Cultivation and identification of blazeispiros of *Agaricus campestroides* and *A. flocculosipes*

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Abstract

The genus *Agaricus* contains many edible and medicinal mushrooms; and some of them are used for commercial purposes. Most *Agaricus* species have been described from temperate geographic areas, but their occurrence in the tropics is still underexplored. The present study describes the successful cultivation of two wild strains from Thailand, *A. flocculosipes* and *A. campestroides* for the first time. The optimal conditions for mycelial growth have been evaluated for both strains, formation of basidiomata on compost could only be observed for *A. campestroides*. In the present study, mycelial growth and mushroom production conditions were optimized. The temperature range of 25–30 °C with 80–90% humidity was assessed as preferred conditions for primordia and basidiomata formation in a compost rice straw medium with sandy-soil casing layer. Blazeispiros (blazeispirol A and D), a family of triterpenoids that is known to act as selective agonists of Liver X receptor alpha with beneficial effects in vivo in a murine model, were detected in crude extracts from submerged cultures of *A. campestroides* and *A. flocculosipes*.

Keywords – Bioactive compounds – edible mushroom – mushroom farming – Thailand

Introduction

*Agaricus* L. (Linnaeus, 1753) is a genus of agaricoid fungi that was erected quite early in mycological history and even today it still forms the core of the *Agaricaeae* (*Basidiomycota*). Approximately 500 species have been described in this genus, and they are quite common in various habitats, such as grassland and forests (Callac & Chen 2018, He et al. 2019). The genus *Agaricus* is characterized by basidiomata having a fleshy pileus with free lamellae that produce brown spores, and an annulate stipe. *Agaricus bisporus* (J.E. Lange) Imbach is actually the most extensively cultivated mushroom in the world, accounting for 38% of world’s mushroom production (ISMS Edible mushrooms 2017, http://www.isms.biz/edible-mushrooms/). For *Agaricus* cultivation, many researchers rely on *A. bisporus* using the local agro-industrial wastes as substrates. The raw materials that are commonly used to prepare the substrate are sugarcane bagasse, various grasses, cereal straw (*Triticum aestivum*, *Avena sativa*, *Oryza sativa*) and manures supplemented with nitrogen sources (soybean, wheat, corn, cottonseed meal, urea, and ammonium...
sulfate) with sources of phosphorus and calcium (Zied et al. 2011a,b). Experiments performed in China showed the possible use of cottonseed hulls, rice hulls, asparagus straw and soybean cake for compost (Wang et al. 2010, Zhou et al. 2010). Cattle bedding compost/sawdust/cereal bran (Pokhrel & Ohga 2007) and chicken manure/wheat straw (Gregori et al. 2008) were also used for *Agaricus* production in temperate countries. Peat, local soils, with or without the addition of vegetal charcoal, have been used in the casing layer for cultivation (Cavalcante et al. 2008, Siqueira et al. 2009, Colauto et al. 2010, Zied et al. 2010)

Another popular edible mushroom, *Agaricus subrufescens* Peck was cultivated first in the late 1800s in eastern North America (Kerrigan 2005). From a nomenclatural point of view, taxonomists agree that the species has been incorrectly referred as *A. blazei* Murrill. Therefore, it was proposed by Wasser et al. (2002) as a new species, *Agaricus brasiliensis* Wasser, Didukh, de Amazonas & Stamets, and then synonymised by Kerrigan (2005) with *A. subrufescens* which has priority since it is older. The type consists partly of cultivated material and partly of field-collected specimens and has been produced on a commercial scale in Brazil since the early 1990s (Kerrigan 2005, Braga et al. 2006) and exported to several countries. Nowadays it is also cultivated at industrial level in China, France, Japan, Korea, Taiwan and USA. *Agaricus flocculosipes* was first described from Thailand, but it is also known from Mayotte Island (Zhao et al. 2012) and China (Gui et al. 2015). The cultivation of *A. flocculosipes* and *A. subrufescens* in Thailand were successfully cultivated on standard compost based on wheat straw and horse manure but the yields were low (Thongklang et al. 2014). Based on our current information, 22 mushrooms were successfully cultivated in Thailand for commercial use (Thawthong et al. 2014). Several hybrid strains of *Agaricus* from Thailand have been developed, focusing on their favourable agronomic behaviour and morphological characteristics (Jatuwong et al. 2014).

Species of *Agaricus* have high potential for cultivation as many species are edible and possess medicinal properties. A substance class already known from the genus *Agaricus* are the blazeispirols which were isolated from cultured mycelium of *A. blazei* (= *A. subrufescens*) (Hirotani et al. 1999, 2000, 2001, 2002). Moreover, Studler et al. (2005) reported a series of triglycerides of chlorinated phenols from mycelial part of liquid cultures of *Agaricus macrosporus* and several other species in the genus *Agaricus*. The aromatic metabolites showed potential analgesic effects, owing to their strong inhibitory activity against neurolysin. The blazeispirols from *A. subrufescens* were discovered as strong and selective agonists of the Liver X receptor (LXR). Significant in vivo effects of blazeispirols in a mouse model were observed which might give rise to the development of a new anti-hypercholesterolemic and antidiabetic agent from cultures of a medicinal mushroom (Grothe et al. 2011).

In this study, *Agaricus campestroides* and *A. flocculosipes* were selected for cultivation. *Agaricus campestroides* belongs to “Tropical clade b” sensu stricto (Zhao et al. 2011) is known as wild edible species from Ethiopia (Dejene et al. 2017). *Agaricus flocculosipes* is also an edible species belongs to section *Arvenses* (Zhao et al. 2012). Those mushrooms are potentially cultivable and might be a promising source with potential commercial value for Thailand and other countries. We aim in this study at finding optimal conditions for mycelial growth and the spawn production for mushroom cultivation that can further used for industrial production. The present study also describes the identification of blazeispirols in liquid cultures of *Agaricus campestroides* and *A. flocculosipes* which were collected from Thailand.

**Materials & methods**

**Sample collection of *Agaricus* strains**

Two specimens of *A. campestroides* and *A. flocculosipes* were collected in Mae Fah Luang university campus, Chiang Rai province, Northern Thailand during the rainy season from June to August 2014. The specimens were dried at 45 °C by hot air dryer and kept in zip-lock plastic bags containing dehydrated silica gel as a desiccant to control humidity. The dried specimens were deposited in the herbarium of Mae Fah Luang University (MFLU herbarium), Chiang Rai, Thailand.
with the accession numbers: MFLU 14–0872 (A. campestroides) and MFLU 14–0874 (A. flocculosipes).

Isolation of mycelial cultures

Pure cultures were aseptically obtained by transferring sections of internal tissue from wild basidiomata of Agaricus onto potato dextrose agar (PDA) after incubation at 25 °C in a dark room for 14 days. The isolates were deposited in the culture collections of Mae Fah Luang University (MFLUCC) with the numbers MFLUCC 14–1024 (MFLU 14–0872) and MFLUCC 14–1026 (MFLU 14–0874). The cultures were kept at 4 °C, 25 °C, and -20 °C for further studies.

DNA extraction, PCR, and sequencing

Genomic DNA was extracted from dry specimens (ca. 10 mg) using a CTAB isolation procedure adapted from Doyle & Doyle (1990). The ITS region was amplified by polymerase chain reaction (PCR). For PCR amplification of the ITS1–5.8S–ITS2 region of rDNA, ITS1-F (Gardes & Bruns 1993) and ITS4 (White et al. 1990) were used as primers. Raspé et al. (2016) and Chuankid et al. (2019) were followed for the PCR amplification, purification and sequencing of ITS region. The sequences of Thai strains of A. campestroides and A. flocculosipes were deposited in the NCBI database under the accession numbers MZ596239 and MZ596240, respectively.

Optimization of culture conditions for mycelial growth

Four different media were used for optimizing mycelial growth rates, including malt extract agar (MEA), malt yeast peptone agar (MYP), potato dextrose agar (PDA), and potato dextrose yeast agar (PDYA), and pH of the media were adjusted to 6.3. After incubation for 10 days, the growing edge of each colony from pure cultures on PDA was cut out by using a cork-borer (8 mm in diameter) and placed on the centre of each optimization medium in 9 cm petri dishes. Three replicates of each medium were incubated in the dark at 25 °C for 30 days. Mycelial characteristics, such as colour, margin, and shape on the agar surface were recorded. Biomass of dry mycelium was measured by melting agar medium, drained away the liquid phase of medium, and dried the mycelia at 40 °C for 24 h. The mycelial dried weight was recorded using an electronic scale (gram). The biomass was measured every three days for 30 days. The day after mycelium fully colonised was selected for statistical analysis.

Mycelial disks (8 mm) from the colony edge of 10-day-old petri dish cultures of each strain were transferred to optimal medium. The optimal temperature for mycelial growth was determined by using four different temperatures: 20, 25, 30 and 35 °C. Three replicates of each strain were incubated in a dark room for 30 days. The biomass was measured as described above.

The medium exhibiting the highest growth rate and optimal temperature was used to evaluate the optimal pH. The pH was adjusted to 5, 6, 7, and 8 with hydrochloric acid (HCl) and sodium hydroxide (NaOH). The pH range of the media were measured using a digital pH meter before autoclaving. The biomass of mycelium was measured for each pH value as described above.

Spawn production

Spawn is the medium for mushroom mycelium that provides the backbone to any mushroom growing operation (seeds for cultivation). The preparation of spawn was adapted from Royse & Beelman (2016). Spawn was made by sterilized a mixture of wheat grain mixed with water, 1% calcium carbonate, and 0.5% calcium sulfate. The spawn jars containing 100 g of wheat grain were inoculated with a 1/4 colony from actively growing mycelia of a 9 mm diameter petri dish and incubated at 25 °C until mycelium fully colonised.

Cultivation substrate

The compost was prepared using the outdoor composting method, including 25 days of phase I and 7 days of phase II, totalling 32 days. The formulation used had a dry weight of 1,080 kg of Oryza sativa, 5.5% of rice bran, 1.1% of urea, 2.2% calcium oxide, and 2.2% gypsum. The bulk
materials (O. sativa straw and rice bran) were moistened for 9 days and rotated every 2 days. The concentrated materials (urea, calcium oxide, and gypsum) were added after each turning operation throughout the composting phase I (Table 1).

The casing mixture used contains 15% peat, 23% limestone, 37% fine sand, and 25% soil, as recommended by Zied et al. (2011a). When the mycelia were fully developed, the compost was pressed and levelled to facilitate the addition of the casing layer to a height of 5 cm. After casing, the trays were watered once a day. The mushroom production was carried out with four replicates (Royse 2010, Llarena-Hernández et al. 2011).

Conditions for fructification

Four trays filled with 5 kg of compost were inoculated with 5% spawn and incubated for 30 days in a grow room at 25 °C with 90% relative humidity. Then 5 cm of casing layer was added, and the trays were left under the same environmental conditions for a 7-day post-incubation period. The grow room was then set at 20 °C with 80% humidity. Time for mushroom forming was the number of days between casing and the first picking. The numbers and fresh weight of basidiomata were recorded every 5 days after casing. The experiments were performed according to a completely randomized design with four replicates per strain.

Investigation of the secondary metabolite production

The production of secondary metabolites in submerged cultures was examined in strain A. campestroides (MFLUCC 14-1024) and A. flocculosipes (MFLUCC 14-1026) using four different media: yeast malt extract medium (YM), sugar malt extract medium (ZM), cotton seed meal medium (Q6 ½), and mannitol salt medium (MMK). The mycelium was inoculated in 100 mL of each medium in 250-mL Erlenmeyer flasks at 24 °C and placed on a 140-rpm rotary shaker. After 5 days following the inoculation, the free glucose was measured with glucose test strips daily until the free glucose was consumed and the pH was checked with a pH meter (method adapted from Kuhnt et al. 2014). To qualitatively analyse the produced metabolites, an ethyl acetate extraction procedure was used for mycelia and submerged culture supernatant. Liquid media and biomass from liquid cultivation on different media were separated by filtration. Liquid media were discarded due to the lack of compounds. Each mycelium was extracted with acetone (200 mL) in an ultrasonic bath at room temperature for 30 min. After, the crude extract was filtered and redissolved in deionized water (100 mL), then partitioned with ethyl acetate (1 × 100 mL). The ethyl acetate layers were dried over anhydrous Na2SO4 and evaporated to dryness (for the yields see Table 2). Due to a lack of standards, no quantification could be performed. All extracts were prepared to 1 mg/mL and subjected to analytical HPLC (Agilent 1260 Infinity with diode array detector and C18 Acquity UPLC BEH column (2.1 × 50 mm, 1.7 μm) from Waters; solvent A: deionized water (Milli-Q, Millipore, Schwabach, Germany) + 0.1% formic acid, solvent B: acetonitrile + 0.1% formic acid, gradient system: 5% B for 0.5 min increasing to 100% B in 19.5 min, maintaining 100% B for 5 min, flow rate = 0.6 mL min⁻¹, UV detection [190–600 nm], coupled to an ion trap MS (amaZon speed™, Bruker). Mass spectrometry (MS) and ultraviolet (UV) data of the crude extracts were analysed and compared with the literature data using substance databases (CRC Dictionary of Natural Products, CAS SciFinder, Wiley-VCH Antibase).

Blazeispiroils A (1) and D (2) were identified by HPLC-MS comparison (MS, UV/Vis, tR) with an authentic sample (for detailed data see in supporting information; SI).

Blazeispiroil A (1): UV/Vis: λqual = 234, 269, 308 nm; analytical HPLC (tR) = 15.6 min; ESI(+)-MS: m/z (%) = 819 (5) [2M+Na]⁺, 381 (100) [M+H−H2O]⁺.

Blazeispiroil D (2): UV/Vis: λqual = 227, 272, 312 nm; analytical HPLC (tR) = 12.7 min; ESI(+)-MS: m/z (%) = 367 (100) [M+H−H2O]⁺.
Statistical analysis

The optimum growth parameters and mushroom production data were subjected to statistical analysis. The mycelial growth values for growth rate, medium type, biomass, temperature, and pH optimization were compared to obtain a mean separation using Tukey’s test (p<0.05), followed by post hoc tests. The results are expressed in a one-way analysis of variance (ANOVA) analysis using the SPSS program version 19 (IBM Corp.).

Results

Confirmation of the identity of the cultivated species


*Index Fungorum number:* IF292252

*Pileus* at first hemispherical, becoming convex, broadly convex to plano-convex; surface white to cream-colour, ochraceous, with greyish to pale ochraceous fibrils darker to margin, dry; margin incurved at first, finally expanding to decurved. *Lamellae* free, dark brown, crowded, with lamellulae. *Stipe* central, cylindrical; surface white and silky, white appressed fibrils. *Spores print* dark brown, anise-like smell.


*Culture characters:* After 21 days of incubation in PDYA medium, the agar surface was fully colonized with a white mycelium. The mycelium is linear to thread-like hyphae.

*Material examined:* Thailand, Chiang Rai, Muang Chiang Rai, Mae Fah University, on grassland, 20 June 2014, B. Chuankid, BC079 (MFLU 14–0872); living culture, MFLUCC 14–1024.

*Hosts:* On grassland soil—(Dejene et al. 2017; this study).

*Distribution:* Ethiopia, Thailand—(Dejene et al. 2017; this study).

*GenBank accession numbers:* ITS: MZ596239.

*Notes:* The nrITS nucleotide sequences of MFLU 14–0872 revealed that the most similar sequences were from *A. campestroides* IF727842 from Togo (Zhao et al. 2011), at 99.03% similarity with 96% query cover. From Index Fungorum, this species was listed in *Micropsalliota campestroides* (Heinem. & Gooss.-Font.) Heinem. The comparison between the type specimen and our specimen were not similar. The shapes of basidia and cystidia of our specimens were subspherical or clavate, while the type specimen is another shape (claviforms basidia and inconspicuous cystidia). Pileipellis also not incrusted with pigments like other *Micropsalliota*. From this, our specimen fits to descriptions of *Agaricus*.


*Index Fungorum number:* IF561690

The basidiomata of MFLU 14–0874 that grew in the wild was confirmed as the same species as *A. flocculosipes*. The floccose stipe, almond odour, well-developed squamules on the pileus, two-layered annulus, and catenulate cheilocystidia were typical of the species (Thongklang et al. 2014).

*Basidiospores* 5.0–6.5 × (3.5–4.0) µm, *Q* = 1.61, ellipsoid, with apical endosporium, smooth, reddish brown, thick walled. *Basidia* 10–13 × 4–6 µm, clavate, 4-spored. *Cystidia* 40–46 × 8–12 µm, narrowly conical or fusiform to narrowly utriform, hyaline, smooth. *Pileipellis* a cutis; hyphae 6–8 µm in diameter, cylindrical, straight or curved, with brown pigments. *Stipitipellis* a cutis of hyphae 5–8 µm in diameter, cylindrical, smooth, hyaline. Clamp connections absent.
Culture characters: After 21 days of incubation in MYPA medium, the agar surface was colonized with a white mycelium. The mycelium is circular with smooth hyphae.

Material examined: Thailand, Chiang Rai, Muang Chiang Rai, Mae Fah University, on grassland, (date) 9 Jul 2014, B. Chuankid, MFLU 14–0874.

Hosts: on rich soil or over heavily rotted wood in forests or grassland soil—(Zhao et al. 2012; Gui et al. 2015; this study).

Distribution: Thailand and China—(Zhao et al. 2012; Gui et al. 2015; this study).

GenBank accession numbers: ITS: MZ596240.

Notes: The collection has identical morphology with A. flocculosipes, nrITS nucleotide sequence was subjected to a BLAST search against the NCBI database. The nrITS nucleotide sequences in the NCBI database revealed that the most similar sequences were from A. flocculosipes KP705076 from Thailand (Thongklang et al. 2016), at 100% similarity with 96% query cover.

Fig. 1 – a Agaricus campestroides (MFLU 14–0872). b Agaricus flocculosipes (MFLU 14–0874) in the wild. c The mycelial growth of A. campestroides on PDYA medium, d Agaricus flocculosipes on MYPA medium after 30 days.

Effect of media on mycelial growth

In A. campestroides, the largest radial mycelial growth was observed on PDYA, MEA, PDA, and MYPA, respectively. The colour characteristics of the surface mycelium were not different for the four media types. The morphology of the colony was filamentous on all media. The maximum yield by dry weight mycelium occurred on PDYA, followed by MEA, PDA, and MYPA. The same mycelial character was found on A. flocculosipes (MFLU 14–0874). The maximum yield by dry weight mycelium occurred on MYPA, followed by PDYA, PDA, and MEA but the data were not significantly different among four media. The effect of four agar media on mycelial growth (biomass, mg) are given in Table 1.

Effect of temperature on mycelial growth

The optimal temperature for mycelial growth of A. campestroides is between 20 to 30 °C and A. flocculosipes was growing well at 25 °C. (Table 1).
Effect of pH on mycelial growth

The optimal pH of *A. campestroides* was pH 5 with the optimal medium and temperature. The result form this study showed that optimal pH range for mycelial growth of *A. flocculosipes* (MFLU 14–0874) was between 5 to 8. The effects of pH on mycelial growth (biomass, mg) are given in Table 1.

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**Fig. 2** – Microscopic characteristics of *Agaricus campestroides* (MFLU 14–0872). a Basidiospores in KOH. b Basidiospores in Melzer’s reagent. c Basidia. d Cystidia. e Pileipellis. f Stipitipellis. Bars: a, b = 10 µm; c, d, f = 5 µm; e = 20 µm.

**Table 1** Effect of different agar media, temperature, and pH on mycelial growth (biomass, mg) of *A. campestroides* and *A. flocculosipes*. Biomass data = mean weight of dried mycelium± standard deviation. Values with the same letter are not significantly different (*p*<0.05) by Tukey’s test.

<table>
<thead>
<tr>
<th>Medium name</th>
<th><em>A. campestroides</em></th>
<th><em>A. flocculosipes</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>MEA</td>
<td>109.99±0.07898&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>11.23±0.00250&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>PDA</td>
<td>23.58±0.01838&lt;sup&gt;c&lt;/sup&gt;</td>
<td>13.47±0.00347&lt;sup&gt;a&lt;/sup&gt;</td>
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<tr>
<td>MYPA</td>
<td>96.12±0.06923&lt;sup&gt;b&lt;/sup&gt;</td>
<td>21.07±0.00196&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>PDYA</td>
<td>115.51±0.08054&lt;sup&gt;a&lt;/sup&gt;</td>
<td>19.17±0.00151&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Temperature (°C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>50.30±0.05030&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.90±0.00490&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>25</td>
<td>73.53±0.07353&lt;sup&gt;a&lt;/sup&gt;</td>
<td>10.83±0.01083&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>30</td>
<td>80.67±0.08067&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.20±0.00320&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>35</td>
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<td>3.10±0.00310&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>pH</td>
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<tr>
<td>5</td>
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<td>15.67±0.009&lt;sup&gt;a&lt;/sup&gt;</td>
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<tr>
<td>8</td>
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<td>17.87±0.001&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>
Spawn production

Mycelium started to colonize the substrates after five days of incubation, and fully grown within two weeks (Fig. 4). Interestingly, the primordia started to form in three or four weeks of incubation. After one week, the pinning and mushrooms formed in wheat grain spawn.

Fig. 3 – Microscopic characteristics of *Agaricus flocculosipes* (MFLU 14–0874). a Basidiospores in KOH. b Basidia. c Cystidia. d Pileipellis. e Stipitipellis. Bars: a, b = 5 µm; c, e = 10 µm; d = 20 µm.
Harvesting/production
Approximately 30 days after, mycelium colonized the entire compost. Pre-primordia and mature basidiomata formed on the surface of the casing layer after 20 days at 25 °C. Dense hyphal growth always formed on the edge of casing where basidiomata produced. Successfully cultivated Thai strain of A. campestroides is shown in Fig. 5, but the yields were still very low. Unfortunately, the Thai strain of A. flocculosipes could not be cultivated.

Fig. 4 – Mycelial growth of Agaricus campestroides on wheat grain spawn after 14 days. a, c Mycelium fully colonised on substrate in different container. b, d A week after mycelium fully colonised, the primordia and mushrooms formed in spawn.

Fig. 5 – The basidiomata of A. campestroides (MFLU 14–0872) on compost medium after 20 days of casing at 20 °C with 80% humidity.

HPLC profiling and detection of blazeispirol
The crude extracts of A. campestroides and A. flocculosipes were screened for their secondary metabolites in four different liquid media by HPLC-UV/Vis and HPLC-MS. Investigation on the crude extracts indicated the presence of the triterpenoids blazeispirol A (1) and D (2) (Fig. 6), which could be detected from the mycelia of both strains. The compounds were identified based on
their spectral data [UV/Vis, ESI(+)-MS] and HPLC comparison with an authentic sample. Herein, blazeispirol A (1) was observed as the main component of the crude extracts, while blazeispirol D (2) could only detected in small amounts (Fig. 7). Although blazeispirol A (1) could be identified in both mycelium and supernatant of the crude extracts, but the larger amounts was significantly present in mycelium. Comparison of the metabolic profiles of different liquid media demonstrated that the highest production of the compounds (1 and 2) was found in mannitol salts medium (MMK) for both strains and YM in case of A. flocculosipes (Table 3).

Whereas blazeispirol 1 has not yet been detected in the basidiomata, it is produced in large quantities by liquid mycelium fermentation of A. blazei (Grothe et al. 2016). Further investigations hypothesized that blazeispirol D (2) and Z seemed to be the actual intermediates of the blazeispirol A biosynthesis, which might be derived from ergosterol (Hirotani et al. 2001).

According to the MS spectra of the observed compounds, the chlorinated triglycerides could not be detected in any extract.

![Chemical structures of blazeispirol A (1) and D (2).](image)

**Fig. 6** – Chemical structures of blazeispirol A (1) and D (2).

![HPLC-UV/Vis chromatogram of the crude extract of A. flocculosipes cultivated in YM medium.](image)

**Fig. 7** – HPLC-UV/Vis chromatogram of the crude extract of A. flocculosipes cultivated in YM-medium (190–600 nm).

**Discussion**

This present study demonstrates the successful cultivation and domestication of A. campestroides, collected in northern Thailand. For this purpose, the optimal culture medium, pH, temperature, and yield were investigated for the best mycelial growth. For the two strains, used in this study, the best growth rates were obtained using PDYA for A. campestroides and all media were suitable for A. flocculosipes. The two strains of Agaricus showed optimal temperature between 25–30 °C for mycelial growth which is consistent with the previous study by Zied et al.
Agaricus campestroides was growing well at pH 5 while A. flocculosipes was able to grow at pH 5–8. Park (2001) reported the optimal pH for the substrate reaction range between 6–7, high acid or basic condition inhibited mycelial growth. Kopytowski et al. (2008) reported that the optimal pH range for compost substrate and casing layer is pH 7.0–7.5. It is suggested that neutral pH was the suitable pH for these mushroom strains.

The two strains of Agaricus were cultivated by using rice straw as the main substrate mixed with the ingredients. Only A. campestroides were produced at the temperature of 20 °C with 80% relative humidity 14 days after the casing layer was applied. The mushroom requires high temperature and high humidity to produce mushrooms like A. subrufescens (Dias et al. 2004, Mantovani et al. 2007, Dias 2010). The humidity during the formation of primordia was at 80–90%, while 75–80% relative humidity was maintained during the development of mushrooms (Iwade & Mizuno 1997, Stamets 2000). According to Chang (2008), humidity of 70–85% was the optimum for mushroom formation. The wild strains of A. campestroides showed a potential to grow in northern Thailand since the optimal conditions for growing these stains were temperature range from 25–30 °C with 80–90% relative humidity that is related with the weather in northern Thailand (Thai Meteorological Department, 2015).

Moreover, the study of secondary metabolites of A. campestroides and A. flocculosipes has led to the identification of blazeispirols A (1) and D (2) in cultured mycelia of both species.

So far, blazeispirols (A, B, C, D, E, F, G, X, Y, Z) are a class of unique spiro-triterpenoids, isolated from cultured mycelia of Agaricus blazei (=A. subrufescens) (Hirotani et al. 1999, 2000, 2001, 2002) Therefore, the detection of blazeispirols in other species of Agaricus extends the natural occurrence of these type of compounds. Since it was assumed that their occurrence is restricted to A. subrufescens, blazeispirol A (1) was constituted as a phylogenetic and chemotaxonomical marker for this taxon (Thongklang et al. 2017).

Table 2 Yields of crude extracts from the mycelia of A. campestroides and A. flocculosipes cultivated in different media.

<table>
<thead>
<tr>
<th>Strains</th>
<th>MMK</th>
<th>Q6 ½</th>
<th>YM</th>
<th>ZM</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. campestroides</td>
<td>3.2</td>
<td>10.6</td>
<td>1.9</td>
<td>1.8</td>
</tr>
<tr>
<td>A. flocculosipes</td>
<td>3.3</td>
<td>NG</td>
<td>4.5</td>
<td>4.5</td>
</tr>
</tbody>
</table>
(NG): not growth.

Table 3 Occurrence of compounds 1 and 2 in the mycelia of A. campestroides and A. flocculosipes cultivated in different media.

<table>
<thead>
<tr>
<th></th>
<th>A. campestroides</th>
<th>A. flocculosipes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MMK</td>
<td>Q6 ½</td>
</tr>
<tr>
<td>Blazeispirol A (1)</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>Blazeispirol D (2)</td>
<td>+</td>
<td>-</td>
</tr>
</tbody>
</table>
(-): not observed, (+): observed in traces, (++): observed in small amounts, (+++): observed in high amounts, (NG): not growth.

Conclusion

The two strains of Agaricus from northern Thailand were cultivated by using rice straw as the main substrate mixed with the ingredients. The basidiomata were only produced from A. campestroides at the temperature of 20 °C with 80% relative humidity 14 days after the casing layer was applied. The best rates of mycelial growth were obtained using PDYA medium adjusted to pH 5 for A. campestroides and incubated between 20 to 30 °C. All media were suitable for mycelial growth of A. flocculosipes. The mycelial of A. flocculosipes was growing well at 25 °C and at pH 5–8. The study of secondary metabolites of A. campestroides and A. flocculosipes has led
to the identification of blazeispirols A (1) and D (2) in cultured mycelia of both species. *Agaricus* cultivation conditions will benefit Thai farmers. Compost production from rice straw would help to reduce the burning of rice straw in Thai rice fields. Furthermore, *A. campestrisoides* and *A. flocculosipes* could be served as additional sources for blazeispirols and their potential medicinal properties.

**Acknowledgements**

Financial support by the Research and Researchers for Industries grant (PHD57l0015) and Thailand Science Research and Innovation (TSRI) grant, Macrofungi diversity research from the Lancang-Mekong watershed and surrounding areas (Grant No. DBG6280009). Many thanks to Olivier Raspé and Luis A. Parra for their suggestions on species identification. The help of Anan Thawthong and Kritsana Jatuwong at Mae Fah Luang University in mushrooms collecting and cultivation is acknowledged. Thanks to Sebastian Pfütze who gave us the analytical data of the compounds. Finally, B.C. would like to thank Benjarong Thongbai for her support during the stay in Germany.

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Supplementary data

Supporting information:

**Blazeispirol A (1)** [authentic sample]: UV/Vis: \( \lambda_{\text{qual}} = 236, 268, 308 \) nm; analytical HPLC \((t_R) = 15.20 \text{ min}; \) ESI(+)−MS: \( m/z \) (%) = 819 (5) \([2\text{M+Na}]^+\), 381 (100) \([\text{M+H–H}_2\text{O}]^+\).

**Blazeispirol D (2)** [authentic sample]: UV/Vis: \( \lambda_{\text{qual}} = 229, 270, 310 \) nm; analytical HPLC \((t_R) = 12.40 \text{ min}; \) ESI(+)−MS: \( m/z \) (%) = 791 (6) \([2\text{M+Na}]^+\), 367 (100) \([\text{M+H–H}_2\text{O}]^+\).

**Identification of blazeispirol A (1) and D (2) in Agaricus flocculosipes (MFLU 14-0874):**

Blazeispirol A (1): UV/Vis: \( \lambda_{\text{qual}} = 234, 269, 308 \) nm; analytical HPLC \((t_R) = 15.6 \text{ min}; \) ESI(+)−MS: \( m/z \) (%) = 819 (5) \([2\text{M+Na}]^+\), 381 (100) \([\text{M+H–H}_2\text{O}]^+\).

![HPLC Chromatogram](BC081_Mycelium_MMK_GA7_01_17088.d: UV Chromatogram, 190-600 nm)

**Fig. 1** – HPLC chromatogram of blazeispirol A (1) identified from the crude extract of *A. flocculosipes* (mycelium) cultivated in MMK-medium.

![ESI-MS Spectrum](BC081_Mycelium_MMK_GA7_01_17088.d: +MS, 15.6-15.6min #1957-1965)

**Fig. 2** – ESI-MS spectrum of blazeispirol A (1) identified from the crude extract of *A. flocculosipes* (mycelium) cultivated in MMK-medium.
Blazeispirol D (2): UV/Vis: $\lambda_{\text{qual}} = 227, 272, 312$ nm; analytical HPLC ($t_R$) = 12.70 min; ESI(+)-MS: $m/z$ (%) = 367 (100) [$\text{M}+\text{H}–\text{H}_2\text{O}]^+$. 

**Fig. 3** – UV/Vis spectrum of blazeispirol A (1) identified from the crude extract of *A. flocculosipes* (mycelium) cultivated in MMK-medium.

Blazeispirol D (2): UV/Vis: $\lambda_{\text{qual}} = 227, 272, 312$ nm; analytical HPLC ($t_R$) = 12.70 min; ESI(+)-MS: $m/z$ (%) = 367 (100) [$\text{M}+\text{H}–\text{H}_2\text{O}]^+$. 

**Fig. 4** – HPLC chromatogram of blazeispirol D (2) identified from the crude extract of *A. flocculosipes* (mycelium) cultivated in YM-medium.
Fig. 5 – ESI-MS spectrum of blazeispirol D (2) identified from the crude extract of *A. flocculosipes* (mycelium) cultivated in YM-medium.

Fig. 6 – UV/Vis spectrum of blazeispirol D (2) identified from the crude extract of *A. flocculosipes* (mycelium) cultivated in YM-medium.