**Bipolaris spicifera** isolates with unusual conidial germination pattern on sunflower from Iran

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*Bipolaris* isolates with unusual conidial germination pattern were isolated from sunflower leaves in Northern Iran. Conidia germinated to form swollen germ tubes from both ends and swollen germ tubes underwent multiple branching soon after formation. The species was identified as *Bipolaris spicifera* based on phenotypic characteristics as well as sequence data from *gpd* and ITS-rDNA regions. Artificial inoculation of 4-week-old seedlings of sunflower under green house conditions induced chlorotic spots on sunflower leaves. This is first report on the occurrence of *Bipolaris spicifera* on sunflower in Iran. Our results on the unusual germination pattern of *B. spicifera* conidia, further stress the need for the implementation of molecular data for the identification of plant pathogenic *Bipolaris* species.

**Key words** – *gpd* – ITS – leaf spot – sunflower – unusual conidium germination pattern

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**Introduction**
In a survey of leaf-infecting fungi in northern Iran, *Bipolaris spicifera* Shoemaker isolates were recovered from cultivated sunflower (*Helianthus annuus* L.). The genus *Bipolaris* was erected by Shoemaker to accommodate species formerly included in *Helminthosporium* Link (Tsuda & Ueyama 1981). The genus is characterized by generally solitary, distosepate conidia, sympodial proliferation of conidiophores, and bipolar germination of conidia (Alcorn 1983). Until now three *Bipolaris* species viz., *B. spicifera* (Bainier) Subram, *B. sorokiniana* (Sacc.) Shoemaker and *B. zeicola* (Stout) Shoemaker have been reported to occur on sunflower leaves or seeds (Chidambaram et al. 1973, Pandy & Saharan 2005). Herein we document *Bipolaris spicifera* to occure on cultivated sunflower leaves in Iran and further discuss morphology and ecology of this species.

**Methods**

**Isolates and morphology**
Infected leaves of sunflower were collected during August-September 2010 from Salmas, West Azerbaijan, Iran. Fungi were isolated from leaf lesions following surface sterilization in 70% ethanol, and plating on potato carrot agar (PCA; 20 g L\(^{-1}\) potato, 20 g L\(^{-1}\) carrot, 20 g L\(^{-1}\) agar). To obtain pure cultures a mass of conidia were transferred to 2% malt extract agar (MEA; Fluka, Hamburg, Germany) plate containing 10 ml distilled water, supplemented with 100 mg/L streptomycin sulphate and 100 mg/L ampicillin. The resulting spore suspension was streaked on MEA. Conidia were allowed to germinate, and single-spore transfers were made. The cultures
were deposited in the living culture collection of Tabriz University (CCTU), Tabriz, Iran. Description of morphology was made based on 7-day cultures on PCA, exposed to 8/16 h of alternate cool white fluorescent light and darkness at room temperature. Three plates of PCA were produced for cultural and microscopic characterization. Microscopic structures were mounted in lactic acid for measurements. Thirty measurements were made for each microscopic element. Photographs were captured with a Leica camera system.

**Pathogenicity test**

Koch's postulates were conducted in the greenhouse on 4-week-old seedlings. A suspension of inoculum containing $10^5$ conidia ml$^{-1}$ was prepared from 10-day-old cultures. Seedlings were sprayed to runoff with conidial suspension containing 0.04% of Tween 80, and control plants were inoculated with sterile water containing Tween 80. The inoculated seedlings were given a 48 h dew period. The notation of results was made during 10 days after inoculation. Inoculated leaves were detached and cut into small pieces, surface sterilized, and plated on PCA plates for fungal re-isolation.

**DNA phylogeny**

DNA extraction was performed with the protocol of Moller et al. (1992). The 3' end of the 18S rRNA gene, ITS1, 5.8S rDNA, ITS2 and the 5' end of 28S rRNA gene regions were amplified with primers ITS1 and ITS4 (White et al. 1990). PCR was performed on a GeneAmp PCR System 9700 (Applied Biosystems, Foster City, CA). Cycling conditions consisted of 5 min at 95°C, followed by 36 cycles of 30 s at 94°C, 30 s at 52°C and 1 min at 72°C, followed by one cycle of 7 min at 72°C. Glyceraldehyde-3-phosphate dehydrogenase (gpd) gene was amplified with gpd1 and gpd2 primers (Berbee et al. 1999). Amplification reaction mixture contained 1X PCR buffer, 1 mM MgCl$_2$, 60 µl of 1 mM dNTPs, 0.2 pM of each primer, 0.5 U of Taq polymerase, 0.5 µl DSMO, and 10–15 ng of fungal genomic DNA. The final reaction volume was adjusted to 12.5 µl by adding sterile distilled water. The thermal cycling program consisted of a 5 min initial denaturation at 94°C, which was followed by 40 cycles of 30 s denaturation at 94°C, 30 s primer annealing at 57°C, and 30 s extension at 72°C, and 7 min at 72°C for a final extension. DNA sequencing was performed using the BigDye Terminator v3.1 (Applied Biosystems, Foster City, CA) Cycle Sequencing Kits and analyzed on an ABI Prism 3700 (Applied Biosystems, Foster City, CA) according to the recommendation of the vendor. For the species identification, sequence similarity searches were performed for each of the sequences using Megablast search option at NCBI’s GenBank nucleotide database.

**Results**

**Morphological and cultural features**

Colonies (Fig. 2) expanding velvety, brownish grey, with olive green margins, flat, circular, entire, attaining a diam. of 7 mm after 7 days. Hyphae branched, septate, sub-hyaline to pale brown, 4-7 µm diam. Conidiophores arising singly, lateral or terminal, simple or rarely branched, septate, smooth-walled, straight or flexuous, repeatedly geniculate, mid brown, the upper fertile region up to 100 µm long. Conidiogenous cells (Fig. 7) polytretic, integrated, terminal, sympodial, sub-cylindrical, cicatrizied, with several (mostly 6–8) geniculation, dark brown. Conidia (Figs 3, 5–7) acrogenous, straight, sub-cylindrical, smooth walled, tapering towards paler and obtuse ends, 27–34 $\times$ 9–11 µm, typically with 3 and rarely with 6 distosepta, juvenile conidia pale brown, becoming yellow-brown at maturity, hilum conspicuous, the first conidial septum median or sub-median, the second septum delimits basal cell, the third septum distal. Germination bipolar, basal germ tube semi axial, apical germ tube axial, both germ tubes swelling and branching close to the conidium (Fig. 4).

**Pathogenicity test**

In field-grown crops, the fungus was associated with *Alternaria* leaf spot. Symptoms on inoculated leaves appeared 7 days after inoculation, and were characteristically small (up to 4 mm), chlorotic spots, presumably due to toxin production (Fig. 1). The fungus was re-isolated from the inoculated leaves.
Figs 1–7 – Bipolaris spicifera. 1 Symptoms developed on in vitro inoculated sunflower leaves, seven days after inoculation. 2 7-day-old colony on PCA. 3 Conidia. 4 Germinated conidium. 5–7 Conidia and conidiophores. Bars = 10 µm.

DNA phylogeny
Sequence comparisons of ITS-rDNA region showed 100 % identity with known B. spicifera gene sequences available in GenBank (GenBank Accession No. JX070077). There was no sequence data available for gpd gene of B. spicifera in GenBank; hence, the gpd sequence was compared with the sequence data set of Dr Hugo Madrid at CBS-Fungal Biodiversity Centre, Utrecht, The Netherlands. The gpd sequence generated in this study (GenBank Accession No. JX070078) was identical to those of B. spicifera.

Discussion
In 1959 Shoemaker originally proposed the generic name Bipolaris to accommodate some species of Helminthosporium which, unlike Helminthosporium spp., have interminate conidiophores (Goh et al. 1998). The genus is allied with Drechslera, Exserohilum and Curvularia which are, respectively, associated with Pyrenophora, Setosphaeria and Cochliobolus teleomorphs (Goh et al. 1998, Zhang & Berbee 2001). The asexual states of these fungi can be separated based on the following taxonomic criteria: conidium shape, hilum morphology, origin of the basal germ tube and its relative position to the conidial axis, conidial septum sequence, and the delimitation of the basal cell (Alcorn 1983).

Bipolaris and other allied genera encompass considerable morphological similarities. Recently, molecular studies have supported these groups as monophyletic lineages. For instance, phylogenetic relationships among Cochliobolus and related species of Pyrenophora and Setosphaeria have been established based on analysis of the internal transcribed spacer (ITS) and glyceraldehyde-3-phosphate dehydrogenase (gpd) data. This analysis clearly delimited these three genera. Two genera composed a monophyletic Setosphaeria-Cochliobolus clade with Pyrenophora as the sister taxon (Berbee et al. 1999).

Until now over 100 species of Bipolaris have been described (Brecht 2005, Crous et al. 2004). The present species is similar to B.
spicifera in all essential features, but it differs from the type specimen of this fungus in conidial germination pattern; in our isolates conidia germinate to form swollen germ tubes from both ends and swollen germ tube undergo multiple branching soon after formation. The phenomenon of dimorphic germination in B. spicifera suggests that other factors may also operate in mediating this response. For example, it has been suggested that conidial germination type in B. oryzae (Breda de Haan) Shoemaker is culture medium dependent (Dela Paz et al. 2006). However, a phylogeny inferred based on ITS sequence data placed all of B. oryzae isolates with different germination pattern as a single clade (Dela Paz et al. 2006). Our results on the unusual germination pattern of B. spicifera further stress the need for the implementation of molecular data for identification of plant pathogenic fungi, especially Bipolaris species. We compared our isolates with the other morphologically and phylogenetically similar species in this genus such as B. australiensis (Ellis) Tsuda & Ueyama. These two species can be differentiated based on conidial dimensions (Tsuda and Ueyama 1981). B. spicifera is probably heterothallic as no teleomorph was formed in monosporic cultures.

Several species of the genus Bipolaris have been reported on different plant species in Iran (Ershad 2009). A number of species such as B. oryzae, B. sorkiniana (Sacc.) Shoemaker and B. maydis (Nisik. & Miyake) Shoemaker are responsible for economically important diseases on agricultural crops such as rice, barley and maize, respectively. B. spicifera has already been isolated from over 70 host plants, from air and soil (Ahmadpour et al. 2011). Furthermore, the fungus is being recognized with increasing frequency as a causal agent of human disease (Padhye et al. 1986, Eghtedari & Pakshir 2006). To the best of our knowledge, this is the first report of sunflower leaf colonisation by B. spicifera in Iran.

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