Fruiting Body Production and Aroma Profile Analysis of *Agrocybe aegerita* Cultivated on Different Substrates

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Abstract

The industrial cultivated basidiomycete *Agrocybe aegerita*, also known as “Pioppino” in Italy or “Samthaube” in Germany, is a high quality mushroom with a delicious aroma. Cultivation of *A. aegerita* on wheat straw supplemented with different residues of the food industry showed the highest yields in fruiting body production with a biological efficiency of 36% when black tea pomace was added. The addition of other substrates resulted in biological efficiencies of 23% to 33%. Besides the mushroom yields, the number and size of fruiting bodies harvested on the different substrates were determined. A comparison of the aroma profiles of *A. aegerita* grown on wheat straw and on wheat straw supplemented with black tea by means of GC/MS/MS/O is presented.

Keywords  
*Agrocybe aegerita*, Aroma Profile, Mushroom Production, GC/MS/MS/O

1. Introduction

Since thousands of years, mushrooms are favoured due to their delicious taste and their nutritive and healthy values [1] [2]. The most commonly cultivated mushrooms are *Agaricus bisporus*, *Pleurotus ostreatus* and *Lentinula edodes* counting for more than 80% of the worldwide mushroom production [1]. Besides, several other tasty mushrooms can be cultivated and have attracted increasing interest in recent years. One of them, the black poplar mushroom *A. aegerita* (syn. *A. cylindracea*) is a well-known high quality mushroom with a pleasant aroma [3]. The fruiting bodies of *A. aegerita* are known for biosynthesis of several secondary metabolites that have been studied in more detail regarding antioxidative and antimicrobial activity [4] as well as tumor pro-

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In general, the quality of mushrooms depends essentially on the aroma of the fruiting bodies. The aroma profile varies between species and may be influenced by the culture conditions or by addition of aroma precursors like unsaturated fatty acids [6] [7]. Furthermore, cooking modifies the aroma profile and causes the formation of new volatiles [8]. Among the different volatile compounds, mushrooms form a series of C8-compounds of which oct-1-en-3-ol is the most abundant one [6].

Growth of *A. aegerita* on lignocellulosic material, such as wheat straw, cotton waste, sawdust, corn cobs and peanut shells has been reported [8]-[11]. Additionally, many other lignocellulosic waste materials have been suggested for edible mushroom cultivation [12], but have not been used for *A. aegerita* so far. Compared to *L. edodes* and *P. ostreatus* the cultivation yields of *A. aegerita* are typically low [3] and the productivity of the commercial strains decreases after consecutive cultivation rounds [13]. Besides the optimal cultivation conditions, the choice of the production strain is an important factor for the yields.

In the European Union each year nearly 90 million tons of food waste is produced and 38% directly accrues at the food industry [14]. In the current study, the utilisation of residual materials from the food industry was tested for the production of fruiting bodies of *A. aegerita*. The spent substrates may be further used after mushroom production as animal feed, because degradation of lignocelluloses by mushrooms improves the availability of nutrients [15].

## 2. Materials and Methods

### 2.1. Cultivation of *Agrocybe aegerita*

Mushroom spawn of *A. aegerita* (strain 4022) was provided by Sylvan Inc. (Horst, The Netherlands). The spawn was stored at 4°C until inoculation.

For cultivation of *A. aegerita* wheat straw supplemented with different residual materials from the food industry was used. *A. aegerita* was cultivated on 100% wheat straw, 100% cocoa shells and on wheat straw supplemented with either cocoa shells (17%), citrus pellets (17%), carrot mesh (17%) or black tea pomace (17% and 45%). The humidity of the different substrates was determined by an infrared moisture analyser (MA 35, Sartorius, Göttingen, Germany) at 105°C. The dry weights of these substrates were used to calculate the amount of water to adjust the humidity of the non-dried substrates to 75%. The substrates were mixed in a cement mixer (GBM 125, DEMA, Übrigshausen, Germany). Autoclavable polyethylene culture bags (Unicorn Imp. & Mfg. Corp., Culemborg, Netherlands) were filled with 1.5 kg of the moistened substrate. For sterilisation, the bags were autoclaved at 121°C for 1 h. After the substrate was cooled down, the bags were inoculated with 10% (w/w) *A. aegerita* spawn. All substrates were produced in four replications. For mycelia growth, the bags were incubated at 24°C in darkness at 75% relative air humidity for four weeks in a climate chamber (Vötsch VB 1514, Minicon, Balingen-Frommern, Germany) until the mycelium was grown through the whole substrate.

For induction of fructification, the bags were sliced (width 1 cm) at six positions, and the culture conditions were modified to 90% relative humidity, 20°C, and incandescent light (400 lux) for 12 h in the climate chamber. The fruiting bodies were harvested when the caps were arcuate down and the lamellas were still not visible. The biological efficiency (BE) was calculated as percentage yield of fresh fruiting bodies of *A. aegerita* in relation to dry weight of the substrate.

\[ \text{BE}\% = \frac{\text{Weight of fresh fruiting bodies}}{\text{Weight of dry substrate}} \times 100\% \]

### 2.2. Extraction

Chemicals were provided by Carl Roth (Karlsruhe, Germany), Sigma-Aldrich (Steinheim, Germany), Acros Organics (Geel, Belgium) and Fisher Scientific (Schwerze, Germany). Solvents were distilled before use. Ripe fruiting bodies were harvested and frozen at −20°C. 100 mL methanol, 100 mL water and 1 mL hept-1-en-3-ol solution (5.0833 mg L⁻¹ in ethanol or 5.2025 mg L⁻¹ in methanol) as internal standard were added to 100 g fruiting bodies. The sample was homogenised with a masher (HR1361/00, Philips, Hamburg, Germany) and an Ultra-Turrax (T 25, IKA, Staufen, Germany). After centrifugation (4700 × g, 4°C, 10 min), the supernatant was subjected to liquid/liquid extraction (three times with 200 mL pentane/diethyl ether, 1:1.12 v/v) in a separatory
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funnel. The combined pentane/diethyl ether fractions were dried over anhydrous sodium sulphate and concentrated to a volume of about 1 mL using a Vigreux column (water bath temperature 45°C).

2.3. GC/MS/MS/O Analysis

The analyses were performed by means of a gas chromatograph (7890A, Agilent Technologies, Waldbronn, Germany) equipped with a VF-WAXms column (Agilent Technologies, 30 m × 0.25 mm i.d., 0.25 μm film thickness), operated in splitless mode (splitless time 0.5 min). Inlet temperature was 250°C. Helium (5.0) was used as carrier gas at 1.56 mL·min⁻¹ (constant flow). The oven temperature program was set to: 40°C (3 min), 5°C·min⁻¹ to 240°C (27 min). After the column, the carrier gas was split 1:1 into a triple quadrupole mass spectrometer (7000B, Agilent Technologies; electron impact energy: 70 eV, transfer line: 250°C, source: 230°C, quadrupoles: 150°C, He quench gas: 2.25 mL·min⁻¹, N₂ collision gas: 1.5 mL·min⁻¹, scan in Q1 m/z = 33-300) and an olfactory detector port (ODP 3, GERSTEL, Mülheim an der Ruhr, Germany, transfer line: 250°C, mixing chamber: 150°C, make up gas: N₂). Alternatively, a DB-5MS column (Agilent Technologies, 30 m × 0.25 mm i.d., 0.25 μm film thickness) with analogous parameters was used. The volatile compounds were identified by comparison of the obtained Kováts retention indices (RI) and mass spectra with authentic standards and database entries (NIST 2011). RI were calculated from the retention times of n-alkanes (C8-C30, Sigma-Aldrich, Steinheim, Germany) by linear interpolation. The following standards, from Carl Roth, Acros Organics, AppliChem (Darmstadt, Germany) and Alfa Aesar (Karlsruhe, Germany), were chromatographed under identical conditions: hexane, octan-3-one, oct-1-en-3-one, oct-1-en-3-ol, acetic acid, benzaldehyde, (E)-oct-2-en-1-ol, butanoic acid, isovaleric acid, pentanoic acid and 2-phenylethanol.

2.4. Statistics

Analysis of variance (ANOVA) was calculated by the program OriginPro 9.0 (OriginLab Corporation, Northampton, MA, USA) with Fisher’s least significant difference (LSD) test for comparing the mean values.

3. Results and Discussion

*A. aegerita* showed similar growth on wheat straw and all wheat straw containing substrates. On 100% cocoa shells the fungus showed only marginally growth around the inoculum which might be explained by an anti-microbial activity of the cocoa shells [16]. A second reason might be the inferior oxygen supply in less structured substrates compared to wheat straw, where the loose structure improves the availability of oxygen [17].

In a study by Zervakis et al. [9], different agro-waste materials were used for growing *A. aegerita*. They observed colonisation on the more structured substrates, such as peanut shells and sawdust, whereas on olive press cake no growth was observed. Nevertheless, certain mushrooms do grow on 100% cocoa shells, as shown by Bermúdez et al. [18] who cultivated *Pleurotus ostreatus* on this substrate.

In our case on 100% cocoa shells, no fruiting bodies could be harvested due to reduced growth of mycelia and absent primordia. In all wheat straw derived substrates, primordia formation started five weeks after inoculation. Interestingly, in wheat straw substrate supplemented with cocoa shells, the numbers of *A. aegerita* primordia exceeded those of all other substrates (Figure 1).

Fruiting bodies were harvested in a time span of 20 days starting five weeks after inoculation. The BE of fruiting body production varied from 23% to 36% between the different substrates (Figure 2). The BE on 100% wheat straw (WS) of 33% ± 11% was close to the yields obtained on wheat straw supplemented with cocoa shells (WS/CS; 31% ± 5%) and carrot mesh (WS/CM; 31% ± 7%). Supplementation with citrus pellets (WS/CP) reduced the yields to 23% ± 5%. The most efficient substrate mixture was wheat straw supplemented with black tea pomace. BE of 36% ± 7% and 36% ± 1% were obtained with wheat straw supplemented with 17% black tea pomace (WS/BT) and wheat straw with 45% black tea pomace (BT/WS), respectively. The BE of *A. aegerita* on 100% WS of 29.5% as reported by Schmithals and Schildbach [3] was comparable to the results of our study, while they could nearly double the BE (BE 58.0%) by supplementation of wheat straw with 50% draff. In addition to the different substrate supplements they used, a harvest period of 22 weeks positively affected the BE [17]. In another study, the BE of three different *A. aegerita* strains cultivated on WS were, similar to our observations, between 25.5% and 31.0% [13]. However, two further strains of *A. aegerita* showed higher production yields of up to 49.8%; it should be mentioned that these yields were obtained during two flushes [8]. On the
Figure 1. Primordia formation of *A. aegerita* cultivated on wheat straw supplemented with 17% cocoa shells.

Figure 2. BE of *A. aegerita* grown on wheat straw (WS), WS and cocoa shells (WS/CS), WS and citrus pellets (WS/CP), WS and carrot mesh (WS/CM), WS and 17% black tea pomace (WS/BT) and WS and 45% black tea pomace (BT/WS) with standard deviations.

other hand, also strains with lower BE have been tested as shown for strain 156 with a BE of only 10.1% ± 2.4% [19].

Furthermore, the fruiting bodies were counted and grouped according to their size. The elevated primordia production observed on WS/CS (Figure 1) led to many small fruiting bodies (Figure 3). The largest fruiting bodies were obtained on WS/BT and BT/WS. The yields on wheat straw supplemented with black tea pomace were the highest though the number of fruiting bodies was lower compared to fruiting bodies grown on other substrates. Thus, wheat straw supplemented with black tea pomace seems to be favourable for industrial mushroom production. Beside the fruiting body yield, the aroma is the most important quality factor of mushrooms. Therefore, the effect of black tea pomace supplementation of the growth substrate on the aroma profile of *A. aegerita* was analysed by GC/MS/MS/O (Figure 4).
The overall aroma profile of *A. aegerita* grown on BT/WS showed no significant differences compared to the cultivation on WS. Eleven aroma compounds were detected in both extracts (Table 1). Beside the typical mushroom flavours oct-1-en-3-one and oct-1-en-3-ol, a bloomy, rose like flavour was perceived, which was identified as 2-phenylethanol. Two green (cut grass) flavour compounds, hexane and octan-3-one, were identified as well. Additionally, acetic acid (vinegar flavour), benzaldehyde (almond flavour) and (E)-oct-2-en-1-ol (rancid odour) were identified. These compounds were previously detected by Rapior *et al.* [20] and in a preliminary study of our group [21]. In addition, three acids (butanoic acid, isovaleric acid and pentanoic acid) were tentatively identified, which have not been detected in *A. aegerita* before. While Rapior *et al.* [20] identified 22 volatiles in an organic extract of *A. aegerita*, only eleven compounds were perceived in the current study. A reason for this might be the different extraction methods. In this study, the extraction of volatile compounds was
Table 1. Perceived odour impressions in extracts of *A. aegerita* by means of GC/MS/MS/O; fruiting bodies were grown on wheat straw (WS) and on wheat straw supplemented with black tea pomace (BT/WS); RI: Kováts retention index.

<table>
<thead>
<tr>
<th>Peak</th>
<th>Odour impression</th>
<th>Compound</th>
<th>RI WS</th>
<th>RI BT/WS</th>
<th>RI standard</th>
<th>RI literature</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>VF-WAXms</td>
<td>DB-5MS</td>
<td>VF-WAXms</td>
<td>DB-5MS</td>
</tr>
<tr>
<td>1</td>
<td>Green</td>
<td>Hexanal</td>
<td>1087</td>
<td>801</td>
<td>1087</td>
<td>801</td>
</tr>
<tr>
<td>2</td>
<td>Green</td>
<td>Octan-3-one</td>
<td>1257</td>
<td>992</td>
<td>1257</td>
<td>992</td>
</tr>
<tr>
<td>5</td>
<td>Vinegar</td>
<td>Acetic acid</td>
<td>1458</td>
<td>&lt;800</td>
<td>1458</td>
<td>&lt;800</td>
</tr>
<tr>
<td>6</td>
<td>Almond</td>
<td>Benzaldehyde</td>
<td>1523</td>
<td>n.d.</td>
<td>1523</td>
<td>968</td>
</tr>
<tr>
<td>7</td>
<td>Rancid</td>
<td>(E)-oct-2-en-1-ol</td>
<td>1615</td>
<td>1073</td>
<td>1615</td>
<td>1073</td>
</tr>
<tr>
<td>8</td>
<td>Cheese</td>
<td>Butanoic acid</td>
<td>1635</td>
<td>n.d.</td>
<td>1632</td>
<td>n.d.</td>
</tr>
<tr>
<td>10</td>
<td>Cheese</td>
<td>Pentanoic acid</td>
<td>1742</td>
<td>n.d.</td>
<td>1742</td>
<td>n.d.</td>
</tr>
</tbody>
</table>

The compounds were detected in *A. aegerita* extracts from fruiting bodies grown on WS as well as on BT/WS. Furthermore, volatiles were confirmed by comparing RI with literature data and sensory evaluation by olfactometry analysis; n.d.: no distinct MS spectrum on DB-5MS column.

performed three times, each step lasting approximately five minutes, whereas Rapior *et al.* macerated the fruiting bodies for seven days in the organic solvent dichloromethane.

**4. Conclusion**

Cultivation of *A. aegerita* on different substrates based on wheat straw supplemented with different residual materials from the food industry was possible. In addition to the exploitation of these waste materials, the yields of fruiting bodies could be increased by supplementing wheat straw with black tea pomace, a by-product of the beverage industry. The thus produced mushrooms should be as marketable as the ones produced on wheat straw, as a comparison of the aroma profiles of mushrooms grown on both substrates showed no significant differences. Eleven aroma compounds were detected in the extracts of the fruiting bodies of *A. aegerita*: hexane, octan-3-one, oct-1-en-3-one, oct-1-en-3-ol, acetic acid, benzaldehyde, (E)-oct-2-en-1-ol, butanoic acid, isovaleric acid, pentanoic acid and 2-phenylethanol.

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**References**


