = 4.5 × 2) µm, L/W ratio = 2.1.

On surface or submerged in the agar — **Phialides** hyaline, mostly cylindrical, 4–12 × 1–2 (av. = 7.5 ± 1.5) µm. **Conidia** hyaline, mostly allantoid, few reniform, (5–)5.5–7(–10) × 1–2 (av. = 6.5 ± 1.5) µm, L/W ratio = 4.4.
Culture characteristics — Colonies reaching a radius of 9.5–10 mm after 8 d at 25 °C. Minimum temperature for growth 15 °C, optimum 25 °C, maximum 30–35 °C. Colonies on MEA flat, with entire margin; after 8 d and 16 d, white to brownish drab above, buff-yellow to yellowish olive in reverse; on PDA flat, felt-like with few woolly tufts near the centre, with entire margin; after 8 d and 16 d, white to olive-brown or olive-green above, pale brownish drab towards the edge to dark greyish brown in reverse; on OA flat, with entire margin; after 8 d and 16 d white to pale olive-grey above. Colonies producing pale brown pigment on PDA.


Notes — The phylogeny of the combined sequence dataset showed that *Pm. amygdalinum* does not group with any of the known species. A distinguishing morphological feature is the frequent occurrence of polyphialides. Other *Phaeoacremonium* species that also form polyphialides include *Pm. australiense*, *Pm. fuscum*, *Pm. krajdenii*, *Pm. occidentale*, *Pm. pallidum*, *Pm. prunicolum*, *Pm. scolyti*, the anamorph of *Togninia africana* and the anamorph of *T. griseo-olivacea*. *Phaeoacremonium amygdalinum* can be distinguished from these species based on brown colonies, the production of pale brown pigment on PDA, and by the predominance of the type II phialides. This species often produced microcyclic conidia under slide culture conditions.

**DISCUSSION**

This study shows the high incidence and diversity of fungal trunk pathogens associated with wood decay symptoms on almond trees on the island of Mallorca. These include species of *Botryosphaeriaceae*, *Eutypa lata*, *Phaeoacremonium iranii-anum*, *Phomopsis amygdali* and the new species described here, namely *Collophora hispanica* and *Phaeoacremonium amygdalinum*. These species could be distinguished based on DNA sequence data and unique morphological characters.

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**Fig. 5** *Phaeoacremonium amygdalinum*. a–c. Sixteen-day-old colonies incubated at 25 °C on MEA (a), PDA (b) and OA (c); d–j. aerial structures on MEA; d–f. conidiophores with polyphialides (indicated by arrows); g, h. type III phialides; i. type III and type II phialides (indicated by arrow); j. type I phialides; k–t. aerial structures by using slide culture technique; k–m. branched conidiophores and type I phialides (indicated by arrows); n–p. type II phialides; q. type I phialides; r–s. Microcyclic conidiation; t. conidia; u–w. structures on the surface of and in MEA; u–v. adelophialides with conidia; w. conidia. — Scale bars: d = 10 µm; scale bar for d applies to d–w.
Several species of Botryosphaeriaceae were isolated from wedge-shaped wood necroses on almond trees. The majority of Botryosphaeriaceae isolates belonged to Neofusicoccum australe, N. parvum and Diplodia olivarum, while Botryosphaeria dothidea and D. seriata were only occasionally isolated. Botryosphaeria dothidea causes canker diseases in a broad range of woody plants, including several Prunus spp. (English et al. 1966, Sutton 1980). A canker of trunk and scaffold branches of almond trees was reported by English et al. (1966). This disease, sometimes called ‘band canker’, was first noted in 1959 occurring in several counties in California. However, some of these reports need to be viewed with care since many species have been relegated incorrectly to the name B. dothidea. For instance, N. ribis (as B. ribis) was previously regarded as a synonym of B. dothidea (von Arx & Müller 1954). Neofusicoccum parvum (as B. parva) was often not distinguished from N. ribis and consequently treated as B. dothidea (Slippers et al. 2004). In Spain, B. dothidea, together with D. seriata and N. parvum, is considered as the most common species associated with grapevine (Vitis vinifera) decline syndrome (Armengol et al. 2001, Aroca et al. 2006). Additionally, this species has recently been isolated from olive fruits in southern Spain showing symptoms of dalmatian disease (Moral et al. 2010). Botryosphaeria dothidea was not found during surveys on stone and pome fruit trees in South Africa (Damm et al. 2007a, Slippers et al. 2007), however it was confirmed to be associated with band canker of almond trees in California (Inderbitzin et al. 2010). This study also represents the first record of D. olivarum on almond. This species was recently associated with diseased olive drupes (Lazzizera et al. 2008a) and carob trees (Ceratonia siliqua) (Granata et al. 2011) in Italy.

The low incidence of D. seriata agrees with the results of Inderbitzin et al. (2010) from almond and peach trees in California. In contrast, this species was the most frequently isolated Botryosphaeriaceae species (43 of 67 isolates) on apricot, nectarine, peach and Japanese plum in South Africa (Damm et al. 2007a). It was also the dominant species in a study on stone and pome fruit trees in South Africa by Slippers et al. (2007), which represented over 90 % of the isolates collected over a 5-year period and in a recent study on pome fruit trees in South Africa by Cloete et al. (2011). Diplodia seriata is known to occur on a wide range of hosts (Funke et al. 1973) and to cause several diseases in some host plants, such as apple (Malus domestica) or peach (Britton & Hendrix 1982). Farr et al. (2008) listed 264 hosts under its former name of Botryosphaeria obtusa. In Spain, D. seriata also occurs on olive drupes (Moral et al. 2007).

Neofusicocum australe was frequently isolated from almond trees studied here. This species was reported from almond and plum in the Western Cape, South Africa, by Slippers et al. (2007) who considered it to be infrequent and of minimal importance on stone fruits. However, Damm et al. (2007a) found N. australae commonly on three Prunus species (peach, Japanese plum and apricot) and in different locations in this region of South Africa. This fungus is the dominant Botryosphaeriaceae species infecting native Eucalyptus species in Western Australia (Burgess et al. 2005). Neofusicocum australe was recently reported from Eucalyptus and pistachio (Pistacia vera) trees in Spain (Armengol et al. 2008), from olives in Italy (Lazzizera et al. 2008b), from avocado (Persea americana) in California (McDonald et al. 2009) and from grapevines in Australia (Taylor et al. 2005), New Zealand (Ampornsh et al. 2009), South Africa (van Nierkerk et al. 2004) and Spain (Aroca et al. 2010).

Neofusicocum parvum was the most frequently isolated Botryosphaeriaceae species in this study. Recently, this fungus was reported affecting almond trees in California (Inderbitzin et al. 2010). Neofusicocum parvum is a common pathogen of pome and stone fruit trees world-wide (Slippers et al. 2007), and judging from the frequency of isolation, it seems to be one of the most common causes of wood decay of almond trees in Spain. Therefore, it should also be taken into account with the development of disease control measures. In Spain, N. parvum has also been isolated from English walnut (Juglans regia) and Japanese plum trees (Moral et al. 2010).

Several strains of Eutypa lata were isolated during this study. This fungus is a major pathogen of cultivated crops such as apricot and grapevine and has been found all over the world (Carter 1957). In almond, the occurrence of a perithecial stroma on the dead stump of a tree was first reported by Carter (1960) in Australia. This species has also been recorded from necrotic vascular tissue associated with cankers in almond trees in Greece (Carter 1982, Rumbos 1985).

Phaeoacremonium iranianum has previously been reported affecting kiwifruit in Italy (Mostert et al. 2006) and grapevines in several countries, such as Italy (Essakhi et al. 2008), Iran (Mostert et al. 2006) and Spain (Gramaje et al. 2005a). This is the first report of Pm. iranianum on almond. This species was recently isolated from necrotic wood of apricot in South Africa (Damm et al. 2008b). However, the impact of Pm. iranianum on dieback disease on almond trees in Spain is uncertain, since only one isolate was obtained. In contrast, the new species, Pm. amygdalinum was found to be more frequently associated with wood decay symptoms of almond trees during this study.

Phomopsis amygdali was isolated from affected shoots of almond trees. This species was recently reported affecting almond branches in Portugal (Diogo et al. 2010) and Tunisia (Rhouma et al. 2008). However, Ps. amygdali is not restricted to Prunus spp. but has also been isolated from grapevines in South Africa (Mostert et al. 2001).

The fungi reported in this study were isolated from necrotic wood tissue of almond trees on the island of Mallorca. We did not determine their pathogenicity, but, most of these fungi had previously been reported to be pathogenic or potentially pathogenic to Prunus spp., such as B. dothidea (English et al. 1966, Inderbitzin et al. 2010), D. seriata (Britton & Hendrix 1982, Britton et al. 1990, Damm et al. 2007a, Inderbitzin et al. 2010), N. australae (Damm et al. 2007a), N. parvum (Inderbitzin et al. 2010), Eutypa lata (Carter & Moller 1971, English & Davis 1978, Carter 1982, Rumbos 1985), Pm. iranianum (Damm et al. 2008b) and Ps. amygdali (Diogo et al. 2010).

The results of the isolations made during this study show the complex situation generated by many fungal trunk pathogens on almond trees on the island of Mallorca, which is in agreement with previous reports on almond and several Prunus spp. in other regions in the world (Damm et al. 2007a, 2008b, 2009, 2010, Slippers et al. 2007, Inderbitzin et al. 2010).

In Spain, as well as in other countries, commercial Prunus orchards are often planted adjacent to vineyards. Most of the species isolated from almond trees in this study are known grapevine pathogens in different regions of the world. For example, species of Botryosphaeriaceae are important pathogens of grapevine, causing cankers and other dieback symptoms in all major viticulture regions worldwide (van Nierkerk et al. 2004). Eutypa lata is the causal agent of eutypa dieback, an important perennial canker disease that occurs in most countries where grapevine is cultivated (Munkvold et al. 1994). Species of Phaeoacremonium have been associated with very destructive grapevine decline diseases such as Petri disease and esca (Mostert et al. 2006). Phomopsis cane and leaf spot is an important disease of grapevines, causing serious losses to the wine industry (Mostert et al. 2001). Therefore, these fungal species could have spread from grapevine plants to almond trees. Conversely, almond orchards should be considered as...
potential sources of viable inoculum for trunk disease pathogens from which grapevines could be infected and almond trees could serve as an additional mode of pathogen survival in the absence of grapevines or Prunus plants. Further studies on the adjacent stands to the ones sampled here could provide new insights on the epidemiology of fungal trunk pathogens.

Disease management practices employed on farms where vineyards are planted in close proximity to Prunus orchards are therefore crucial for disease prevention. Since these trunk pathogens mainly infect fresh woods such as pruning wounds via air- and waterborne inoculum (Trese et al. 1980, Hewitt & Pearson 1988, Pscheidt & Pearson 1989, Larignon & Dubos 2000), having a low inoculum pressure would be a logical condition for preventing infections. Methods by which the infection of wounds could be prevented or at least reduced include the removal of dead wood from vineyard or Prunus orchard floors, in order to minimise possible infection sources, avoiding pruning immediately following rainfall and applying fungicides or biological control agents to wound surfaces immediately after pruning.

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