Penicillium menonorum, a new species related to P. pimiteouiense

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Abstract: *Penicillium menonorum* is described as a new monoverticillate, non-vegetative species that resembles *P. restrictum* and *P. pimiteouiense*. On the basis of phylogenetic analysis of DNA sequences from four loci, *P. menonorum* occurs in a clade with *P. pimiteouiense*, *P. vinaceum*, *P. guttulosum*, *P. rubidurum*, and *P. parvum*. Genealogical concordance analysis was applied to *P. pimiteouiense* and *P. parvum*, substantiating the phenotypically defined species. The species *P. rubidurum*, *P. guttulosum*, and *P. menonorum* were on distinct branches statistically excluded from inclusion in other species and have distinct phenotypes.

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INTRODUCTION

In the course of a screening program to find useful fungi for conversion of organic matter into high-value products such as lipid precursors to biofuels and animal feed formulations, a *Penicillium* isolated from garden soil in southern California was obtained that could not be placed with confidence in any described species. Sequencing of the ITS region was performed, with sequence analysis showing that this isolate is phylogenetically related to *P. pimiteouiense*. DNA distance from *P. pimiteouiense* suggested that it might be an undescribed species.

Additional gene loci (β-tubulin, calmodulin, and DNA replication licensing factor Mcm7) were amplified and sequenced for this isolate and for phylogenetically and phenotypically similar species. On the basis of the phenotypic and phylogenetic distinctions, this isolate is described as a new species.

MATERIALS AND METHODS

Cultures (Table 1) may be obtained from the Agricultural Research Service Culture Collection (NRRL), Peoria, IL (http://nrrl.ncaur.usda.gov). The *P. menonorum* culture ex-type is available from the Agricultural Research Service Patent Culture Collection (http://nrrl.ncaur.usda.gov). Cultures were maintained on potato-dextrose agar (PDA) during the course of this study. Colony descriptions were based on 7 d growth of cultures on Czapek’s yeast autolysate agar (CYA), malt extract agar (MEA), and glycerol nitrate agar (G25N) at 25 °C, and on CYA at 5 °C and 37 °C as detailed by Pitt (1980). Some color names are taken from Ridgway (1912) and are designated with an upper case R and a plate number.

Microscope slides were made by teasing apart bits of mycelium in a drop of lactic acid with cotton blue. A Zeiss axioscope with DIC optics was used for microscopic observations. Photomicrographs were taken with a Kodak 14n digital camera attached to the microscope. Micro- and macro-photographs were sized and placed in a plate using Adobe Photoshop v. 6.0.1.

Biomass for DNA extraction was grown in 125 mL flasks containing 25 mL malt extract (ME) broth incubated at 25 °C on a rotary platform (200 rpm). Biomass ca. 0.5 g wet weight was collected by vacuum filtration, placed in micro centrifuge tubes, and freeze-dried. Freeze-dried mycelium was ground to a powder with a sterile pipette tip and DNA was extracted from the powdered biomass using the CTAB method. Purified DNA was stored in TE buffer (Tris 10 mM, EDTA 1 mM, pH 8.0) at -20 °C until needed. DNA was amplified using the primers and conditions detailed in Peterson et al. (2010). Amplified DNA was prepared for sequencing using ExoSAP-IT (www.usbweb.com). DNA sequences were produced using DyeDeoxy v. 3.1 reagents and an ABI 3730 DNA sequencer (www.appliedbiosystems.com). Complementary strand sequences were assembled and corrected using Sequencher (www.genecodes.com). Finished sequences were aligned using CLUSTALW (Chenna et al. 2003), and maximum parsimony trees and bootstrap proportions were calculated using PAUP v. 4.0b10 (Swofford 2003). MrBayes v. 3.12 (Huelsenbeck & Ronquist 2001, Ronquist & Huelsenbeck 2003) was used to calculate Bayesian posterior probabilities. DNA sequences used in this
study are deposited in GenBank (www.ncbi.nlm.nih.gov) with accession numbers HQ646566–HQ646603, AF033460–AF033462, AF033464, AF037431, and AF037434. Data sets and tree diagrams are deposited at TREEBASE (www.treebase.org).

The initial search to find phylogenetically related species was performed by BLAST searches of GenBank using the ITS sequence from the new species.

RESULTS

Penicillium menonorum S.W. Peterson sp. nov.
MycoBank MB519297
(Fig. 1A–D)

Etymology: Named for Menon & Associates whose scientists isolated the fungus.

A speciebus alis conidiophoris brevibus, conidiis scaberulis, colore in substrato nutritorio CYA pallide caesio atque augmento in temperatura 37 °C distinguendum.

Typus: USA: California: isolated from garden soil, 2009 (BPI 881018 – holotypus; culture ex-holotype NRRL 50410).

Colonies on CYA (Fig. 1A) attaining 17–20 mm diam after 7 d growth at 25 °C, velutinous-silky, radially sulcate peripherally, centrally raised ca. 2–3 mm, sporulation moderate, central region pale bluish gray (court gray R-47), peripheral area white; no exudate or soluble pigments; no sclerotia or ascomata; reverse yellowish brown centrally (buckthorn brown R-15) to pale brownish-yellow (warm buff, R-15) peripherally. On MEA (Fig. 1B) attaining 17–19 mm diam after 7 d growth at 25 °C, mycelium loosely woven, woolly, umbonate 3–4 mm deep centrally, sporulation moderate, white peripherally, court gray (R-47) centrally; no exudate or soluble pigments; no sclerotia or ascomata; reverse yellowish brown centrally to brownish yellow peripherally. On G25N attaining 8–10 mm diam after 7 d growth at 25 °C, umbonate, woolly 1–2 mm deep, white to court gray; no exudate or soluble pigment; no sclerotia or ascomata; reverse white to buff. Incubation for 7 d on CYA at 5 °C produced no growth or germination of conidia. Incubation for 7 d on CYA at 37 °C produced colonies of 29–32 mm diam, resembling growth on CYA at 25 °C, but clear exudate moderately abundant, the reverse color is a darker, more uniform shade of brown. Conidiophores (Fig. 1C) smooth-walled, hyaline, 5–15(–20) × 1.5–2.0 µm, non-vesiculate, with an apical whorl of (1–)2–5 phialides 5–7(–9) × 2.5–3.5 µm, conidia spherical to subspherical, (2–)2.5–3.5 µm (Fig. 1D), with roughened to rugose surface.

Table 1. Provenance of isolates used in this study.

| Species                  | NRRL Accession No. | Origin                                                                 |
|--------------------------|--------------------|-------------------------------------------------------------------------|
| Penicillium erubescens   | MB335726           | South Africa: Pretoria: isolated from nursery soil, 1967, culture ex-type |
| Penicillium guttosum     | MB266689           | USA: Utah: isolated from soil, 1927, culture ex-type                     |
| Penicillium menonorum    | MB519297           | USA: California: isolated from garden soil, 2009, culture ex-type        |
| Penicillium parvum       | MB289101 (syn. E. parvum) | Papua-New Guinea: isolated from soil, ca. 1973, S. Udagawa, culture ex-type of P. papuanum MB319290 |
| Penicillium pimiteouensi | MB460126           | New Guinea: isolated from tent cloth, ca. 1944, G.W. Martin              |
| Penicillium rubidurum    | MB319295 (syn. E. rubidurum) | USA: Illinois: Peoria: isolated from human kidney cell culture plate, April 1996, J.T. Hjelle, culture ex-type |
| Penicillium vinaceum     | MB281754           | USA: Utah: isolated from soil, 1927, culture ex-type                     |
| Penicillium pimiteouensi | MB460126           | Unknown: obtained from M.B. Morrow, 1936                                 |

*MB=MycoBank (http://www.mycobank.org/).
DNA sequences from the β-tubulin locus included all or part of 4 exon and 4 intron regions. After alignment the data set included 703 base positions. The calmodulin data included all or part of 4 exon and 3 intron regions and aligned with 726 base positions. The ID regions included the ITS1, ITS2, 5.8S rDNA, and ca. 650 bases from the 28S rDNA in an alignment of 1141 bases. DNA replication licensing protein (Mcm7) was composed of an amino acid coding region of 616 bp length. Penicillium erubescens was chosen as the outgroup on the basis of phylogenetic trees previously published (Peterson et al. 1999, Peterson 2000).

The most parsimonious trees, bootstrap proportion and Bayesian posterior probabilities for individual data sets were determined and the trees were compared for strongly supported contradictory branch points. Strongly supported nodes are those with > 90 % of the bootstrap sample and a Bayesian posterior probability of > 0.90. The individual locus trees contained no strongly supported contradictions that would preclude combining the data. The data from the four loci were combined to calculate a single phylogenetic tree (Fig. 2).

The five isolates of P. pimiteouiense occur on a single strongly supported branch; three isolates of P. parvum and the single isolate of P. papuanum occur on a different strongly supported branch, and the two P. vinaceum isolates occur on another strongly supported branch. Penicillium rubidurum and P. guttolusosum are most closely related to each other and form a sibling group to P. pimiteouiense, while P. menonorum is positioned basal in the tree to this three species branch.

**DISCUSSION**

*Penicillium menonorum* is similar phenotypically to *P. pimiteouiense, P. restrictum, P. striatisporum, P. vinaceum, P. rubidurum, P. erubescens, and P. parvum. Penicillium restrictum, P. malacaense, P. kurssanovii, P. griseolum, and P. striatisporum*, which phenotypically resemble *P. menonorum,* are phylogenetically positioned in different clades (Peterson & Horn 2009). Other species bearing some resemblance to *P. menonorum* are either not represented by extant extype cultures or the type cultures are not readily available. *Penicillium menonorum* differs from *P. pimiteouiense* by producing conidiophores in a basal layer rather than from aerial hyphae and a bluish gray (Court gray R-47) color on CYA versus white in *P. pimiteouiense.* Additionally, *P. pimiteouiense* produces yellow exudate and a brown soluble pigment, neither of which appear in *P. menonorum* after 7 d incubation. On different media (e.g., yeast extract malt agar incubated at 25 °C) or after extended incubation, a clear to rosy exudate often appears in *P. menonorum.* *Penicillium restrictum* produces somewhat longer conidiophores (up to 60 µm) and has smaller colonies (< 10 mm diam) at 37 °C than *P. menonorum* (29–32 mm diam). *Penicillium striatisporum* produces rosy colored colonies on Czapek’s agar and has striate conidia. *Penicillium vinaceum* produces copious exudate in yellow to vinaceous colors, yellow to brown soluble pigments, and a dark brown colony reverse on CYA, and colonies grown at 37 °C are somewhat smaller (8–20 mm diam) than those of *P. menonorum. Penicillium parvum* typically has mycelium that varies from white to yellow to red in color, while the *P. menonorum* mycelium is uniformly white. *Penicillium parvum* usually makes brown or purple-brown exudate, a brown soluble pigment, and has a colony reverse that is deep reddish-brown versus *P. menonorum,* which has no exudate or soluble pigments and a yellow brown colony reverse after 7 d incubation. *Penicillium rubidurum* produces white to orange or rosy-buff mycelium, red-brown exudate, a dark brown colony reverse, and produces conidia on M40Y medium but not on CYA. *Penicillium menonorum* produces no exudate or soluble pigment and has a yellow brown reverse and has abundant conidiogenesis on CYA. *Penicillium erubescens* produces white, pink or flesh color mycelium, reddish-brown exudate, and gray-red to magenta to vinaceous purple soluble pigments, with colony reverse either similarly colored or brown. Each of these species is...
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Ima funGus easily distinguished from *P. menonorum* on these bases. Raper & Thom (1949) regarded *P. guttulosum* to be a synonym of *P. janthinellum*, differing primarily by the production of copious amounts of exudate. *Penicillium guttulosum* as represented by Gilman & Abbott’s ex-type strain is distinct from *P. janthinellum* as well as the species studied here. *Penicillium guttulosum* cultures on CYA resemble the cultures of *P. vinaceum*, differing most noticeably in the production of dark purple exudate in large quantities, while *P. vinaceum* exudate is more red in color. *Penicillium rubidum* colonies also resemble *P. vinaceum* and *P. guttulosum* but produce pale yellow exudate. Pitt (1980) treated *P. papuanum* as a

Combined data from BT2, CF, ID and Mcm7 loci, 3186 total characters, 2798 are constant, 204 are variable not parsimony informative, 200 are parsimony informative; 1 mpt, CI=0.8348, RC=0.7190

**Fig. 2.** Phylogenetic tree calculated using maximum parsimony criterion for the concatenated data set composed of beta-tubulin, calmodulin, ITS and 28S rDNA, and DNA replication licensing protein (Mcm7). Bootstrap proportions/Bayesian posterior probabilities are placed on internodes.
synonym of *P. parvum* and they are in the same strongly supported clade (Fig. 2). Phenotypically, they are very similar to each other. Additional isolates of each species are needed to further assess the phylogenetic and phenotypic distinctions of these species.

Phylogenetic systematics (Hennig 1966) is based on the principle that species must be monophyletic. Taylor et al. (2000) presented the genealogical concordance phylogenetic species recognition (GCPSR) concept as a means of determining the boundaries of species in fungi. Dettman et al. (2006) showed experimentally that GCPSR is effective in recognizing species boundaries in the genus *Neurospora*. GCPSR can be applied to *P. pimiteouiense* and *P. parvum* in this study and the species are supported by the GCPSR principles. *Penicillium vinaceum*, *P. guttulosum*, *P. rubidurum*, and *P. menonorum* are each on distinct branches, but the boundaries of the species cannot be determined from the single isolates available here. Phenotypic distinctions make each of these species recognizable and the phylogenetic placement of the species is consistent with the phenotypic descriptions of the species.

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