Stoichiometry equation to describe the growth of the Pleurotus ostreatus ceba-glie-po-010606 strain

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Abstract
This work was aimed at developing a proximate stoichiometric equation to describe the growth of Pleurotus ostreatus mushroom strain ceba-glie-po-010606 on picking beans (Phaseolus vulgaris) waste. Empirical formulas were established for the residue of fresh dried picking beans (CH1.81O0.81N0.15) and the biomass of the fungal strain (CH1.83O0.84N0.26). The elemental composition of these materials and the ashes were determined. The stoichiometric coefficients obtained further supported the estimation of parameters relevant for fungal growth characterization: theoretical biological efficiency (867.49 g of fungal dry matter (FDM)/kg of substrate dry matter), mean coefficient of breath (0.77 mol CO2/mol O2 consumption), specific air consumption (1.36 m3/kg FDM) and metabolic heat (16 576.47 kJ/kg FDM).

Keywords: Pleurotus ostreatus, solid fermentation, edible fungus, stoichiometry

Introduction
The Earth produces an estimate of 146 billion tons of living matter, known as Biomass, yearly [1]. Those resources have a great potential to face the challenges of decreased availability of traditional non-conventional fuel sources and to prevent the risks to stable food supply in most countries.

One of the possible ways to take advantage of those wastes is to use them for edible fungi production [2]. A quarter of all the cereal wastes annually discarded could be used to produce fresh edible fungi, enough for a daily supply of 250 g to more than four million people [3]. There is a very significant increase in mushroom and truffles production worldwide. According to the analysis by Toland and Lucier [4] on the data provided by the United Nations Food and Agriculture Organization (FAO), the worldwide production of that food rich on proteins and nutrients has exponentially raised from 1961 to 2009, and it duplicates every 1.03 years. Pleurotus ostreatus, also known as oyster mushroom, is one of the most extensively cultivated either in warm or hot climates. However, mushroom cultivation techniques are mostly empirical and many of the methods remain unpublished or are protected by several patents [5-14]. Therefore, mechanistic approaches are demanded to develop more efficient biological processes for that purpose [15]. Such procedures comprise the expression of process’ mathematical models and implementing optimization procedures based on computational modeling [16]. In this sense, process design has become a mathematical programming task, being identified by several authors as a design in transit from in vitro to in silico [15, 17, 18].

One of the primary contributions to bioprocess modeling comes from identifying a stoichiometric model by establishing the mass relationships among the main raw materials [19]. These models result in, either, a simplified analysis expressed in terms of a global stoichiometric equation describing the process from a non-structured perspective, or more complex ones, with a system of several equations describing the balances of cellular metabolism and cell-culture medium interactions [19, 20].

Following this strategy and after an unsuccessful exhaustive review of the literature on this topic, we decided to develop a stoichiometric equation to describe the growth of the P. ostreatus basidiocitete on industrial wastes. Thus, a stoichiometric equation was obtained, describing the growth of the P. ostreatus Ecuadorian strain ceba-glie-po-010606 on picking beans’ harvest waste.

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Materials and methods

Strain
The P. ostreatus Ecuadorian strain ceba-gliiepo-010106 was provided by the Ecuadorian Center of Environmental Biotechnology (CEBA). It was preserved in Agar-Malt extract at 4 °C until use.

Raw materials
Two raw materials were used to develop the empirical formulas: picking beans (Phaseolus vulgaris) harvest wastes, the entire plant, and the P. ostreatus biomass. Picking beans waste samples were used as primary raw material, being collected from the production communities La Concepción, Salinas and Ambuquí at the Imbabura and Carche provinces.

Fresh picking beans waste sampling procedure
Both, the picking beans waste and the mushroom biomass were sampled following the procedures described by Oakland [21]. For three months, three 14-kg samples of picking beans waste were collected every month, sun-dried after harvesting, up to 9. The monthly processing procedure is described in the following. The three samples were separately grinded to a 9-mm size in a hammer hog and further mixed. The mix was extended on the floor in a 1 m² square area. The square was divided in four even quadrants, and a 1 kg portion of the mix was collected from each quadrant, and further grinded with a manual grinder down to a 1 mm in size. It was extended again in the floor in a 1 m² square area, divided in four even quadrants. Samples of 25 g were collected from each quadrant and mixed for a final sample of 100 g for each month. Samples were packed in polystyrene bags and stored at 4 °C.

P. ostreatus biomass sampling procedure
The monthly-harvested picking beans waste was used for mushroom production at the CEBA production plant. The biomass produced by the three first sprouts of the fruiting body was stockpiled, during 51 days. Thereafter, mushroom growth was insignificant. The growth medium for fungal biomass production was used without nutrient supplementation. Three samples 1-kg each were harvested every month, for a final amount of 9 kg. The mushroom biomass obtained every month was dried until constant weight in an oven at 70 °C. It was further grinded in a manual grinder and extended on a 1 m² square table, which was subdivided in four even quadrants. Then, 25 g were collected from each quadrant and mixed to homogeneity. The three final samples of 100 g each were stored at 4 °C.

Biomass culture procedure by solid fermentation
The picking beans waste samples were grinded down to 9 mm and the humidity was set to 70 %. Polypropylene bags were filled with 300 g of substrate and further pasteurized in a sterilizer at 103.46 KPa for 30 min. When the substrate reached room temperature, it was inoculated with 4 % dry base. Samples were incubated at a constant temperature of 20 °C for 51 days. The relative humidity was kept at an average of 85 % to avoid hyphal contamination, and after the sprout of the first primordium, it was raised to 90 %. Fifteen days later, the mushroom was harvested, with an average size of 8 cm. The experiment was conducted with three bags as replicates.

Elemental composition analytical technique
Samples were analyzed at the Center of Research Services and Chemical Analyses (CISAO), of the National University of Loja, Ecuador. The elemental analysis was done with a PerkinElmer, model 2400, series II equipment [22]. This technique provides the total content of carbon, hydrogen, nitrogen and sulfur for a wide range of samples either organic or inorganic, solid or liquid, by using the Pregl-Dumas' classical method [22, 23]. This method consists of combusting a sample of known mass at high temperature (approximately 900 °C) in the presence of pure oxygen. The process releases carbon dioxide, water and nitrogen. The gasses are passed through special columns which absorb the carbon dioxide and water. A column carrying a thermal conductivity detector separates the nitrogen from any carbon dioxide or water residue, and the resulting nitrogen content is measured. The instrument must be previously calibrated by analyzing a pure standard of known nitrogen content. Then, the signal of an unknown nitrogen concentration sample is measured by the thermal conductivity detector, and converted into the equivalent nitrogen content value [24, 25].

Calculation of raw materials empirical formulas
Data corresponding to carbon, hydrogen, nitrogen and sulfur were obtained from fresh picking beans waste and mushroom fruiting body biomass by the technique previously described. Oxygen content was calculated as the remaining elemental composition difference, in disregard of other elements present in significant amounts. Elements' concentrations in the solid phase were determined for the organic fraction once subtracted the ashes present in the sample. These data were used to calculate the relative atomic mass ratio for each element, by the ratio of the element's mass fraction over its atomic mass. The atomic mass ratio could be deduced from these relative atomic mass ratios, by dividing each element value over that of carbon, the final value being expressed per carbon atom-gram. The empirical formulas of the raw materials were determined by the thermal conductivity detector, by using a thermal conductivity detector separates the nitrogen from any carbon dioxide or water residue, and the resulting nitrogen content is measured. The instrument must be previously calibrated by analyzing a pure standard of known nitrogen content. Then, the signal of an unknown nitrogen concentration sample is measured by the thermal conductivity detector, and converted into the equivalent nitrogen content value [24, 25].

Figure. Raw materials. A) Picking beans (Phaseolus vulgaris) waste, whole plant. B) Oyster mushroom (Pleurotus ostreatus). C) Biomass samples ready for elemental analysis.

materials used on this work did not include sulfur, due to its very low proportions and, therefore, its irrelevance for stoichiometric balances.

Proposal of a simplified stoichiometric model

It started considering the aerobic cell growth as a simplified mechanism, expressed through a simplified stoichiometric equation. It describes the transformation of fresh picking beans substrate, together with a nitrogen supplement in form of ammonia and oxygen as reactants, into products due to the presence of the mushroom as biocatalyst. The resulting reaction products considered were the *P. ostreatus* biomass, CO$_2$, and water:

$$\alpha CH_{4x}O_{y}N_{z} + \beta NH_{3} + \gamma O_{2} \rightarrow \chi CH_{2}O_{p} + \nu N_{2} + \delta CO_{2} + eH_{2}O \tag{1}$$

The stoichiometric coefficients were determined from a balance per element from equation (1) [19]:

- $\alpha$: hydrogen content on the molecule per carbon atom-gram.
- $\beta$: oxygen content on the molecule per carbon atom-gram.
- $\gamma$: nitrogen content on the molecule per carbon atom-gram.

For carbon, 4 electrons are free electrons during the full combustion of a given compound to render CO$_2$, H$_2$O and N$_2$. For carbon, electrons are free electrons to be transferred, with 1 for hydrogen, -2 for oxygen and -3 for nitrogen [19]. Positive and negative values stand for electrons either to be donated or accepted, respectively. Accordingly, the reduction degree for CO$_2$, H$_2$O and NH$_3$ is null. An electron balance from equation (1) leads to the following expression:

$$\chi: x_1 + 3 \beta = x_2 + 2 \varepsilon \tag{2}$$

$$O: y_1 \alpha + 27 = y_2 + 28 + \varepsilon \tag{3}$$

$$N: z_1 \alpha + \beta = z_2 \tag{5}$$

Where:
- $\alpha$, $\beta$, $\varepsilon$, $\chi$, $y$ and $z$: stoichiometric coefficients expressed as moles of the respective compound per mol of fungal dry matter (mol/mol FDM).
- $x$: number of electrons to be transferred to oxygen during the reaction [27].
- $y$: air oxygen fraction (mol O$_2$/mol total).
- $\varepsilon$: air density (kg/m$^3$).
- $RC$: respiration coefficient.

Results and discussion

Determining the empirical formulas for biomass of the fresh picking beans waste and the *P. ostreatus* mushroom is highly relevant to establish the stoichiometric equation describing the growth of the selected *P. ostreatus* strain, and also for the stoichiometric equation balance and the calculation of its coefficients. Table 1 summarizes the elemental composition of collected samples measurements and the estimated values for relative atomic mass ratios of the fungal biomass.
The *P. ostreatus* nitrogen levels were among the highest for a biomass tested [28], as expected from a fungal biomass. Instead, carbon levels were low, if we compare them to a list of elemental composition reported for forty agriculture matters [29]. This could explain the relatively high atomic mass ratio calculated for the oxygen, almost twice the value reported for many microorganisms. The ashes content was comparable to the lowest values reported by Sánchez and Mata [30] for eleven edible fungi species, and was closer to that of *Pleurotus sajou-caju* (5.84%).

Results for elemental composition of collected samples measurements and the calculation of relative atomic mass ratios of the fresh picking beans waste are shown in table 2.

The chemical carbon composition of the fresh picking beans waste was in the same order of the values reported by Parikh et al. [28] for biomass grown on waste from different sources. According to that, the carbon content was only higher than those reported for rice hull (40.6 %) and cotton stems (41.3 %). On the contrary, hydrogen and oxygen contents were highest for a biomass tested [28], as expected from rice hull (40.6 %) and cotton stems (41.3 %). As predicted, the calculated elemental composition does not vary for the most dissimilar biomasses; but growth conditions, either nutritional or environmental, are responsible for small variations [28].

There was a marked difference in the elemental composition of *P. ostreatus*, compared to that reported for *Aspergillus niger* (CH\(_{1.72}\)O\(_{0.55}\)N\(_{0.17}\)) [31, 32], which was quite more similar to that of the *Kluyveromyces marxianus* yeast (CH\(_{1.04}\)O\(_{0.16}\)N\(_{0.17}\)) [33]. The elemental composition does not vary for the most dissimilar biomasses; but growth conditions, either nutritional or environmental, are responsible for small variations for certain substrates and strain specific conditions. Several authors have optimized the growth medium for different *P. ostreatus* strains [34]. Nevertheless, the maximal efficiency reported for the process at 20°C was 261.89 g FDM/kg SDM [35]. Hence, these results for the system equations (8) through (12) were used to formulate the stoichiometric equation. Due to its simplicity, only two digits were reported for the stoichiometric coefficients (17):

**Theoretical biological efficiency**

Taking into account the amount of ashes and water of picking beans waste and *P. ostreatus* biomass, it was estimated that the expected biological efficiency from this system is 867.49 g FDM/kg SDM. This result arises as a model for the efficiency that must be achieved for certain substrates and strain specific conditions. Several authors have optimized the growth medium for different *P. ostreatus* strains [34]. Nevertheless, the maximal efficiency reported for the process at 20°C was 261.89 g FDM/kg SDM [35]. Hence, these levels could have been further improved, since they represent just 30.18 % of the estimated theoretical efficiency from tables 2 and 3.

**Table 1. Elemental composition dry base for the Pleurotus ostreatus biomass**

<table>
<thead>
<tr>
<th>Element</th>
<th>Mass percentage (dry base)</th>
<th>Atomic mass (g/mol)</th>
<th>Relative atomic mass ratio (mol/g)</th>
<th>Atomic mass ratio (mol/mol-C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>36.132</td>
<td>12.011</td>
<td>3.080</td>
<td>1.000</td>
</tr>
<tr>
<td>H</td>
<td>5.552</td>
<td>1.008</td>
<td>5.508</td>
<td>1.831</td>
</tr>
<tr>
<td>N</td>
<td>11.066</td>
<td>14.007</td>
<td>0.790</td>
<td>0.263</td>
</tr>
<tr>
<td>S</td>
<td>0.355</td>
<td>32.065</td>
<td>0.011</td>
<td>0.004</td>
</tr>
<tr>
<td>O</td>
<td>40.355</td>
<td>15.999</td>
<td>2.522</td>
<td>0.838</td>
</tr>
<tr>
<td>Ashes</td>
<td>6.540</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Total</td>
<td>100</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

**Table 2. Molar elemental composition of the fresh picking beans (Phaseolus vulgaris) waste**

<table>
<thead>
<tr>
<th>Element</th>
<th>Mass percentage (dry base)</th>
<th>Atomic mass (g/mol)</th>
<th>Relative atomic mass ratio (mol/g)</th>
<th>Atomic mass ratio (mol/mol-C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>39.875</td>
<td>12.011</td>
<td>3.464</td>
<td>1.000</td>
</tr>
<tr>
<td>H</td>
<td>6.047</td>
<td>1.008</td>
<td>6.261</td>
<td>1.807</td>
</tr>
<tr>
<td>N</td>
<td>6.727</td>
<td>14.007</td>
<td>0.501</td>
<td>0.145</td>
</tr>
<tr>
<td>S</td>
<td>0.201</td>
<td>32.065</td>
<td>0.007</td>
<td>0.002</td>
</tr>
<tr>
<td>O</td>
<td>42.980</td>
<td>15.999</td>
<td>2.543</td>
<td>0.809</td>
</tr>
<tr>
<td>Ashes</td>
<td>4.170</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Total</td>
<td>100</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

**Table 3. Empirical formula and molecular masses estimated for the picking beans (Phaseolus vulgaris) waste and oyster mushroom (Pleurotus ostreatus) biomass**

<table>
<thead>
<tr>
<th>Product</th>
<th>Composition (mass percentage, dry base)</th>
<th>Formula</th>
<th>Molecular mass (g/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P vulgaris waste</td>
<td>39.875 C: 6.047 H: 6.727 O: 42.980 S: 0.210</td>
<td>CH(<em>{1.82})O(</em>{0.86})N(_{0.26})S (\pm 0.08)NH(_2) + 0.355CO(_2)+ 0.355H(_2)O (\pm 0.27)</td>
<td>28.804</td>
</tr>
<tr>
<td>P ostreatus</td>
<td>36.132 C: 5.552 H: 11.066 O: 40.356 S: 0.355</td>
<td>CH(<em>{1.83})O(</em>{0.84})N(_{0.26})S (\pm 0.08)NH(_2) + 0.355CO(_2)+ 0.355H(_2)O (\pm 0.27)</td>
<td>30.949</td>
</tr>
</tbody>
</table>

\[ \beta = 0.0790; \gamma = 0.3494; \delta = 0.2689; \varepsilon = 0.3495. \]
value according to our results. Noteworthy, the biological efficiency depends not only on the attained nutritional balance, but also on other environmental aspects such as: the water retention capacity of the substrate, aeration and the relative humidity at different culture phases, among others [3].

**Mean coefficient of breath**

As defined for this parameter, the coefficient reached 0.77 mol CO₂/mol O₂, a relatively low value compared to that reported for the fungal aerobial growth. The breath coefficients commonly derive from increased demands for energy to synthesize the enzyme complexes required to produce simple sugar molecules during growth, as reported for *A. niger* growth on citric wastes [36]. Another plausible explanation comes from the CO₂ retention that occurs in growth bags, which could slightly modify the fungal growth metabolism.

**Specific air consumption**

The air consumption was calculated for normal temperature and pressure conditions, being estimated as 1.36 m³/kg FDM. This indicator demonstrates that the fungal growth demands a non-significant amount of air to produce a ton of products.

**Metabolic heat**

According to equations (10) and (17), metabolic heat was estimated as the heat release equivalent to 16 576.47 kJ/kg FDM. A huge amount of energy is normally released in the form of heat during solid fermentation processes due to metabolic activity [37]. This parameter is scarcely calculated from stoichiometric balances [38] similar to those developed in our work. For example, González *et al.* reported metabolic heat values for the growth of *A. niger* of 16 000 kJ/kg FDM, very similar to ours. This implies that the adjusted stoichiometric equation generates values which are in agreement with those previously reported for fungal growth.

**Conclusions**

A proximate stoichiometric equation was developed to describe the aerobial growth of the Ecuatorian *P. ostreatus* strain ceba-glie-po-010106 on local picking beans (*Phaseolus vulgaris*) harvest wastes. Empirical formulas were established to describe the elemental composition of fresh sun-dried picking beans waste (CH₁.₈₁O₀.₈₁N₀.₁₅) and that of the mushroom biomass (CH₁.₈₃O₀.₈₄N₀.₂₆). In this paper, the results of the proposed stoichiometry supports the estimation of key parameters for the development of fungal solid fermentation processes based on mathematical models with a mechanistic approach. They included the theoretical biological efficiency (867.49 g FDM/kg SDM), the mean respiration coefficient (0.77 mol CO₂/mol O₂), the specific air consumption to grow (1.36 m³/kg FDM) and the metabolic heat (16 576.47 kJ/kg FDM).

**Acknowledgements**

The research resulting on this publication was funded by the National Secretary of Higher Education, Science, Technology and Innovation of Ecuador (SENESCYT), through a Scholarship granted in 2008.

**Declaration of conflict of interests**

The authors declare the absence of conflicts of interests.

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