# Some species of fungi associated with declined Persian oak trees in Ilam province with emphasis on new records to mycobiota of Iran

Received: 09.05.2018 / Accepted: 04.09.2018

Amin Alidadi: MSc Graduate in Plant Pathology, Department of Plant Protection, College of Agriculture and Natural Resources, University of Tehran, Karaj, Iran

- Mohammad Javan-Nikkhah⊠: Prof. of Mycology and Plant Pathology, Department of Plant Protection, College of Agriculture and Natural Resources, University of Tehran, Karaj 31587-77871, Iran (jnikkhah@ut.ac.ir)
- Mojegan Kowsari: Research Assistant Prof., Microbial Biotechnology Department, Agricultural Biotechnology Research Institute of Iran (ABRII), Agricultural Research, Education & Extension Organization (AREEO), Karaj, Iran
- Saadi Karami: MSc Graduate in Plant Pathology, Department of Plant Protection, College of Agriculture and Natural Resources, University of Tehran, Karaj, Iran

Morteza Ebrahimi Rastaghi: Expert, Forests, Range and Watershed Management Organization, Tehran, Iran

## Abstract

Zagros vegetation zone is one of the most important forest regions in Iran, which consists of a diverse group of arboreal species, especially oaks (*Quercus* spp.). Ilam province located in west of Iran and in Zagros vegetation zone which has 641000 ha of oak forests that its dominant species is Persian oak (*Q. brantii*). Oak trees decline is a complicated phenomenon that may result from different kinds of agents such as fungi. In order to study on fungi associated with oak trees decline, different parts of symptomatic Persian oak trees were sampled in different regions of Ilam province during the summer and autumn of 2014–15. Fungal species were identified according to either morphological or molecular characteristics obtained from ITS of ribosomal DNA. Eleven species of eight fungal genera were identified that all of them are reported for the first time as Persian oak-associated species. Also three species including *Immersidiscosia eucalypti*, *Petriella sordida*, and *Neocamarosporium obiones* are reported and fully described here as new records to mycobiota of Iran.

Keywords: Morphology, phylogeny, Quercus spp., ribosomal DNA, Zagros vegetation zone

# برخی قارچهای همراه با درختان بلوط ایرانی با علایم زوال در استان ایلام با تاکید بر آرایههای جدید برای میکوبیوتای ایران\* دریافت: ۱۳۹۷/۰۲/۱۹ / پذیرش: ۱۳۹۷/۰۶/۱۳

امین علیدادی: دانش آموخته کارشناسی ارشد بیماری شناسی گیاهی، گروه گیاه پزشکی، پردیس کشاورزی و منابع طبیعی، دانشگاه تهران، کرج، ایران محمد جوان نیکخواه⊠: استاد گروه گیاه پزشکی، پردیس کشاورزی و منابع طبیعی، دانشگاه تهران، کرج، ایران (jnikkhah@ut.ac.ir) مژگان کوثری: استادیار پژوهش بخش بیوتکنولوژی میکروبی، پژوهشگاه بیوتکنولوژی کشاورزی ایران، سازمان تحقیقات، ترویج و آموزش کشاورزی، کرج، ایران سعدی کرمی: دانش آموخته کارشناسی ارشد بیماری شناسی گیاهی، گروه گیاه پزشکی، پردیس کشاورزی و منابع طبیعی، دانشگاه تهران، کرج، ایران مرتضی ابراهیم رستاقی: کارشناسی ارشد بیماری شناسی گیاهی، گروه گیاه پزشکی، پردیس کشاورزی و منابع طبیعی، دانشگاه تهران، کرج، ایران

خلاصه

ناحیه رویشی زاگرس به عنوان یکی از مهمترین مناطق جنگلی ایران، در بردارنده گونههای درختی مختلفی به ویژه گونههای بلوط (.Quercus spp) میباشد. استان ایلام که در حوزه رویشی زاگرس قرار می گیرد دارای جنگلهای وسیعی بوده که گونه غالب جنگلهای این ناحیه را بلوط (یرانی (brantii)) میباشد. استان ایلام که در حوزه رویشی زاگرس قرار می گیرد دارای جنگلهای وسیعی بوده که گونه غالب جنگلهای این ناحیه را بلوط (یرانی (brantii)) میباشد. استان ایلام که در حوزه رویشی زاگرس قرار می گیرد دارای جنگلهای وسیعی بوده که گونه غالب جنگلهای این ناحیه را بلوط (یرانی (brantii)) میباشد. استان ایلام که در حوزه رویشی زاگرس قرار می گیرد دارای جنگلهای وسیعی بوده که گونه غالب جنگلهای این ناحیه را بلوط به منظور جمع آوری و بررسی عوامل قارچی همراه با علایم زوال درختان بلوط در استان ایلام، نمونهبرداری از مناطق مختلف جنگلی استان ایلام و از اندامهای مختلف درختان بلوط طی تابستان و پاییز سالهای ۱۳۹۳ و ۱۳۹۴ انجام پذیرفت. شناسایی گونههای قارچی براساس خصوصیات ریختشناختی و اطلاعات توالی حاصل از نواحی TS از NA از میراز می مورت پذیرفت. به این ترتیب، تعداد ۱۱ گونه قارچی از هشت جنس مختلف قارچی شناسایی شد. گزارش وجود گونههای قارچی شاسایی شده در این مطالعه، برای نخستین بار از درختان بلوط ایرانی جدید میباشد. علاوهبراین، تمامی گونههای قارچی شناسایی شده در این مطالعه، برای نخستین بار از درختان بلوط ایرانی جداسازی و گزارش می شوند.

**واژدهای کلیدی:** دی.ان.ای ریبوزومی، ریختشناسی، فیلوژنی، مناطق جنگلی، ناحیه رویشی زاگرس

\* مستخرج از پایاننامه کارشناسی ارشد نگارنده نخست به راهنمایی دکتر محمد جوان نیکخواه و دکتر مژگان کوثری ارایه شده به پردیس کشاورزی و منابع طبیعی دانشگاه تهران

#### Introduction

Zagros forest zone is one of the most important forest area in Zagros Mountains located in western parts of Iran and contains about 5.2 billion hectares of Iran's forests area (Jazirehi & Ebrahimi Rostaghi 2013). This ecologically important area is host for a diverse group of plants such as trees and shrubs (Sagheb Talebi *et al.* 2014). Oaks (*Quercus* spp.) are the most dominant tree species present in this area (Sagheb Talebi *et al.* 2014). Among *Quercus* spp., Persian oak (*Quercus brantii*) is more widespread and frequent than other species (Jazirehi & Ebrahimi Rostaghi *l.c.*). Ilam province covers about 641000 ha (10%) of forest regions in Zagros forest zone (Ahmadi *et al.* 2014).

The decline is one of the most important diseases of oak trees around the world especially in Iran that resulted in destruction of oak trees in Zagros forest ecosystems. Oak decline is a complicated phenomenon that is generally caused by a diverse group of biotic and abiotic stresses (Akilli *et al.* 2013). Undoubtedly, biotic stresses play an important role on appearing decline symptoms on their host plants (Akilli *et al.* 1.c.). One of the most major groups of organisms affecting the oaks is those fungi, which can basically take into consideration in any oak decline projects.

A brief overview on the literature reveals that, the high numbers of studies have tried to determine the diversity of fungal communities associated with oak decline symptoms worldwide (Kowalski 1996, Bruhn et al. 2000, Thomas et al. 2002, Ragazzi et al. 2003, Kelley et al. 2009, Henriques et al. 2012, Mirabolfathy 2013, Linaldeddu et al. 2014). For instance, the association of fungal species including Botryosphaeria dothidea. Diplodia corticola. D. seriata. and Neofusicoccum parvum with evergreen oak (Q. ilex), decline has been reported by Linaldeddu et al. (2014). Luque et al. (2000) have isolated Biscogniauxia mediterranea, Botryosphaeria stevensii, Ophiostoma quercus, Phomopsis sp., Graphium sp., and Dendrophoma myriadea from leaves and stems of cork oak (Q. suber). Several species such as Apiognomonia quercina, Colpoma quercinum, Diplodia mutila, and Phomopsis quercina have been isolated from Quercus spp. in Italy (Ragazzi et al. 2003). In investigations on oak trees with decline symptoms in Zagros forests by Mirabolfathi et al. (2013) and Mirabolfathi (2013), two species, Biscogniauxia mediterranea, and Obolarina persica have reported. In addition, several species viz. Cladosporium tenellum, Paecilomyces formosus, Petriella guttulata, Preussia australis, and Sordaria sibutii have also been reported as endophytes of oak trees in Zagros forests (Hajizadeh et al. 2015).

There is limited information concerning the fungal communities along with oak trees with decline symptoms in the Zagros forest region. The aims of this study was: a) to collect the fungal isolates accompanying the oak trees with decline symptoms, and b) to identify recovered fungal isolates based on morphology and DNA sequence data, and reconstruct their phylogenetic relationships.

## **Materials and Methods**

- Sampling and isolation of fungi

In different forest regions of Ilam province (western Iran) including Tang-e Dalab, Chagha Sabz, Mehran, Eyvan (Dareh Deraz), and Malek Shahi during the summer and autumn of 2014-15; several symptomatic samples were collected from different parts (stems, leaves and roots) of Persian oak trees, placed in separate paper bags enclosed by sample information (plant tissues, geographic location and date of sampling). In mycological lab, samples were washed under running tap water to remove dusts or other surface contaminations and were subjected to air dry at room temperature for 2-3 h. Plant tissues were cut into small pieces (about 1-2 cm) and woody samples (branches and roots) were disinfested with 1% sodium hypochlorite for 1 min and ethanol 70% for 30 seconds, and then rinsed twice with sterile water. Leaf samples were surface sterilized using ethanol 70% for 1 min and rinsed twice with the sterile water. In order to isolate of fungi, samples were further cut into smaller segments (about 5

mm) at sterile conditions and paced on 2% Water Agar (2% WA) as well as Potato Dextrose Agar (PDA) and then incubated at 25 °C for 5–7 days. Fungi purification was done by transferring single spore and/or single hyphal tip grown on 2% Water Agar (2% WA) onto Potato Dextrose Agar. Purified isolates were stored on PDA slants at 4 °C for future studies.

# - Morphological diagnosis

Morphological species identification was conducted on six general culture media, including Potato Dextrose Agar (PDA), Oat Meal Agar (OA), Potato Carrot Agar (PCA), Carnation Leaf Agar (CLA), Rice Straw Agar (RSA), and 2% Water Agar supplemented with oak tissues (2% WA + sterile host material including leaves and branches). Macro- and micromorphological characters such as colonies tissue, color and diameter, sexual and asexual states characteristics were recorded and compared with the literature (Barron et al. 1961, Tanaka et al. 2011, Gruyter et al. 2013, Woudenberg et al. 2013, Verkley et al. 2014). Micromeasurements and micrographs were done by using lactophenol and lactophenol-cotton blue slide mounts with Nikon (E600) light microscope. All identified species have been deposited at the Microbial of Agriculture Biotechnology Research Institute of Iran Culture Collections (ABRIICC).

# - DNA extraction, PCR amplification and sequencing

Total genomic DNA was extracted according to the protocol described by Zhong & Steffenson (2001). The PCR amplifications were done with the primer pairs NS1/NS4 for SSU and ITS1/ITS4 for ITS-rDNA (White *et al.* 1990), and LROR/LR5 for LSU (Rehner & Samuels, 1995). The PCR was performed in a final volume of 25 µl containing 17.85 µl deionized water, 2.5 µl PCR buffer 10X (Sinagene, Iran), 1.5 mM MgCl<sub>2</sub>, 0.2 mM dNTPs, 0.75 U of Taq DNA polymerase (Sinagene, Iran), 0.2 pmol of each primer and 10–30 ng/µl DNA template. Conditions for each genomic region consisted of an initial denaturation of 4 min at 94 °C followed by 35 cycles denaturation of 50 s at 94 °C, annealing of 50 s at 57 °C and elongation of 50 s at 72 °C, with a final elongation step of 10 min at 72 °C. The PCR products were visualized on a 1% agarose gel to validate the presence and size of amplicons. The PCR products were purified and sequenced by Macrogen Corporation (South Korea).

# - Alignment and phylogenetic analysis

Newly generated sequences were observed and edited in BioEdit Ver. 7.2.5 (Hall 1999) and were subjected to BLAST search tool in GenBank nucleotide database. Required sequences for phylogenetic analysis were retrieved from GenBank (Table 1) and multiple sequence alignments were generated using Clustal X software (Thompson et al. 1997). Phylogenetic estimates were evaluated using the Maximum Parsimony Analyses (MP) in MEGA 6.0 (Tamura et al. 2013). MP analyses were done by using heuristic searches with 1000 random sequence additions and branch swapping with Tree-Bisection-Reconnection (TBR) algorithm, gaps treated as missing data and the reliability of resultant trees was determined by bootstrap values in 1000 replicates (Felsenstein 1985). All sequences in present study were submitted to GenBank nucleotide database and accession numbers have been recorded.

#### **Results and Discussion**

Ninety-eight isolates of fungi were obtained from twigs, trunck, leaf and root. Finaly, eleven species belong to eight genera including *Alternaria atra*, *A. infectoria*, *A. consortialis*, *A. molorum*, *Chaetomium globosum*, *Epicoccum nigrum*, *Immersidiscosia eucalypti*, *Kalmusia variispora*, *Petriella sordida*, *Neocamarosporium obiones*, and *Sordaria fimicola* were identified by morphological characteristics and molecular data. *I. eucalypti*, *P. sordida*, and *N. obiones* are reported as new records to micobiota of Iran.

## - Phylogenetic analyses

Ninety-eight isolates were morphologically characterized, and were placed into 11 species belonging to eight genera viz. Alternaria, Epicoccum, Kalmusia, Neocamarosporium, Petriella, Immersidiscosia, Sordaria, and Cheatomium. The PCR amplification of ITS-rDNAproduced 500-570 bp DNA fragments. Multiple alignment of ITS-rDNA sequences of 11 isolates of this study together with the ITS-rDNA sequences of 25 species downloaded from GenBank and Paecilomyces divaricatus as the outgroup taxon (Table 1), yielded a 556-characters (nucleotides + gaps) dataset, of which 225, 298 and 276 characters were constant, parsimony uninformative and parsimony informative, respectively. MP analysis of ITS-rDNA confirmed identification of all morphologically identified species (Fig. 1). Phylogenetic analysis of ITS-rDNA revealed all identified species are well-clustered in belonging two highly supported clades to Dothidiomycetes and Sordariomycetes (Fig. 1).

# Toxonomy

*Immersidiscosia eucalypti* (Pat.) Kaz. Tanaka, Okane & Hosoya, in Tanaka, Endo, Hirayama, Okane, Hosoya & Sato, Persoonia, Mol. Phyl. Evol. Fungi 26: 94 (2011). (Fig. 2)

Colonies on OA yellow to pale orange, with white arachnoid mycelia, 27 mm in diam. after 10 days (Fig. 2a), on PDA 37 mm after seven days; pycnidia developed slightly on 2% WA + oak tissues and RSA media after three weeks, immerged globose to ovoid and commonly unilocular; conidiophores cylindrical, branched and up to 40  $\mu$ m in length (Fig. 2b); conidiogenous cells holoblastic, hyaline, cylindrical and  $1.5-2 \times 4-18 \mu$ m in size (Fig. 2c-d); conidia cylindrical, hyaline, with a filiform appendage at the both ends, with three transverse septa and  $2.5-3 \times 14-21 \mu$ m in size; apical and basal cells of conidia  $2.5-3 \mu$ m in length and the assemblage length of two median cells  $2.5-3 \times 14-21 \mu$ m; lengths of apical and basal appendages  $7.5-15 \mu$ m and  $9-15 \mu$ m, respectively (Fig. 2e-f).

Note: The genus Immersidiscosia is monotypic only one species, with i.e. *I*. eucalypti (http://www.indexfungorum.org), which was first described from dead leaves of Laurus nobilis. *al.* 2011). (Tanaka Although, Ouercus myrsinifolia, Eucalyptus sp., and Ardisia japonica are the additional hosts for this species (Tanaka et al. 2011). In the present study, this species was isolated from Persian oak leaves with blight symptoms. The closest genus to the Immersidiscosia is Discosia that can be well distinguished from each other by ITSrDNA (Fig. 1, clade H). In this study, the GenBank blast searches of LSU region sequences for two 17BR1 isolates; 17RA1 (KY825092) and (KY825093) shows high similarity to I. eucaliypti (AB593723) too, (823/823 bp (100%) identity and 819/821 bp (99%) identity, respectively).

Specimens examined: Isolates 17RA1 (ABRIICC 10035) and 17BR1 (ABRIICC 10036), isolated from Persian oak leaves, Ilam province, Tang-e Dalab, N33 42.212 E46 22.739, Sept. 2014 and Oct. 2015, A. Alidadi and S. Karami.

Taxon	Strain	GenBank No.	Reference
Alternaria atra	91RF2	KY751367	This study
A. atra	UAMH 7840	AY625072	Meklin et al. 2004
A. consortialis	45SP	KY751366	This study
A. consortialis	CBS 104.31	KC584247	Vu et al. 2019
A. infectoria	18SA	KY751365	This study
A. infectoria	CBS 112250	FJ214897	Andersen et al. 2009
A. malorum	112SA	KY751368	This study
A. malorum	CBS 900.87	FJ214860	Crous et al. 2009
Ascochyta pisi	CBS 122750	KT389477	Chen et al. 2015
A. rabiei	CBS 237.37	KT389479	Chen et al. 2015
Chaetomium elatum	C29	HM365236	Asgari & Zare 2011
C. globosum	57SA	KY783412	This study
C. globosum	C63	HM365254	Asgari & Zare 2011
C. globosum	CBS 161.52	KM655335	Vu et al. 2019
Discosia pseudoartocreas	CPC 21117	KF777161	Crous et al. 2013
D. pseudoartocreas	346Jb14	KU516455	Crous et al. 2013
Epicoccum nigrum	30SA1	KY783413	This study
E. nigrum	CBS 505.85	FJ426997	Aveskamp et al. 2009
E. nigrum	Zbf-S21	KX065012	Li et al. 2016
Immersidiscosia eucalypti	17RA1	KY783415	This study
I. eucalypti	KT2191	AB594791	Tanaka <i>et al</i> . 2011
I. eucalypti	KT2115	AB594793	Tanaka et al. 2011
Kalmusia italica	MFLUCC 13-0066	KP325440	Thambugala <i>et al</i> . 2015
K. variispora	95SA2	KY783414	This study
K. variispora	CBS 197.82	JX496053	Verkley et al. 2014
K. sarothamni	CBS 116474	KF796676	Zhang et al. 2014
Paecilomyces divaricatus	CBS 284.48	MH856344	Vu et al. 2019
Petriella sordida	90SA3	KY783417	This study
P. sordida	CBS 297.58	AY882359	Rainer et al. 2006
P. setifera	CBS 745.69	AY882350	Vu et al. 2019
Neocamarosporium betae	CBS 523.66	FJ426981	Aveskamp et al. 2009
N. obiones	NBR1	KY783416	This study
N. obiones	CBS 432.77	GU230752	De Gruyter et al. 2012
N. obiones	CBS 786.68	MH859227	De Gruyter et al. 2012
Sordaria fimicola	96SA	KY783418	This study
S. fimicola	CBS 508.50	AY681188	Vu et al. 2019
S. sibutii	CBS 768.96	AY681180	Vu et al. 2019

Table 1. ITS1-rDNA sequences used in phylogenetic analysis



Fig. 1. One of the most parsimonious tree inferred from the ITS rDNA sequences of 31 taxa belong to *Dothideomycetes* and *Sordariomycetes*. The numbers in front of the nodes show the bootstrap values from 1000 replicates. The ITS sequence of *Paecilomyces divaricatus* was used as out group.



Fig. 2. Immersidiscosia eucalypti: a. Colony on OA, b. Conidiomata and conidiophores c-d. Conidiogenous cells, e-f. Conidia.

*Kalmusia variispora* (Verkley, Göker & Stielow) Ariyawansa & K.D. Hyde, in Ariyawansa, Tanaka, Thambugala, Phookamsak, Tian, Campo, Fungal Diversity 68: 85 (2014). (Fig. 3)

Colonies on PDA cottony, olive green at center, white at margin and 30 mm in diam. after 10 days (Fig. 3a); on OA olive green with white and loose aerial arachnoid hyphae (Fig. 3b). Pycnidia developed on OA three weeks after incubation at 25 °C under 12/12 h nUV photoperiod, mostly spherical, dark brown to black, thick-walled with angular tissues and a distinct ostiole, commonly observed in cross-section as multi-locular and/or sometimes with merged locules and seen as unilocular (Fig. 2d). Conidiophores simple or branched with integrated conidiogenus cell (Fig. 3e-f). Conidiogenous cells phialidic, cylindrical to oblong at the apex and 4–15  $\times$  2–4 µm in size. Conidia varied in shape, spherical to ovoid, hyaline at beginning then turning to olive brown to pale brown, aseptate and 2.5–3  $\times$  1–1.5 µm in size (Fig. 3g-h), usually exudate out from pycnidia in a black droplet (Fig. 3c).

Note: Verkley et al. (2014) have first reported this species on Erica carnea, and Vitis vinifera. In present study, this species is reported for the first time on Persian oak. This species was originally known as Dendrothyrium variisporum. In a multi-gene based study by Ariawansa et al. (2014), D. variisporum and D. longisporum were well clustered with K. ebuli in a same clade. ITS-rDNA sequences of the members of this genus are nearly identical and in this study K. variisporum, and K. italica were grouped in the same sub-clade (Fig. 1, clade d). Ariyawansa et al. (2014) separated these species from each other based on analyses of concatenated ITS-rDNA with LSU, SSU and -tubulin gene sequences. In this study, the LSU regions were amplified and sequenced. The blast searches of the LSU region sequences of this species in GenBank (KY825094) revealed 100% identity (873/873 bp) to *K. variiposa* (CBS 121517) as well.

Specimen examined: Isolate 95SA2 (ABRIICC 10037),

isolated from Persian oak branches, Ilam province, Chagha Sabz, N33 35.669 E46 26.411, Oct. 2015, A. Alidadi and S. Karami.



Fig. 3. *Kalmusia variispora*: a. Colony on PDA, b. Colony on OA, c. Pycnidia formed on OA, d. Cross-section of pycnidia, e-f. Conidiogenous cell, g-h. Conidia.

*Petriella sordida* (Zukal) G.L. Barron & J.C. Gilman, Can. J. Bot. 39: 839 (1961). (Fig. 4)

Colonies on PDA white, flat, slow growing and reaching 40 mm in 30 days (Fig. 4a); on OA 30 mm and on CMA 22 mm. This species apparently forms two different kinds of conidial forms; *Graphium* and *Sporotrichum* state. The *Sporotrichum* state develops on PDA two days after incubation, and includes hyaline and spherical to ovoid conidia with  $4-7 \mu m$  in size, produced on hyaline, simple and narrow conidiophores which

resulted terminally from vegetative hyphae (Fig. 4g). The *Graphium*-like conidial state develops on OA, 7–10 days after incubation as straight and pedicellate synnema, dark brown in color and up to 0.5–0.95 mm in length (Fig. 4h). Conidia one-celled, kidney shape or cylindrical and 2.5–4 × 6–10  $\mu$ m in size (Fig. 4i). Ascomata were produced on CLA about 14 days after incubation (Fig. 4b). Perithecia black, globuse to subglobuse at the base, covered with brown, septate hairs and with short perithecial neck covered with short hairs. These hairs are

peritecial neck, and spherical, asymmetric and smaller

curly at the base of perithecia (Fig. 4d). Asci clavate, spherical to ovoid, reddish brown with a lot of oil droplets and  $4.5-5.5 \times 9-11 \,\mu$ m in size (Fig. 4e-f). Note: The most morphologically closest species to *Petriella sordida* is *P. guttulata*. However, these species are differentiated from each other by oblong ellipsoid conidia with truncate at the base, ascomata without

Oct. 2015, A. Alidadi and S. Karami.



Fig. 4. *Petriella sordida*: a. Colony on PDA, b. Perithecia on CLA, c. Synnema formed on OA, d. Perithecia, e-f. Ascospores, g. Conidiophores and conidia of *Sporotrichum* state, h-i. Synnema and conidia of *Graphiam* state.

*Neocamarosporium obiones* (Jaap) Wanas. & K.D. Hyde, in Wanasinghe, Hyde, Crous, Wijayawardene, Jeewon, Jones, Bhat, Phillips, Groenewald, Dayarathne, Phukhamsakda, Thambugala, Bulgakov, Camporesi, Gafforov, Mortimer & Karunarathna, Stud. Mycol. 87: 249 (2017) (Fig. 5).

Colonies on MEA cottony, white to gray and 40 mm after 10 days under darkness at 25 °C and (Fig. 5b). Pycnidia developed on PDA under 12/12 h photoperiod nUV at 25 °C seven days after incubation (Fig. 5c-d), ovoid in shape with distinct ostiole covered with mycelial hairs and 250–260  $\mu$ m in size (Fig. 5e). Pycnidial wall consisted of an outer layer with darkened and thickened pseudoparenchyma cells and a hyaline inner layer including hyaline conidiophores. Conidiphores small, hyaline, produced from pycnidial inner layer, pyriform with a short neck and rarely long, aseptate and 5–10 × 3–4  $\mu$ m in size (Fig. 5f-g). Conidia single, yellow to pale brown, with 1–2 transverse septa, ellipsoid to ovoid and  $10-13(-15) \times 4-5(-6)$  in size (Fig. 5h-i). Chlamydospores were also produce on vegetative hyphae.

Note: This species was formerly placed in Ascochyta (Didymellaceae) and known as A. obiones that was first described on Halimione portulacoides (Dickinson & Morgan 1966). As a result of a phylogenetic study by De Gruyter et al. (2013), this species was clustered with A. hyalospora, A. caulina, and Phoma betea in Pleosporaceae and was named as Pleospora halimiones. The subsequent studies on Pleosporineae (Dothideomycetes) based on multi-gene phylogenetic analysis showed that the P. halimiones clustered with of Neocamarosporium species genus in family. Thus, the the Neocamarosporiaceae

*P. halimiones* is synonymized under *Neocamarosporium obiones* (Wanasinghe *et al.* 2017).

We identified this species according to morphological characteristics as well as phylogenetic analysis based on ITS-rDNA sequences (Fig. 1, clade b-c). In addition, the blast searches of newly generated LSU (KY950252) and SSU (KY950253) sequence were revealed a high identity to *N. obiones* (CBS 432.77) (100% identity). This species is reported for the first time from Persian oak leaves and is also new records to mycobiota of Iran.

Specimen examined: Isolate NBR1 (ABRIICC 10039), isolated from Persian oak leaves, Ilam province, Mehran, N33 32.463 E46 14.996, Sept. 2014, A. Alidadi and S. Karami.



Fig. 5. *Neocamarosporium obiones*: a. Pycnidia on oak leaf surface, b. Colony on MEA, c-d. Pycnidia on PDA surface after seven days, e. Pycnidia with mycelial hairs, f-g. Conidiophores, h-i. Conidia.

*Alternaria atra* (Preuss) Woudenb. & Crous, Stud. Mycol. 75(1): 204 (2013). (Fig. 6a-b)

The main morphological characteristics of this species are the production of conidia with golden yellow, pale brown to brown, with warty outer conidial wall, spherical to ellipsoidal shape and with cruciform conidial septa; 1–2 transverse septa and 1–2 longitudinal septa (Fig. 6a-b).

Note: This species was formerly known as *Ulocladium atrum* (Simmons 1998). In a revision by Woudenberg *et al.* (2013), all Alternarioid fungi were kept under a single generic name as *Alternaria* and resulted in the synonymy of genus *Ulocladium* as section *Ulocladioides* within genus *Alternaria*. In the present study, *A. atra* and *A. consortialis* well clustered with the *Alternaria* species in section *Ulocladioides* (100% bootstrap values; Fig. 1, clade a), showing that, the ITS-rDNA sequences as a universal barcode for fungi, has less phylogenetic information within this genus and needed some more genomic loci to accurate species delineation in the genus (Woudenberg *et al.* 2013). This species has frequently been reported from Iran on different substrates (Ershad 2009, Hergholi *et al.* 2015).

Specimen examined: Isolate 91RF2 (ABRIICC 10033), isolated from Persian oak branches, Ilam province, Chagha Sabz, N33 35.723 E46 26.518, Sept. 2014, A. Alidadi and S. Karami.

*Alternaria consortialis* (Thüm.) J.W. Groves & S. Hughes [as 'consortiale'], in Hughes, Can. J. Bot. 31: 636 (1953). (Fig. 6c, e)

The most important morphological characteristics of this species are the conidia often are produced singly or in short chains (2–3 spores), usually with smooth and rarely verrucous wall surface, 1–5 transverse septa and 1–3 longitudinal septa (Fig. 6d-e).

Note: This species was formerly placed in genus *Ulocladium* as *U. consortiale* (Woudenberg *et al.* 2013). It has transferred to the genus *Alternaria* by the mutli-gene phylogeny, and currently known as *A. consortialis* (Woudenberg *et al.* 2013). This species was first reported

as an endophytic fungus of peach and apricot trees in Iran (Hashemlou *et al.* 2015).

Specimen examined: Isolate 45SA (ABRIICC 10032), isolated from Persian oak roots, Ilam province, Chagha Sabz, N33 35.775 E46 26.497, Sept. 2014, A. Alidadi and S. Karami.

Alternaria infectoria E.G. Simmons, Mycotaxon 25(1): 298 (1986). (Fig. 6f-h)

Colonies on PCA were olive at center and white at margin, sporulation at dense clumps of conidia in short to moderate conidial chains in which the first-formed conidia are robust and large (Fig. 6f-h).

Note: Alternaria infectoria was first reported by Ghosta et al. (2003) from wheat seeds in Iran. In the last revision of the genus Alternaria, 27 sections were determined that A. infectoria is placed in section Infectoriae as the type species (Lawrence et al. 2016). In this study, A. infectoria was isolated from trunks and branches of Persian oak trees.

Specimen examined: Isolate 18SA (ABRIICC 10030), isolated from trunks of Persian oak trees, Ilam province, Tang-e Dalab, N33 42.188 E46 22.684, Oct. 2015, A. Alidadi and S. Karami.

*Alternaria malorum* (Ruehle) U. Braun, Crous & Dugan, in Braun, Crous, Dugan & de Hoog, Mycol. Progr. 2(1): 5 (2003). (Fig. 6i-l)

Sporulation abundant from both agar surface and aerial mycelia in long and branched to unbranched spore chains. Conidiophores were short, straight to somewhat curved. Conidia were cylindrical, pale olive, one-celled with smooth outer wall. Ramoconidia were rectangular with 1–3 even darkened and thickened transverse septa (Fig. 6j-l).

Note: This species has been recognized as *Chalastospora* gossypii (Crous et al. 2009), and currently is placed in section *Chalastospora* within genus *Alternaria* as *A. malourum* (Woudenberg et al. 2013). It has been reported from Iran on barley (Asgari et al. 2004).

However, Hergholi *et al.* (2015) have also reported *A. malorum* as endophyte of *Vitis vinifera*. Specimen examined: Isolate 112SA (ABRIICC 10028), isolated from Persian oak roots, Ilam province, Tang-e Dalab, N33 51.0220 E46 11.938, Oct. 2015, A. Alidadi and S. Karami.



Fig. 6. a-b. *Alternaria atra* (a. Colony on PCA, b. Conidiophores and conidia), c-e. *Alternaria consortialis* (c. Colony on PCA, d. Secondary coniciophores, e. e. Short chains of conidia, f-h. *Alternaria infectoria* (f. Colony on PCA, g. Sporulation on PCA surface, h. Branching chains conidia), i-l. *Alternaria malorum* (i. Colony on PCA, j-k. Branching chains, l. Ramoconidia).

*Chaetomium globosum* Kunze, in Kunze & Schmidt, Mykologische Hefte (Leipzig) 1: 16 (1817). (Fig. 7a-c)

Colonies on MEA cottony, yellowish brown at center and white at margin. Perithecia brown, spherical to ellipsoidal and covered with long and covered with long and curly hairs at the upper half of ascomata. Asci hyaline, clavate, with long pedicel and consisting eight lime shape, one-celled and dark brown to brown ascospores (Fig. 7a-c).

Note: This species has been reported from different hosts including cotton, maize and soybean in Iran (Ershad 2009). Based on the literature, this species has only been reported from three oak species *viz. Quercus germana*, *Q. robur*, and *Q. sartorii* (Heredia 1993, Mulenko 2008), and here is reported on Persian oak.

Specimen examined: Isolate 57SA (ABRIICC 10029), isolated from Persian oak branches, Ilam province, Dareh Draz, N33 46.153 E46 22.205, Sept. 2014, A. Alidadi and S. Karami.

*Epicoccum nigrum* Link, Magazin der Gesellschaft Naturforschenden Freunde Berlin 8: 32 (1816). (Fig. 7d-f).

Colonies on PDA smooth, olive green at center and white at margin. Sporodochia pulvinate and appeared as a black conidial mass on OA. Conidia spherical to pyriform, dark brown with irregular trans- and longitudinal septa (Fig. 7d-f).

Note: *Epiccocum nigrum* is a ubiquitous fungal species and were frequently isolated from a wide variety of substrates such as soils, plants, air, animals, foods, textiles etc. It is commonly known as a potential saprobe associating with plant debris decay; however, it can also see as secondary plant pathogen (Ellis 1971, Domsch *et al.* 2007). In Iran, *E. nigrum* has been isolated from peach, apricot and almond trees (Dokhanchi *et al.* 2014), yew trees (Jam Ashkezari *et al.* 2013), grape vine (Hergholi *et al.* 2015), and here is reported on Persian oak.

Specimen examined: Isolate 30SA1 (ABRIICC 10034), isolated from trunks of Persian oak trees, Ilam province, Tang-e Dalab, N33 41.830 E46 24.014, Sept. 2014, A. Alidadi and S. Karami.

*Sordaria fimicola* (Roberge ex Desm.) Ces. & De Not., Comm. Soc. Crittog. Ital. 1(fasc. 4): 226 (1863). (Fig. 7g-h)

Colonies on PDA white at beginning then subsequently turning to dark brown. Perithecia formed abundantly on PDA. Asci hyaline and 8-spored. Ascospores ellipsoidal, pale yellow when are young and brown to dark brown at maturity, and surrounded by a hyaline gelatinous sheath with a germinating pore at the apex (Fig. 7g-h).

Note: Sordaria fimicola is phylogenetically close to *S. sibutii* based on the ITS-rDNA sequences (Fig. 1, clade e). However, *S. sibutii* is differentiated from *S. fimicola* due to its ascospores shape, dimensions and outer wall smoothness as well as lack of hyaline gelatinous sheath surrounding the ascospores (Huhndorf *et al.* 2004, Cai *et al.* 2006). Sordaria fimicola has been described as endophyte of *Q. ilex* in Spain (Collado *et al.* 1996). Sordaria sibutii, the closest species to *S. fimicola*, has been previously reported on *Q. brantii* in Iran (Hajizadeh *et al.* 2015). However, this study is the first report of *S. fimicola* on branches of Persian oak trees with decline symptoms in Iran.

Specimen examined: Isolate 96SA (ABRIICC 10040), isolated from Persian oak branches, Ilam province, Chagha Sabz, N33 35.694 E46 26.449, Oct. 2015, A. Alidadi and S. Karami.



Fig. 7. a-c. *Chaetomium globosum* (a. Colony on MEA, b. Ascomata, c. Ascospors), d-f. *Epicoccum nigrum* (d. Colony on PDA, e. Sporodochium, f. Conidia), g-h. *Sordaria fimicola* (g. Asci, h. Ascospores with gelatin sheath).

#### Acknowledgments

The authors would like to thank Ebrahim Karimi for his help in sampling, the Agriculture Biotechnology Research Institute of Iran (ABRII), University of Tehran, and also Forests, Range and Watershed Management Organization for their financial supports to this projects.

## References

- Ahmadi, R., Kiadaliri, H., Mataji, A. & Kafaki, S. 2014. Oak forest decline zonation using AHP model and GIS technique in Zagros Forests of Ilam province. Journal of Biodiversity and Environmental Sciences 4: 141–150.
- Akilli, S., Uluba Serçe, Ç., Katırcıo lu, Y.Z. & Maden, S. 2013. Does *Pythium anandrum* contribute to the dieback of sessile oak (*Quercus petraea*) in Turkey?. Forest Pathology 43: 505–508.
- Andersen, B., Sørensen, J.L., Nielsen, K.F., van den Ende, B.G. & de Hoog, S. 2009. A polyphasic approach to the taxonomy of the *Alternaria infectoria* species-group. Fungal Genetics and Biology 46: 642–656.
- Ariyawansa, H.A., Tanaka, K., Thambugala, K.M., Phookamsak, R., Tian, Q., Camporesi, E., Hongsanan, S., Monkai, J., Wanasinghe, D.N., Mapook, A. & Chukeatirote, E. 2014. A molecular phylogenetic reappraisal of the *Didymosphaeriaceae* (= *Montagnulaceae*). Fungal Diversity 68: 69–104.
- Asgari, B., Zare, R. & Payghami, E. 2004. Hyphomycetous fungal community of barley phylloplane in East Azarbaijan province with emphasis on new taxa for Iranian fungal flora. Rostaniha 5: 171–197.
- Aveskamp, M.M., Verkley, G.J., de Gruyter, J., Murace, M.A., Perello, A., Woudenberg, J.H., Groenewald, J.Z. & Crous, P.W. 2009. DNA phylogeny reveals polyphyly of *Phoma* section *Peyronellaea* and multiple taxonomic novelties. Mycologia 101: 363–382.
- Barron, G.L., Cain, R.F., & Gilman, J.C. 1961. The genus *Microascus*. Canadian Journal of Botany 39: 1609–1631.
- Bruhn, J.N., Wetteroff, J.J., Mihail, J.D., Kabrick, J.M. & Pickens, J.B. 2000. Distribution of *Armillaria* species in upland Ozark Mountain forests with respect to site, overstory species composition and oak decline. Forest Pathology 30: 43–60.

- Cai, L., Jeewon, R. & Hyde, K.D. 2006. Phylogenetic investigations of *Sordariaceae* based on multiple gene sequences and morphology. Mycological Research 110: 137–150.
- Chen, Q., Jiang, J.R., Zhang, G.Z., Cai, L. & Crous, P.W. 2015. Resolving the *Phoma enigma*. Studies in Mycology 82: 137–217.
- Collado, J., Platas, G. & Peláez, F. 1996. Fungal endophytes in leaves, twigs and bark of *Quercus ilex* from Central Spain. Nova Hedwigia 63: 347–360.
- Crous, P.W., Braun, U., Wingfield, M.J., Wood, A.R., Shin, H.D., Summerell, B.A., Alfenas, A.C., Cumagun, C.J.R. & Groenewald, J.Z. 2009. Phylogeny and taxonomy of obscure genera of microfungi. Persoonia 22: 139.
- Crous, P.W., Wingfield, M.J., Guarro, J.,
  Cheewangkoon, R., Van der Bank, M., Swart,
  W.J., Stchigel, A.M., Cano-Lira, J.F., Roux, J.,
  Madrid, H. & Damm, U. 2013. Fungal planet
  description sheets: 154–213. Molecular
  Phylogeny and Evolution of Fungi. Persoonia 31: 188–296.
- De Gruyter, J., Woudenberg, J.H.C., Aveskamp, M.M., Verkley, G.J.M., Groenewald, J.Z. & Crous, P.W. 2013. Redisposition of *Phoma*-like anamorphs in *Pleosporales*. Studies in Mycology 75: 1–36.
- Dickinson, C.H. & Morgan-Jones, G. 1966. The mycoflora associated with *Halimione portulacoides*: IV. Observations on some species of *sphaeropsidales*. Transactions of the British Mycological Society 49: 43–55.
- Dokhanchi, H., Arzanlou, M. & Babai-Ahari, A. 2014. Identification of the fungal species associated with trunk diseases of stone fruit trees in East and West Azerbaijan provinces. Applied Researches in Plant Protection 2: 30–45.
- Domsch, K.H., Gams, W. & Anderson, T.H. 2007. Compendium of Soil Fungi. 2nd Ed., IHW Verlag, Eching bei München, 672 pp.

- Ellis, M.B. 1971. Dematiaceous *Hypomycetes*. Commonwealth Mycological Institute Publication, Kew. Surrey. England, 608 pp.
- Ershad, J. 2009. Fungi of Iran. Iranian Research Institute of Plant Protection Press. Tehran, 531 pp.
- Felsenstein, J. 1985. Confidence limits on phylogenies: an approach using the bootstrap. Evolution 39: 783–791.
- Ghosta, Y., Ershad, D., Zare, R. & Goltapeh, E.M. 2003. A taxonomic study on *Alternaria* species in Iran (2). Rostaniha 4: 105–122.
- Hajizadeh, A., Amini, J. & Abdollahzadeh, J. 2015. New records of endophytic fungi isolated from oak trees in Kurdistan province (Iran). Rostaniha 16: 109–122.
- Hall, T.A. 1999. BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucleic Acids Symposium Series 41: 95–98.
- Henriques, J., Inácio, M.L., Lima, A. & Sousa, E. 2012. New outbreaks of charcoal canker on young cork oak trees in Portugal. IOBC/wprs Bulletin 76: 85–88.
- Heredia, G. 1993. Mycoflora associated with green leaves and leaf litter of *Quercus germana*, *Quercus sartorii* and *Liquidambar styraciflua* in a Mexican cloud forest. Cryptogamie Mycologie 14: 171–183.
- Hergholi, N., Ghosta, Y., Javan-Nikkhah, M., Campisano, A. & Pancher, M. 2015. New species of endophytic fungi from grapevine (*Vitis vinifera*) in Iran. Rostaniha 16: 17–35.
- Huhndorf, S.M., Miller, A.N. & Fernández, F.A. 2004. Molecular systematics of the *Sordariales*: the order and the family *Lasiosphaeriaceae* redefined. Mycologia 96: 368–387.
- Jam Ashkezari, S., Fotouhifar, K.B. & Farzaneh, M. 2013. Introduction of some endophytic fungi of common yew (*Taxus baccata*) in Iran. Rostaniha 14: 184–197.

- Jazirehi, M.H. & Ebrahimi Rostaghi, M. 2013. Silviculture in Zagros. University of Tehran Press. 2nd Ed., 560 pp.
- Kelley, M.B., Fierke, M.K., & Stephen, F.M. 2009. Identification and distribution of *Armillaria* species associated with an oak decline event in the Arkansas Ozarks. Forest Pathology 39: 397–404.
- Kowalski, T. 1996. Oak decline II. Fungi associated with various types of lesions on stems and branches of young oaks (*Quercus robur*). Österreichische Zeitschrift für Pilzkunde 5: 51–63.
- Lawrence, D.P., Rotondo, F. & Gannibal, P.B. 2016. Biodiversity and taxonomy of the pleomorphic genus *Alternaria*. Mycological Progress 15: 1–22.
- Li, P., Wu, Z., Liu, T. & Wang, Y. 2016. Biodiversity, phylogeny, and antifungal functions of endophytic fungi associated with *Zanthoxylum bungeanum*. International Journal of Molecular Sciences 17: 1541.
- Linaldeddu, B.T., Scanu, B., Maddau, L. & Franceschini, A. 2014. *Diplodia corticola* and Phytophthora cinnamomi: the main pathogens involved in holm oak decline on Caprera Island (Italy). Forest Pathology 44: 191–200.
- Luque. J., Parladé, J. & Pera, J. 2000. Pathogenicity of fungi isolated from *Quercus suber* in Catalonia (NE Spain). Forest Pathology 30: 247–263.
- Meklin, T., Haugland, R.A., Reponen, T., Varma, M., Lummus, Z., Bernstein, D., Wymer, L.J. & Vesper, S.J. 2004. Quantitative PCR analysis of house dust can reveal abnormal mold conditions. Journal of Environmental Monitoring 6: 615–620.
- Mirabolfathy, M. 2013. Outbreak of charcoal disease on Quercus spp. and Zelkova carpinifolia trees in forests of Zagros and Alborz mountains in Iran. Iranian Journal of Plant Pathology 49: 77–79.
- Mirabolfathy, M., Ju, Y.M., Hsieh, H.M. & Rogers, J.D. 2013. *Obolarina persica* sp. nov., associated with dying *Quercus* in Iran. Mycoscience 54: 315–320.

- Mulenko, W., Majewski, T. & Ruszkiewicz-Michalska,
  M. 2008. A Preliminary Checklist of *Micromycetes* in Poland. W. Szafer Institute of Botany. Polish Academy of Sciences 9: 752.
- Ragazzi, A., Moricca, S., Capretti, P., Dellavalle, I. & Turco, E. 2003. Differences in composition of endophytic mycobiota in twigs and leaves of healthy and declining *Quercus* species in Italy. Forest Pathology 33: 31–38.
- Rehner, S.A. & Samuels, G.J. 1995. Molecular systematics of the *Hypocreales*: a teleomorph gene phylogeny and the status of their anamorphs. Canadian Journal of Botany 73: 816–823.
- Rainer, J. & De Hoog, G.S. 2006. Molecular taxonomy and ecology of *Pseudallescheria*, *Petriella* and *Scedosporium prolificans* (*Microascaceae*) containing opportunistic agents on humans. Mycological Research 110: 151–160.
- Sagheb-Talebi, K., Sajedi, T. & Pourhashemi, M. 2014. Forestsof Iran- a treasure from the past, a hope for the future. Springer, Dordrecht Heidelberg, New York, London.
- Simmons, E.G. 1998. Multiplex conidium morphology in species of the *Ulocladium atrum* group. Canadian Journal of Botany 76: 1533–1539.
- Tamura, K., Stecher, G., Peterson, D., Filipski, A. & Kumar, S. 2013. MEGA6: molecular evolutionary genetics analysis version 6.0. Molecular Biology and Evolution 30: 2725–2729.
- Tanaka, K., Endo, M., Hirayama, K., Okane, I., Hosoya, T. & Sato, T. 2011. Phylogeny of *Discosia* and *Seimatosporium*, and introduction of *Adisciso* and *Immersidiscosia* genera nova. Persoonia 26: 85–98.
- Thambugala, K.M., Hyde, K.D., Tanaka, K., Tian, Q., Wanasinghe, D.N., Ariyawansa, H.A., Jayasiri, S.C., Boonmee, S., Camporesi, E., Hashimoto, A. & Hirayama, K. 2015. Towards a natural classification and backbone tree for *Lophiostomataceae*, *Floricolaceae*, and *Amorosiaceae* fam. nov. Fungal Diversity 74: 199–266.

- Thomas, F.M., Blank, R. & Hartmann, G. 2002. Abiotic and biotic factors and their interactions as causes of oak decline in Central Europe. Forest Pathology 32: 277–307.
- Thompson, J.D., Gibson, T.J., Plewniak, F., Jeanmougin, F. & Higgins, D.G. 1997. The CLUSTAL\_X windows interface: flexible strategies for multiple sequence alignment aided by quality analysis tools. Nucleic Acids Research 25: 4876–4882.
- Verkley, G.J.M., Dukik, K., Renfurm, R., Göker, M. & Stielow, J.B. 2014. Novel genera and species of Coniothyrium-like fungi in *Montagnulaceae* (Ascomycota). Persoonia 32: 25–51.
- Vu, D., Groenewald, M., De Vries, M., Gehrmann, T., Stielow, B., Eberhardt, U., Al-Hatmi, A., Groenewald, J.Z., Cardinali, G., Houbraken, J. & Boekhout, T. 2019. Large-scale generation and analysis of filamentous fungal DNA barcodes boosts coverage for kingdom fungi and reveals thresholds for fungal species and higher taxon delimitation. Studies in Mycology 92: 135–154.
- Wanasinghe, D.N., Hyde, K.D., Jeewon, R., Crous, P.W.,
  Wijayawardene, N.N., Jones, E.B.G., Bhat, D.J.,
  Phillips, A.J., Groenewald, J.Z., Dayarathne,
  M.C., Phukhamsakda, C. & Phukhamsakda, C.
  2017. Phylogenetic revision of *Camarosporium* (*Pleosporineae*, *Dothideomycetes*) and allied
  genera. Studies in Mycology 87: 207–256.
- White, T.J., Bruns, T., Lee, S.J.W.T. & Taylor, J.W. 1990. Amplification and Direct Sequencing of Fungal Ribosomal RNA Genes for Phylogenetics. PCR Protocols: A Guide to Methods and Applications 18: 315–322.
- Woudenberg, J.H.C., Groenewald, J.Z., Binder, M. & Crous, P.W. 2013. *Alternaria* redefined. Studies in Mycology 75: 171–212.
- Woudenberg, J.H.C., Groenewald, J.Z., Binder, M. & Crous, P.W. 2013. *Alternaria* redefined. Studies in Mycology 75: 171–212.

- Zhong, S. & Steffenson, B.J. 2001. Virulence and molecular diversity in *Cochliobolus sativus*. Phytopathology 91: 469–476.
  Zhang, Y., Zhang, J., Wang, Z., Fournier, J., Crous, P.W., Zhang, X., Li, W., Ariyawansa, H.A. & Hyde, K.D. 2014. Neotypification and phylogeny
  - of Kalmusia. Phytotaxa 176: 164-173.

Rosth