Muscodor cinnamomi, a new endophytic species from Cinnamomum bejolghota

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Abstract — Muscodor cinnamomi is described as a new species, endophytic within leaf tissues of Cinnamomum bejolghota (Lauraceae) in Doi Suthep-Pui National Park, Northern Thailand. Molecular analysis indicated differences from the five previously described Muscodor spp. Volatile organic compounds analysis showed that M. cinnamomi produced azulene (differentiating it from M. crispans) but did not produce naphthalene (differentiating it from M. albus, M. roseus, and M. vitigenus).

Key words — sterile ascomycete, cinnamon, endophytes, volatile compounds

Introduction

Plants are reservoirs of untold numbers of endophytic organisms (Bacon & White 2000). By definition, these microorganisms (mostly fungi and bacteria) reside in the tissues beneath the epidermal cell layer and cause no apparent harm to the host (Azevedo et al. 2000, Hyde & Soytong 2008). Endophytes from rainforest and medicinal plants have been studied for their volatile antibiotic and other medicinal characteristics (Strobel et al. 2003, Huang et al. 2008, 2009, Mitchell et al. 2008, Tejesvi et al. 2009, Aly et al. 2010). Five endophytes characterized by sterile mycelium that have recently been described as novel fungi are Muscodor albus isolated from Cinnamomum zeylanicum (Lauraceae) in Honduras (Worapong et al. 2001), M. roseus from Grevillea pteridifolia (Proteaceae) in the Northern Territory of Australia (Worapong et al. 2002), M. vitigenus from Paullinia paullinioides (Sapindaceae) in Lake Sandoval (Daisy et al. 2002), M. crispans from Ananas ananassoides (Bromeliaceae) in the Bolivian Amazon (Mitchell et al. 2008), and M. yucatanensis from Bursera
**Fungal isolation**

Ten healthy leaves of *Cinnamomum bejolghota* were collected from plants growing in Doi Suthep-Pui National Park, Northern Thailand (alt. 950 m) during May 2008. Totally, 250 tissue squares (5 x 5 mm) were cut from the leaf samples. All leaf tissues squares were surface sterilized in 75% ethanol for 30 s, 2% sodium hypochlorite for 3 min and 95% ethanol for 30 s under a laminar flow hood (Nuangmek et al. 2008). The sterilized samples were placed in Petri dishes containing 2% malt extract agar, 0.05% streptomycin sulfate and 0.03% rose bengal (Bussaban et al. 2001). Petri dishes were sealed with Parafilm and incubated at room temperature (25±2°C) for one week. The fungi growing out from the samples were aseptically transferred to two culture media, potato dextrose agar (PDA) and malt extract agar (MA); pure isolates were maintained in corn meal agar (CMA) slants. Various methods were tried to stimulate spore production (Guo et al. 1998).

**Scanning electron microscopy**

Scanning electron microscopy was preformed on isolate CMU-Cib 461 following procedures described by Castillo et al. (2005). A piece of agar with fungus was placed in a filter paper packet and then placed in 2% glutaraldehyde vapor, a wetting agent, and aspirated over night. Samples were then dehydrated in an ethanol series (15 mins at 5, 10, 15, 20, 40, 50, 70, 80, 95 and 100%) and in an acetone series (10 mins at 10, 15, 20, 40, 50, 70, 80, 95 and 100%). The fungal material was critically point dried, gold sputter coated, and images observed under a JEOL JSM-5910LV SEM using a high vacuum mode.

**Qualitative analysis of CMU-Cib 461 volatiles**

CMU-Cib 461 was grown in 5 ml Agilent® clear glass vials containing PDA for 10 days at room temperature (25±2°C). Volatile compounds produced by the fungus were analyzed on an automatic Agilent Technologis GC 7890 gas chromatograph column.
containing a HP-5MS 30 m × 0.25 mm I.D. × 0.25 μm. The column was temperature programmed as follows: 32°C for 2 min followed to 220°C at a rate of 5°C/min. The carrier gas was ultra high purity helium released at a rate of 1.5 mL/min. Prior to trapping the volatiles, the fiber was conditioned at 250°C for 39.6 min under a flow of helium gas. The gas chromatograph was interfaced to a MSD 5973 (EI) mass selective detector (mass spectrometer) operating at unit resolution. Acquisition and processing data were performed on the MSD 5973 (EI) software system. Initial identification of the volatile compounds produced by CMU-Cib 461 was made through library comparison using the NIST database, and compared with the original isolates, M. albus strain 620 (Strobel 2006) and strain E-6 (Strobel et al. 2007).

Fungal cultures and DNA extraction
Genomic DNA was extracted by a modified SDS-CTAB method (Bussaban et al. 2005). Strain CMU-Cib 461, isolated from C. bejolghota leaves, was subcultured onto PDA and incubated for 10 days. Mycelium was harvested, freeze dried, and ground into a fine powder with a pestle and mortar. About 15 mg of powdered mycelium was suspended in 1 mL of ice-cold lysis buffer (150 mM NaCl, 50 mM EDTA, 10 mM Tris-HCl, pH 7.4, 20 mg/mL proteinase K), transferred into 1.5 mL Eppendorf tube and kept at 4°C to prevent endonuclease activity during rehydration of the sample. SDS was added to a final concentration of 2%, vortexed and incubated 30 min at 65°C. After centrifugation for 15 min at 14,000 rpm, the supernatant was transferred to a new sterile 1.5 mL Eppendorf tube. The volume of supernatant was measured and the NaCl concentration was adjusted to 1.4 M, and one-tenth volume of 10% CTAB buffer (10% CTAB, 500 mM Tris-HCl, 100 mM EDTA, pH 8.0) was added. The solution was thoroughly mixed and incubated for 10 min at 65°C. After cooling for 2 min at 15°C, an equal volume of chloroform: isoamyl alcohol (24:1 v/v) was added, thoroughly mixed and the tube was centrifuged 15 min at 14,000 rpm. The extraction was repeated until the interface was clear. The supernatant was removed to a new Eppendorf tube, containing 2 volumes of cold 100% ethanol. After DNA precipitation, the pellet was centrifuged for 15 min at 14,000 rpm and 4°C. The pellet was washed with 70% ethanol and dried at room temperature. It was resuspended in 100 mL of 0.002% RNase (5 mg/mL) in TE buffer and incubated for 1 h at 37°C. The suspension was stored at -20°C pending use for PCR amplification.

Fungal ITS regions sequencing and phylogenetic analysis
The internal transcribed spacer (ITS) regions 1 and 2, including 5.8S rDNA were separately amplified in a 25 mL reaction on a GeneAmp 9700 thermal cycler (Applied Biosystems) under these reaction conditions: 1 mL of template DNA extraction, 0.2 mM dNTP, 0.2 mL of FastTaq (Applied Biosystems), 0.2 mM each of primers, 2.5 mL of the supplied 103 PCR buffer with MgCl₂, and sterile water to bring the volume to 25 mL. The ITS regions were amplified by using ITS4 and ITS5 primers. Amplification of ITS regions was for 30 cycles (initial denaturation at 95°C for 2 min, denaturation at 95°C for 30 s, annealing at 50°C for 30 s, and extension at 72°C for 1 min, with a final extension at 72°C for 10 min). PCR products were analyzed by electrophoresis in 1% agarose gels in TAE buffer (20 mM Tris-Acetate, 1 mM EDTA, pH 8.0) and viewed by staining with ethidium bromide. PCR products were purified using PCR clean up Gel extraction
NucleoSpin® Extract II purification Kit (Macherey-Nagel, Germany) following the manufacturer’s protocol. The purified PCR products were directly sequenced. Sequencing reactions were performed and sequences determined automatically in a genetic analyzer (1st Base, Malaysia) using the PCR primers mentioned above. Sequences obtained in this study were compared to those from GenBank database using the BLAST software on the NCBI website: (http://www.ncbi.nlm.nih.gov/BLAST/). After multiple alignment of selected sequencer with Clustal X. Phylogenetic trees were constructed using the PUAP beta 10 software version 4.0 (Swofford 2002).
Results

**Taxonomic description**

*Muscodor cinnamomi* Suwannarach, K.D. Hyde & Lumyong, sp. nov. Figs 1–5

MycoBank # MB518008, GenBank # GQ848369

_Fungus in natura cum Cinnamomi bejolghota consociatus et est deuteromycete mycelii sterilibus pertinens. Coloniae fungales est luteus in vitro examinati in loco cum sol lux. Sporae velcorpora fructificantia substatibus ullis non observata. Hyphae (0.9–5.2 μm) vulgo ramificantes et convolventes, fila stripformia et spiras perfectas (4.5–12 μm) formantes. In vitro examiniter corpores colifloriform (6.3–14 μm) e repletus forma hyphae._

Etymology: _cinnamomi_, from the name of the host plant.

Holotype: Thailand, Doi Suthep-Pui National Park; from a leaf of *Cinnamomum bejolghota* (Lauraceae), May 2008, Nakarin Suwannarach; holotype – dried culture, SDBR CMU-Cib 461. (Living culture, BCC38842).

Teleomorph: Unknown.

In nature, the fungus is associated with *Cinnamomum bejolghota* and it is an ascomycete with sterile mycelium. Fungal colonies whitish on all media (PDA, MA and CMA) when grown in darkness (Fig. 1), pale orange when grown in natural light. Hyphae (0.9–5.2 μm thick) commonly appearing as fused rope-like strands, branching (Fig. 4); with coils (4.5–12 μm diam.; Figs 2, 3) and cauliflower-like bodies (6.3–14 μm; Fig. 5). Mycelium on PDA reaching 9 cm in 2–3 weeks and producing a fruity odor. Spores and other fruiting bodies did not develop under any conditions tested.

**Molecular phylogeny of Muscodor cinnamomi CMU-Cib 461**

Partial ITS1 5.8 ITS2 rDNA sequences of *M. cinnamomi* were obtained and compared with GenBank database. After searching the ITS-5.8S rDNA sequences, 635 bp of *M. cinnamomi* (GQ848369) was subjected to an advanced BLAST search. The ITS1 5.8S ITS2 rDNA sequences of *M. cinnamomi* blasted five type strains of *Muscodor* species. The result showed that there was a 99, 99, 99, 98 and 90% similarity with *M. albus* (AF324336), *M. roseus* (AY034665), *M. crispans* (EU195297), *M. vitigenus* (AY100022) and *M. yucatanensis* (FJ917287), respectively.

Parsimony analysis of the alignment yielded 100 most parsimonious trees with total length of 873 steps (CI = 0.705, RI = 0.746, RC = 0.526, HI = 0.294), one of which is shown in Fig. 6. *Muscodor cinnamomi* and *Muscodor* species from GenBank formed a monophyletic clade (clade I) with a high bootstrap support (99%), and formed a sister group to *Anthostomella* (clade II) with 83% bootstrap support. *Muscodor* species are more closely related to the *Xylariaceae* than *Amphisphaeriaceae* with 100% bootstrap support.

**Volatile compounds from M. cinnamomi (CMU-Cib 461)**

*Muscodor cinnamomi* (CMU-Cib 461) produced at least 11 volatile compounds. These could be positively identified on the basis of a GC/MS
comparison with authentic standards obtained from commercial sources as well as organic synthesis. The compounds were identified primarily on the basis of their mass spectral properties when compared to the NIST database. Of the compounds produced by this organism the most abundant were propanoic acid, 2-methyl, methyl ester, butanoic acid, 2-methyl, methyl ester and cis-2,4-dimethylthiane,S,S-dioxide with total area higher than 10% (Table 1). A number of other volatiles appeared that were unique to this isolate, including cis-2,4-dimethylthiane,S,S-dioxide; β-humolene; cyclopentane; eudesma4(14),11-diene and 1,1,1,5,7,7,7-heptamethyl-3,3-bis(trimethylsiloxy) tetrasiloxane compounds. In addition, the fungus produced azulene, but no naphthalene compounds.
Table 1. GC/MS analysis of the volatile compounds produced by *Muscodor cinnamomi* (CMU-Cib 461) culture in 5.0 mL clear glass vial Agilent® for 10 days.

<table>
<thead>
<tr>
<th>RT (mins)</th>
<th>Total area (%)</th>
<th>Analysis compound</th>
<th>M/z</th>
</tr>
</thead>
<tbody>
<tr>
<td>3:15</td>
<td>1.10</td>
<td>(S)-(+) 5-methyl-1-heptanol</td>
<td>130</td>
</tr>
<tr>
<td>3:32</td>
<td>5.49</td>
<td>ethyl acetate</td>
<td>88</td>
</tr>
<tr>
<td>4:35</td>
<td>32.26</td>
<td>propanoic acid,2-methyl,methyl ester</td>
<td>102</td>
</tr>
<tr>
<td>5:38</td>
<td>11.35</td>
<td>cis-2,4-dimethylthiane,S,S-dioxide*</td>
<td>162</td>
</tr>
<tr>
<td>5:41</td>
<td>7.69</td>
<td>cyclopentane*</td>
<td>70</td>
</tr>
<tr>
<td>6:38</td>
<td>14.90</td>
<td>butanoic acid,2-methyl,methyl ester</td>
<td>116</td>
</tr>
<tr>
<td>9:29</td>
<td>3.12</td>
<td>1-butanol,3-methyl,acetate</td>
<td>130</td>
</tr>
<tr>
<td>27:42</td>
<td>3.23</td>
<td>β-humulene*</td>
<td>204</td>
</tr>
<tr>
<td>30:89</td>
<td>8.58</td>
<td>azulene,1,2,3,5,6,7,8,8a-octahydro-1,4-dimethyl-7-(1-methylethenyl)-.[1S-(1α, 7α, 8a,β)]</td>
<td>204</td>
</tr>
<tr>
<td>30:90</td>
<td>7.32</td>
<td>Eudosma-4(14),11-diene*</td>
<td>107</td>
</tr>
<tr>
<td>34:48</td>
<td>2.66</td>
<td>1,1,1,5,7,7,7-heptamethyl-3,3-bis (trimethylsiloxy) tetrasiloxane</td>
<td>444</td>
</tr>
</tbody>
</table>

Abbreviations: * = compounds found in *M. cinnamomi* but not in other *Muscodor* species; RT = retention time; M/z = mass to charge ratio.

**Discussion**

*Muscodor cinnamomi* is introduced as a new species based on differences in colony characteristics, growth rate, ITS sequence data and volatile compounds produced. *Muscodor cinnamomi* (CMU-Cib 461) produced a white mycelium on a PDA. Spores or fruiting structures did not develop on any media including ones containing the host plant material, cinnamon leaves. In this respect it is similar to other *Muscodor* species. The hyphae tend to intertwine to form rope-like strands. Other species of *Muscodor* also have this tendency (Worapong et al. 2001, 2002). The fungus also produces cauliflower-like structures, which is similar to *M. crispans*. The features of *M. cinnamomi* (CMU-Cib 461) are similar to *M. albus*, *M. crispans*, *M. vitigenus* and *M. yucatanensis* which produce whitish mycelium on all media tested in artificial light (Worapong et al. 2001, Daisy et al. 2002, Mitchell et al. 2008, González et al. 2009). *Muscodor cinnamomi* developed a pale orange coloured mycelium in natural light, while *M. crispans* produces a pale pink mycelium in natural light (Mitchell et al. 2008). Phylogenetic analysis of the sequences of ITS1, 5.8S, and ITS2 showed that *M. cinnamomi* was closely related the other *Muscodor* species, which are related to family Xylariaceae (Worapong et al. 2001, 2002).

When measured by GC/MS, the fungus consistently produced alcohols, esters and small molecular weight acids, in the gas phase, when grown on PDA. *Muscodor cinnamomi* produces propanoic acid,2-methyl,methyl ester, which is similar to other *Muscodor* species. However, there are differences in other
compounds produced by the different *Muscodor* species (Table 2). The volatile compounds showed inhibition ability and lethal activity against a number of plant and human pathogens (Strobel et al. 2001, Worapong & Strobel 2009). Details on the bioactivities of this interesting genus appear elsewhere (Worapong et al. 2001, 2002, Daisy et al. 2002, Ezra et al. 2004, Strobel 2006, Strobel et al. 2007, Mitchell et al. 2008). The strain CMU-Cib 461 shared all of the common features of previously described *Muscodor* species but there were a number of different aspects to the taxon that distinguished it from other *Muscodor* species.

**Acknowledgments**

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