Improvement of Nutritional Values of *Agaricus Bisporus* Mushroom Stalk and Cap through Solid-State Fermentation Using *Aspergillus niger*

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**ABSTRACT**

This study aimed to investigate the effect of solid-state fermentation on the nutritional composition of *Agaricus bisporus* mushroom stalk and cap. Four homogenous test samples were prepared from the fermented mushroom stalk (Fermented MS), unfermented mushroom stalk (MS), fermented mushroom cap (Fermented MC), and unfermented mushroom cap (MC); each was independently analyzed in triplicates, providing a total of 12 independent measurements of all treatments. Nutritional changes in the mushrooms were determined by analyzing crude protein (CP), ether extract (EE), crude ash (CA), crude fiber (CF), hemicellulose, neutral detergent fiber (NDF), acid detergent lignin (ADL), and acid detergent fiber (ADF) contents. The best improvement in nutritional composition was obtained from the MS with increasing the crude protein (35%). The results related to other parameters are similar among all the treatments, except for CF. To conclude, the fermented MS can be considered as an alternative protein resource in animal nutrition.

**Keywords:** Mushroom, Nutrient value, Fermentation, Animal nutrition, Waste product

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**Introduction**

For centuries, people have used natural resources like plants, minerals, and animal products in treatment and prevention of many diseases of themselves and their animals. Mushroom is one of the natural resources that used as medicines or functional food on the world. Because they have beneficial effects on health, such as antibacterial, antifungal, anti-oxidant, hypoglycemic activities (Wang and Ng, 1999) and produced some protein such as lectins (Wang et al. 2002), ribosome-inactivating proteins, antifungal proteins (Lam and Ng, 2001) and rib nucleases (Kobayashi et al. 1992).

Mushroom production has been rapidly raised in Turkey about more than 40,712 tons and in the world more than 10 million tons annually (TUİK, 2017). Mushrooms are an ideal food for people due to the high content of fiber and protein or low content of fat (Wong et al. 2003). Therefore, edible mushrooms have the important potential in filling the nutritional gap in human nutrition. On the other hand, it would be difficult to say that mushrooms are quite cheap feedstuff for animal nutrition in the world, except for by-products of mushrooms such as stalk, the surplus of need, near to shelf life of mushroom, etc. Although there have been so many studies which were reported the effects of mushrooms powder (intact form) or extracts on performance, immune traits, meat quality, gastrointestinal traits in broilers, laying hens or pigs (Guo et al., 2004a; Guo et al., 2004b; Giannenas et al., 2010a; Giannenas et al., 2010b), there has been limited studies about the basic chemical composition, the nutrient digestibility, the metabolizable energy content of mushroom stalk meal or cap. For example, Buwjoom et al., (2004) reported that there was difference between stalk meal and cap of *L. edodes* in terms of crude protein (CP), ether extract (EE), crude fiber, gross energy, calcium, and phosphorus contents, and *L. edodes* stalk meal could be used as an alternative feedstuff for poultry. Lee et al., (2005) suggested that if these by-products were evaluated in animal feeds and feed additives or any field, they would cause environmental pollution such as a bad odor in the future.
Recently, solid-state fermentation has been considered as a useful and inexpensive method to improve the nutritional composition of feedstuffs and to eliminate its antinutritional factors. Indeed, improvement of both protein quantity and quality (Mathivanan et al., 2006; Zhang et al., 2012), degradation of non-starch polysaccharides to monosaccharides such as glucose and eliminating of possible antinutritional factors (Zhao et al., 2013) in feedstuffs or agricultural residues. However, there has been limited information about the nutrient values or zootechnical performance traits of fermented mushrooms stalk or cap (Chu et al., 2012). Furthermore, there is no report about the nutrient composition of fermented A. bisporus stalk or cap. In light of all the above information, the study was conducted to determine whether mushrooms stalk and cap is suitable for SSF and the changes in their nutrient composition.

Materials and Methods

Preparation of A. bisporus

Mushrooms (A. bisporus) were obtained from a local mushroom producer (Samsun, Turkey) and the intact mushrooms were divided into two parts as stalk and cap before drying. Then, each part of mushrooms was sundried for three days to a stable moisture content of about 10-20% and the soils attached to mushroom stalk were removed. Samples were oven-dried at 60-70°C for 48 h (Nadubuyu et al., 2010) and were milled a 2 mm sieve for chemical analysis (Retch ZM200, Haan, Germany). Four homogenous test samples were prepared from the fermented mushroom stalk (Fermented MS), unfermented mushroom stalk (MS), fermented mushroom cap (Fermented MC), and unfermented mushroom cap (MC); each was independently analyzed in triplicates, providing a total of 12 independent measurements of all treatments.

Aspergillus niger

Aspergillus niger (ATCC 9142) was obtained from the American Type Culture Collection (ATCC, USA). A. niger cultures were incubated at 24°C for 7 days in potato dextrose agar (PDA, MERCK) medium according to agar plate technique. After incubation, A. niger spores were harvested by inverting and tapping the top of the plate, slightly. Spores count was determined by Fuch-Rosenthal technique with hematocytometer. The spores were inoculated into the solid fermentation medium on the same day.

Solid-State Fermentation

After mushrooms and nutritional salt were autoclaved at 121°C for 15 minutes, one hundred g mushroom stalk or cap and 160 ml potato dextrose broth (Merck, Germany) was mixed in anaerobic bags. The pH during fermentation was calibrated to 5 using 1 N NaOH and HCl. Starting humidity was 60% and for each g of the solid environment 1×10^6 A. niger spores were inoculated into a sterile chamber and left to incubate at 28–30°C. After incubation, fermented mushroom stalk or cap were placed in plastic containers, gently pressed, and left for 48 hours at ambient temperature. Since A. niger is a micro aerobic organism, there were sufficient micro aerobic conditions for its growth and development in this closed environment (David et al., 2003). After 48 hours, samples were spread over polyethylene paper and left for 6 days at 30–40°C until reaching approximately 90% of dry matter, after which they were splintered into 0.15-mm pieces.

Nutrient Composition of A. bisporus

Crude ash content was determined by incineration in a muffle furnace at 550°C for 8 hours. Nitrogen (N) contents were analyzed using the Kjeldahl method according to the AOAC (1998) procedure. Crude protein (CP) was calculated as N×6.25. The ether extracts (EE) content was determined by using AnkomXT15 analyzer (AOAC, 1998). Crude fiber content was determined by using The ANKOM2000 Fiber Analyzer. The analyses of neutral detergent fiber (NDF), acid detergent lignin (ADL), and acid detergent fiber (ADF) contents of the leaves were based on the method of Van Soest et al. (1991) using an Ankom fiber analyzer. Hemicellulose was calculated as NDF minus ADF. Nitrogen-free extract (NFE) was estimated on a dry weight basis by subtracting the percentages of CP, EE, CF and CA from 100%.

Statistical Analysis

The data were analyzed by using Windows version of SPSS 21.0 (SPSS Inc., NY, and USA) statistical package program. T-test was used to compare the differences between groups after Shapiro-Wilk test was used for the normality assumption of data.

Results and Discussion

The nutrient composition of MS and fermented MS before and after fermentation was presented in Table 1. According to the data, the CF (P<0.01), CP, CA, NDF and ADF of the fermented MS were higher than those of MS (P<0.001). There were no differences between treatments in terms of EE and ADL (P>0.05). The variation in nutrient composition of MC or the fermented MC was generally similar to those of MS or the fermented MC (Table 2). While the CP, CA, CF, ADF, and ADL of fermented MC were increased, its NFE and hemicellulose were decreased (P<0.05). The EE and NDF contents of both MC and fermented MC were not found differences. In general, these findings have been consistent with studies reported that fermented feed raw materials, agricultural wastes, and agricultural by-products were better nutrient composition according to non-fermented products (Mathivanan et al., 2006; Zhang et al., 2012; Zhao et al., 2013). On the other hand, the finding regarding the increase in cellulose content of MS after fermentation has been contradicted with the findings of many studies indicating that cellulose content decreases after fermentation (Ihuyemi et al., 2006; Zhang et al., 2007; Lawal et al., 2010; Güngör et al., 2017; Aktop et al., 2018; Jannathulla et al., 2018). These incompatibilities among the studies may be due to the chemical properties (cellulose content) of the substrates used, the type of microorganisms selected (fungi, bacteria, yeast, etc.), fermentation ambient conditions (humidity, temperature, O2) or variability in fermentation times.

In this study, an increase in protein content of fermented MS was an expected finding. Indeed, the protein content of the fermented product was higher than those of non-fermented products in previous studies (Zhang et al., 2007; Güngör et al., 2017; Jannathulla et al., 2018; Aktop et al., 2018). The EE content of fermented MS was higher than those of MS. Crude protein (CP) was calculated as N×6.25. The ether extracts (EE) content was determined by using AnkomXT15 analyzer (AOAC, 1998). Crude fiber content was determined by using The ANKOM2000 Fiber Analyzer. The analyses of neutral detergent fiber (NDF), acid detergent lignin (ADL), and acid detergent fiber (ADF) contents of the leaves were based on the method of Van Soest et al. (1991) using an Ankom fiber analyzer. Hemicellulose was calculated as NDF minus ADF. Nitrogen-free extract (NFE) was estimated on a dry weight basis by subtracting the percentages of CP, EE, CF and CA from 100%.

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et al., 2018). This was associated with an increased amount of fungus or bacteria during fermentation process (Raimbault, 1998; Nair et al., 2008). Bacteria or fungi selected for fermentation have been also microbial protein sources and their protein contents were high. Therefore, the increase in protein content of fermented MC could be explained with the increased amount of A. niger during the fermentation process in the present study. On the other hands, the cellulose content of fermented MS was increased, contrary to other studies (Iluyemi et al., 2006; Zhang et al., 2007; Lawal et al., 2010; Güngör et al., 2017; Altop et al., 2018; Jannathulla et al., 2018). This could be closely related to the increased amount of A. niger in the fermentation process like the increase in protein content. In A. niger cell, its cell wall usually have exhibited a multilaminar and fibrillar feature, and this could increase the stability of the cell wall. Aspergillus niger has contained polysaccharides (about 80%), protein (5-15%) and lipids (3-10%). Among the polysaccharides, glucan, galactose, chitin, chitosan, mannan, and cellulose are the most abundant (Arda, 2000). Considering the changes in the EE and CP content of the fermented MS in the study, it could be said that the increase in CF content was due to the increase in A. niger number during fermentation process.

Table 1  Changes in nutrient composition of Agaricus bisporus mushroom stalk before and after fermentation (based on DM %)

<table>
<thead>
<tr>
<th>Composition, %</th>
<th>MS</th>
<th>Fermented MS</th>
<th>SEM</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>CP</td>
<td>22.36</td>
<td>30.40</td>
<td>1.798</td>
<td>***</td>
</tr>
<tr>
<td>EE</td>
<td>0.87</td>
<td>1.17</td>
<td>0.167</td>
<td>NS</td>
</tr>
<tr>
<td>CA</td>
<td>11.42</td>
<td>14.74</td>
<td>0.749</td>
<td>***</td>
</tr>
<tr>
<td>NFE</td>
<td>51.42</td>
<td>29.22</td>
<td>4.979</td>
<td>***</td>
</tr>
<tr>
<td>CF</td>
<td>13.93</td>
<td>24.47</td>
<td>2.392</td>
<td>***</td>
</tr>
<tr>
<td>NDF</td>
<td>41.06</td>
<td>44.96</td>
<td>0.902</td>
<td>NS</td>
</tr>
<tr>
<td>ADF</td>
<td>19.49</td>
<td>28.54</td>
<td>2.029</td>
<td>**</td>
</tr>
<tr>
<td>ADL</td>
<td>6.42</td>
<td>6.59</td>
<td>0.493</td>
<td>NS</td>
</tr>
<tr>
<td>Hemicellulose</td>
<td>21.57</td>
<td>16.43</td>
<td>1.153</td>
<td>***</td>
</tr>
</tbody>
</table>

Table 2  Changes in nutrient composition of Agaricus bisporus mushroom cap before and after fermentation (based on DM %)

<table>
<thead>
<tr>
<th>Composition</th>
<th>MC</th>
<th>Fermented MC</th>
<th>SEM</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>CP</td>
<td>33.19</td>
<td>37.15</td>
<td>0.904</td>
<td>***</td>
</tr>
<tr>
<td>EE</td>
<td>1.62</td>
<td>1.30</td>
<td>0.201</td>
<td>NS</td>
</tr>
<tr>
<td>CA</td>
<td>11.10</td>
<td>13.54</td>
<td>0.553</td>
<td>***</td>
</tr>
<tr>
<td>NFE</td>
<td>36.37</td>
<td>25.06</td>
<td>2.539</td>
<td>***</td>
</tr>
<tr>
<td>CF</td>
<td>17.72</td>
<td>22.94</td>
<td>1.178</td>
<td>**</td>
</tr>
<tr>
<td>NDF</td>
<td>51.87</td>
<td>40.66</td>
<td>3.741</td>
<td>NS</td>
</tr>
<tr>
<td>ADF</td>
<td>19.76</td>
<td>28.08</td>
<td>1.867</td>
<td>**</td>
</tr>
<tr>
<td>ADL</td>
<td>5.51</td>
<td>8.69</td>
<td>0.730</td>
<td>NS</td>
</tr>
<tr>
<td>Hemicellulose</td>
<td>32.11</td>
<td>12.58</td>
<td>5.130</td>
<td>*</td>
</tr>
</tbody>
</table>

In the present study, the amount of CA of the fermented MC and MS were increased. These results are consist in previous studies related to fermentation (Güngör et al., 2017; Altop et al., 2018). The increase in CA could be attributed to the phytase enzyme produced by the fermenter during the process (Selle et al., 2007; Adeola, 2010). In order to confirm this idea, it is important to determine the phosphorus content of the samples after fermentation. Therefore, more detailed studies are required. However, the presence of soil dust, etc. over the sample may affect its ash content. In the study, the higher CA content of the fermented MS as compared to MS could be attributed to the compost on the mushroom stalk could not be completely cleaned. On the other hands, the increase in CA content both of the fermented MS and MC confirmed the idea that content could be as a result of the growth and multiplication of the microorganism in the fermentation medium (Ahaotu et al., 2013)

The EE content of both the fermented MS and MC was not affected. This finding is disagree with the previous studies reported that the EE content of the product after fermentation increased or decreased (Ahmed et al., 2016; Güngör et al., 2017; Altop et al., 2018). Oboh et al (2002) suggested that the increase in the EE content might due to the increase in the microbial mass, activities of the lipolytic microorganism to secrete an extracellular enzyme (lipase), secretion of microbial oil into the fermenting medium and other products from metabolism. On the contrary, Vries and Visser (2001) suggested that the decreases in EE contents may be attributed to the degradation by extracellular enzymes secreted by A. niger; this strain secretes a number of enzymes including alpha-galactosidase, hemicellulose, and pectinase. During fermentation, the enzymes secreted are known to metabolize carbohydrate as a carbon source for the growth of the strain (Nigam and Singh, 1994). Thus, it could be attributed that there was no differences between EE contents of fermented MC and MS because the fermenters used in this study had low lipase enzyme production potential or absence of lipolytic microorganism.

Conclusion

According to the results, it is possible to say that (1) both A. bisporus stalk and cap are suitable substrates for solid-state fermentation process, (2) the fermented mushroom products could be higher nutritive values than those of non-fermented. Thus, A. bisporus stalk (especially fermented stalk), is by product in mushroom production, has a potential nutritional value that can be considered as a protein source in farm animal nutrition.
References


