Influence of mashing profile curve and addition of proteases on the composition of the wort and beer

Abstract

Beer is the product resulting from the fermentation by brewer’s yeast of the wort obtained from malted cereal (barley), which may or may not be added by other cereals or sugar sources (called ‘adjuncts’), with the addition of hops. Depending on the raw materials used and the execution of its production process, it is possible for the consumer to appreciate different types and styles of beer, in addition to possible changes in the quality of the drink.1-3

For a long time, beer production on artisanal scale was sufficient to supply the drink demand. However, throughout its history beer became a popular drink, becoming today the most consumed alcoholic beverage in the world, having as a consequence the production of beers in large scales, a factor that characterizes the brewer market until nowadays. Due to the great competition in the market and the need to supply to the demand with agility, the large industries usually make diverse adjustments, aiming at the reduction of the process costs or of the production times.

As an example, significant changes in the mash curve profile, such as suppression of the time of protease activity to reduce time and energy expenditure in the process. The profile of the mash curve has a great influence on the composition of the wort and beer, affecting the nitrogen content and the type of nitrogen compounds present, with effect on several characteristics of the drink, such as the flavor (products and byproducts of the fermentation and permanence of non-hydrolyzed insoluble substances); the color (mainly by the release of soluble nitrogen and reducing sugars that will undergo subsequent parallel reactions); the texture (by the presence of proteins and dextrins); foam stability and beer turbidity (also influenced by the protein profile present); caloric power; the alcohol content; and the beverage carbonation.4-5

The reduction of mashing time usually promotes problems for the wort and beer, and it is common supplementing the wort with the addition of exogenous enzymes to avoid it. Mainly microbial amylase and proteases are added to improve the hydrolytic process.6 Thus, there are important changes in the process, such as: increase in the yield of the mash and the free amino nitrogen content (FAN) present in the wort, as well as increased ethanol production and fermentation productivity.6-12

The objective of this study was to evaluate the chemical and physicochemical composition of sweet worts and beers produced from different mash profiles, with and without the addition of exogenous enzyme extract, rich in proteolytic activity.

Materials

The following raw materials, additives and fermentation agents were used to obtain worts and beers: water filtered on activated charcoal, Pilsen malt (Agromalte®), hops (Hallertau Perle, HGV®), 9.7% α-acids), proteolytic extract of microbial origin, obtained in previous work (Mathias et al., 2017) and bottom fermenting yeast-Saccharomyces cerevisiae (W-34/70, Fermentis®).
For the amino acid quantification in sweet wort, by capillary electrophoresis, namely cysteine (Cys), histidine (His), phenylalanine (Phe), lysine (Lys), tryptophan (Trp) and arginine (Arg), respective analytical standards were used (≥99%, USP, USA) and ultrapure water (resistivity of 18.2 MΩcm) produced by the Arium Confort II system (Sartorius, Germany). Sodium phosphate, cetyltrimethyl ammonium bromide (CTAB) and sodium hydroxide (Merck, Brasil) were used in the preparation of the working electrolyte (BGE).

The pH of the BGE was adjusted using a pH meter (DM-22, Digimed, Brasil). All solutions (BGE, standard solutions and samples) were filtered using syringe filters (PTFE 0.45μm, Agilent, Germany) prior to introduction into the capillary electrophoresis system.

**Methodology**

**Wort and beer production**

All brewer’s worts production was carried out on a laboratory scale from a compilation of scientific literature data with a 1:4 (cereal: water) mixture, enriched with calcium chloride (0.125g/L). The pH of the medium had its value adjusted to range from 5.2 to 5.4 by the addition of lactic acid P.A. when necessary.

The influence of the addition of exogenous proteolytic extract as well as step suppression of protease activity (inherent to the malt) during the mash was verified. For this, a factorial experimental design was used, with two factors (proteolytic step and protease addition) at two levels (with step/without step - with enzyme/without enzyme), resulting in four experiments, as shown in Table 1. Concentration of enzyme extract was fixed to confer a standard amount of enzyme activity (0.75U/g malt) and the experimental conditions varied only qualitatively by its addition or not. Figure 1 briefly shows the mashing curves.

**Table 1** Experimental design for brewery worts production

<table>
<thead>
<tr>
<th>Proteolytic Step</th>
<th>Proteolytic Extract</th>
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<tbody>
<tr>
<td></td>
<td>t (°C) t (min)</td>
</tr>
<tr>
<td>Beer 1</td>
<td>45 15</td>
</tr>
<tr>
<td></td>
<td>add (+)</td>
</tr>
<tr>
<td>Beer 2</td>
<td>55 15</td>
</tr>
<tr>
<td></td>
<td>add (+)</td>
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<tr>
<td>Beer 3</td>
<td>45 15</td>
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<tr>
<td></td>
<td>not add (-)</td>
</tr>
<tr>
<td>Beer 4</td>
<td>55 15</td>
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<td></td>
<td>not add (-)</td>
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</table>

The mashout was established by the iodine test (alcoholic iodine solution in 2% potassium iodide), when it was considered the disappearance of the intense purple coloration (which characterizes the presence of starch) and appearance of yellowish color, typical of iodine solution. After mashing, the wort was clarified using a sieve (0.6-0.7mm), and the brewer spent grain was washed with heated water (78°C), with volume equal to mash water initially used. The wort was centrifuged at 1300g for 10min for further procedures.

The clarified wort obtained was send to boiling step during 60minutes with the heating power adjusted to promote no more than 10% (v/v) water loss by evaporation. Simultaneously, hops were added and hop mass was calculated as a function of the volume of wort at the start of the boiling, based on Klopper’s empirical table, which shows the theoretical equivalence between the concentrations of α-acids added to the wort and the bitterness (in BU) generated in the drink, considering an isomerization efficiency of 30%. It was planned to obtain a wort with 9° Plato and 12BU, which characterize a typical American Light Lager beer.

**Figure 1** Mash profiles. (A) with proteolytic step, to obtaining Beers 1 and 3; (B) without proteolytic step, to obtaining Beers 2 and 4.

After the boiling time, the protein coagulation (hot trub) was deposited by strong manual stirring in a circular movement, to act the centripetal force, reproducing the effect of the equipment adopted in the industrial practice (whirlpool).

After removal of the hot trub, by filtering, the wort was cooled and inoculated with lyophilized brewer’s yeast, at a concentration of 1g/L of wort, properly hydrated in 10ml of cold and sterile water, for each gram of yeast. The fermentation was conducted for 6days at a temperature of approximately 15°C. After this time, the decanted yeast mass was removed and the green beer matured for 7days at low temperatures (4°C).

**Analysis of worts and beers**

The sweet worts produced from different time/temperature combinations and added or not of exogenous proteolytic extracts, were submitted to the following analyses: °Plato by refractometric measure; extract (% m/m of dissolved solids); total nitrogen content by Kjeldahl method; FAN-free amino nitrogen content by ninhydrin method. In addition, the nitrogen content lost by coagulation during boiling was determined by the difference between the total nitrogen contents before and after this stage.

The amino acids cysteine, histidine, phenylalanine, lysine, tryptophan and arginine were quantified in sweet worts by capillary electrophoresis in commercial EC 7100 equipment (Agilent, Germany) equipped with a diode arrangement spectrophotometric detector. Separations were conducted in a 60cm total length fused silica capillary (51.5cm to the detector-effective length) and 50μm internal diameter. Samples were introduced by hydrodynamic mode at 50mbar pressure for 15s and optimized conditions of the method were: 50mol L^-1 phosphate buffer containing 0.4mol L^-1 CTAB with pH adjusted to 12.5 as BGE; applied potential of-20kV and temperature of 20°C. The wavelengths used for detection were 200nm (Lys and Arg); 220nm (Trp, His and Phe) and 230nm (Cys).
The capillary electrophoresis method validation was performed according to the International Conference on Harmonization ICH Q2. The ANVISA Resolution No. 089/2011 and DOQ-CGCRE-008 (“Guidance on validation of analytical methods”) of the National Institute of Metrology, Standardization and Industrial Quality (INMETRO). The beers obtained were analyzed for alcohol content; real extract; and fermentation efficiency in specific equipment (Alcolyzer Beer Analyzer, Anton Paar, DMA 4500M). In addition, total nitrogen content were determined by Kjeldahl method. The addition of exogenous proteolytic enzymes had a greater effect when the worts were not submitted to the step of proteases (from the malt) during the mash. With this result, it is assumed that the proteolytic step is sufficient by itself, and the addition of exogenous enzymes is necessary when it is suppressed, aiming to reduce the process time. It is observed that with the exogenous proteolytic extract and without the protease step (B2 wort), an extract value similar to that produced with no exogenous proteolytic extract and protease step was obtained, with a reduction of the process time in 30 minutes.

For the total nitrogen content, there is a small increase, although statistically significant (Turkey’s test, Table 2), of the B2 wort relative to the B4 wort. This suggests that the added proteases promoted greater extraction of the nitrogen compounds from the malt to the wort. Even with the low performance of the proteases on the insoluble protein fraction of the malt, the addition of the exogenous proteolytic enzymes promoted changes in the composition of the sweet wort, fundamental for the profile of the soluble nitrogenous compounds present.

The comparative analysis of free amino nitrogen between B2 and B4 worts shows that the highest value was determined when there was supplementation of proteases in the sample (330 and 212 mg/L, respectively), indicating a compensation for proteolytic step suppression. In addition, observing the FAN results for B2 and B3 worts, an expressive increase of this content was observed, even with a reduction of 30 minutes of the processing time. In general, in the condition of low amino acid availability, brewer’s yeast presents metabolic alterations, which results in the generation of different fermentation by-products. In this way, the supplementation of proteases on mash may have a positive effect on sensorial quality of the beer.

Differential losses of total nitrogen during boiling of sweet worts are also evidenced. In the presence of exogenous proteases (B1 and B2) there were smaller percentage losses of this content (Table 2). It is known that during boiling stage, high molar mass proteins tend to lose salvation water, coagulating, in addition to being complexes with each other, with the formation of the so-called hot trub, insoluble, that precipitates and is removed from the wort. In this way, the addition of exogenous proteolytic enzymes during mash may have promoted a reduction in the molar mass of the proteins present, reducing the intensity with which this coagulation occurred at high temperature. The lower the loss of nitrogen compounds during boiling, the better the nutritional quality of the wort, the performance of the brewer’s yeast, and the final product.

In addition, specific amino acids (cysteine, histidine, phenylalanine, lysine, tryptophan and arginine) were determined in sweet worts by capillary electrophoresis. These amino acids are assimilated by yeast, and the final product.

Figure 2 illustrates the different amino acid compositions in the worts observed in this study. The results indicate that the wort B1 has
higher concentrations of the amino acids studied; in addition, the amino acid cysteine was only detected in this wort. The concentrations of the amino acid tryptophan in wort B1 and B3 are statistically equivalent, according to the comparison between the mean experimental concentrations by Student’s t-test (95% confidence level). The amino acid content in B2 and B4 worts shows that the highest values were determined in the wort produced with the addition of exogenous proteases (B2 wort), the same result observed when comparing the B1 (with protease) and B3 (without protease).

These results confirm that the addition of the proteolytic extract had an effect on the protein fraction of the malt, increasing the content of amino acids and/or other nitrogen compounds of low molar mass, or changing the profile of the nitrogen compounds.

Table 3 presents the parameters of analysis of the four beers produced. There is little variation between the final extracts, although the alcoholic contents were very similar. These values allowed calculating the fermentation efficiencies, whose highest values corresponded to conditions B2 (without proteolytic step and with enzyme supplementation) and B3 (with proteolytic step and without protease supplementation). Additionally, comparing B2 and B4, which were differentiated by the addition of exogenous protease, an increase in fermentation efficiency was observed. Indicating, again, the improvement of the process when there are proteases be they exogenous or endogenous. It is noteworthy that, for B2 beer, the same fermentation efficiency was observed, with a process time reduced by 30 minutes (in relation to B3 wort).

It is evident the correlation between total nitrogen and FAN content, with the highest values for beers B1 and B3, both obtained from wort without protease supplementation. However, the highest consumption of amino acids in the fermentation was observed for beer B2, produced by mashing without proteolytic step and with protease supplementation.

Table 3 indicates that the greater mass of coagulable matter by boil was observed for B4 beer (without proteolytic step and without protease supplementation), followed by beers B3, B2 and B1, in this order, the last three being very close values. The coagulable matter can be associated to the coagulation of proteins of high molar mass, being the reduction of its concentration beneficial for the colloidal stability of the beverage, avoiding its early turbidity during the period of transport and storage. Therefore, the results, already expected, indicate the positive effect of both the step of proteases and the supplementation with exogenous proteases on the nitrogenous compounds present in the wort and beer.

**Table 3 Characterization of beer**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Beers**</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>B1</td>
</tr>
<tr>
<td>Er (% m/m)</td>
<td>3,1</td>
</tr>
<tr>
<td>N_total (%)</td>
<td>0,027</td>
</tr>
<tr>
<td>FAN (mg/L)</td>
<td>118</td>
</tr>
<tr>
<td>Ethanol (%v/v)</td>
<td>3,6</td>
</tr>
<tr>
<td>Coagulable matter (g/L)</td>
<td>4,5</td>
</tr>
<tr>
<td>Fermentation efficiency (%)</td>
<td>92</td>
</tr>
</tbody>
</table>

Some authors have evaluated the commercial proteases supplementation for hydrolysis of amylaceous raw materials, whether for the production of beverages or ethanol fuel, in the high gravity process. In the case of wheat, corn or traditional barley malt, responses obtained were: increase in amino acid concentration; increase in the mashing yield; increased efficiency and productivity of alcoholic fermentation; and, in some cases, the increase of the cellular biomass at the end of the fermentation.

**Conclusion**

In summary, the results indicate that the addition of the proteolytic extract had an effect on the protein fraction of the grain, with significant changes in the characteristics of the wort and beer. Its addition promoted increase of the extract, total nitrogen and amino acid content in the sweet wort, and a higher fermentation efficiency. Additionally, there was a reduction of the nitrogen loss in the boiling stage and of the coagulable matter content in beer.

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**Conflict of interest**

The author declares no conflict of interest.

**References**


22. ICH. Validation of Analytical Procedures: Text and Methodology, Q2 (R1); 2005.


