Study of antioxidant activity during the malting and brewing process

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Abstract In this study the evolution of antioxidant activity was investigated during malting of different barley cultivars, and during the production of different types of beers on laboratory scale and in pilot brewery. Samples were taken at technologically important points of productions. Malts were produced from 3 spring and 3 winter barley cultivars. Two types of beers were brewed under laboratory conditions, and two in a pilot brewery. For the determination of antioxidant activity five commonly used assays were applied such as ABTS Radical Scavenging Activity, Cupric Reducing Antioxidant Capacity, DPPH Radical Scavenging Activity, Ferric Reducing Antioxidant Power and Total Polyphenol Content. Prior to malting it was observed that there are orders of magnitude differences between the antioxidant activities of the barley varieties. During malting, the biggest increase was noticed during steeping. Spring and winter cultivars showed similar trends during steeping and germination, but kilning had different effect on antioxidant activity of the varieties. The antioxidant activity of malts was always higher than the corresponding barleys. During the brewing process antioxidants were released to the highest extent during the early stages of mashing. Adequate sparging and hop boiling could further improve the antioxidant potential of the wort. Furthermore, differences between the equipment used for wort separation and hop boiling under laboratory conditions and in the pilot brewery had effect on antioxidant activity. In the course of malting and brewing by selecting the appropriate raw materials and technological parameters, the conditions for the release and retention of antioxidants can be optimized.

Keywords Beer · Malts · Brewing · Malting · Antioxidant activity

Introduction

Technological steps of malting and brewing do have a significant effect on the composition of malts and beer made of them. These steps not only influence the extract, alcohol or protein content of the final product but the bioactive components like antioxidants, as well. This study focuses on the evolution of antioxidant activity during malting of different barley cultivars, and during the production of beers made of different types of malts in laboratory and in pilot brewery.

Antioxidants

Antioxidants are important compounds which help us to retain our health. The main role of antioxidants in human health is to attenuate oxidative stress. Oxidative stress arises from overproduction of reactive oxygen or nitrogen species (ROS/RNS). These free radicals are produced under normal physiological and pathological conditions in our organism and play an important role in pathological
processes and regulatory activities. Antioxidants can act in different ways, they can scavenge free radicals, inhibit pro-oxidative enzymes, chelate metal ions, among others. Grains are excellent sources of antioxidants, such as vitamin E, polyphenols, phytic acid, folates and microelements (e.g. zinc, selenium) (Shahidi et al. 2012).

**Determination of antioxidant activity**

According to Huang et al. (2005) there are two categories of assays which are applicable to determine antioxidant activity, these categories are hydrogen atom transfer reaction-based assays and single electron transfer reaction-based assays. In this study electron transfer based assays were applied. These methods involve two components in the reaction mixture, antioxidants and oxidant. They are based on a reaction in which the probe (oxidant) is reduced by the antioxidant (present in the sample) that cause color change which can be measured by a spectrophotometer. The degree of the color change is proportional to the antioxidant concentrations. The reducing capacity of the antioxidant is usually expressed as Ascorbic acid equivalent (AAE), Trolox equivalent (TE) or gallic acid equivalent (GAE). These assays have the limitation that they are not selective to certain compounds, only suitable for determining the reducing capacity of the sample. On the other hand, significant correlation can be found between antioxidant activity and certain groups of antioxidants, e.g. polyphenol content in beer (Zhao et al. 2010).

**Antioxidant potential of barley and malt**

Malting barley is used in the second highest proportion after water in brewing. The most significant antioxidants in barley are polyphenols. Many scientific papers have reported that the long-term consumption of plants rich in polyphenols protects us against development of cancers, cardiovascular diseases, osteoporosis, diabetes and neurodegenerative diseases (Graf et al. 2005). Phenolic substances are an important part of the natural defense system of plants, protecting them from bacterial and fungal pathogens, insects and herbivores (Zimmermann and Galensa 2007). The phenolic content of barley is influenced by biotic and abiotic factors, which affect plant physiology and secondary metabolites (Mikkelsen et al. 2015). Phenolic compounds are predominantly found in the outer layers of the grain (husk, pericarp, testa, aleurone cells) bound to cell wall polysaccharides (Naczk and Shahidi 2004; Kähkönen et al. 1999). During malting, the extractability of these compounds is increasing mainly due to enzymatic processes and better friability. The other group of potential antioxidant compounds is developed during the heat treatment of green malt. In case of special malts (e.g. caramel, coloring) which are treated at high temperature (150–200 °C) at the end of the malting process, Maillard reaction takes place intensively resulting in products that do have antioxidant activity (Carvalho et al. 2015).

**Antioxidant potential of hops**

Hops, the other important ingredient of beer regarding to antioxidants, contain higher concentrations of polyphenols than barley malt (barley malt 50–100 mg/100 g, hops up to 4 g/100 g), but barley malt contributes 70–80% of total polyphenols in traditional beers (Almaguer et al. 2014; Narziss 1976).

**Effect of technological steps during brewing on antioxidants**

There have been studies investigating antioxidants from many aspects related to brewing. Schwarz, Boitz and Methner (2012) studied how the mashing-in temperature influences the release of polyphenols. They found that 40–45 °C is ideal for phenolic acid release from malt, while at temperatures above 65 °C no enzyme activity related to release of phenolic acids was detected. Fumi et al. (2011) studied polyphenols in all-malt worts and in maize adjunct worts, and their fate during the main brewing steps. They observed higher phenolic content in all-malt worts than in worts with maize adjunct, furthermore they reported that the overall brewing process reduces by 50% the initial content of total phenols. Zhao (2015) investigated the effects of processing stages on the profile of phenolic compounds from barley to the final product. It was found that their amount had generally increased significantly during malting and mashing but decreased markedly during the subsequent fermentation and storage. Pascoe, Ames and Chandra (2003) studied the effect of critical stages of the brewing process on antioxidant activity. They observed a decrease after beer filtration, and increase in levels of antioxidant activity after mashing, boiling, fermentation and chill-lagering.

The number of studies focusing on antioxidants related to brewing have increased in recent years proving the importance of this topic. With this study we would like to widen knowledge on the subject by comparing the antioxidant activity of spring and winter barley cultivars, and its evolution during the malting process. Furthermore, antioxidant activity during the whole brewing process of different beer types, produced using both basic and special malts, under laboratory conditions and in pilot brewery have not been investigated and compared yet.
Materials and methods

Barley samples

Six barley cultivars were involved in the study. Malts were produced from all of them. Among them there were 3 spring barley cultivars: Quench, Malz and Kangoo, and 3 winter barley cultivars: Casanova, Vanessa and Wintmalt.

Malting

The malting technology was as follows: steeping consisted of 3 sessions, each session lasted for 8 h (4 h of wet period and 4 h of air rest) at 16 °C. The germination lasted for 4 days, the temperature was 18 °C on the first day, 20 °C on the second day, 22 °C on the third day and 20 °C on the fourth day. Water was sprayed 5 times per hour onto the green malt during germination, and 30 rotations were set in every 2 h. The kilning began at 40 °C, the temperature was raised to 48 °C in 2 h, and finally cooled to 30 °C in 10 h.

Brewing

Four types of beers were brewed, two under laboratory conditions and two in a 50-liter capacity pilot brewery.

Beers brewed in laboratory

In laboratory two types of lagers (Pilsner and Vienna lager) were brewed. Samples were taken at the end of all enzymatic rests during mashing, after first wort separation, after hop boiling, on every day during germination and after kilning.

The malting technology was as follows: steeping consisted of 3 sessions, each session lasted for 8 h (4 h of wet period and 4 h of air rest) at 16 °C. The germination lasted for 4 days, the temperature was 18 °C on the first day, 20 °C on the second day, 22 °C on the third day and 20 °C on the fourth day. Water was sprayed 5 times per hour onto the green malt during germination, and 30 rotations were set in every 2 h. The kilning began at 40 °C, the temperature was raised to 48 °C in 2 h, and finally cooled to 30 °C in 10 h.

The brewing technology of beers brewed under laboratory conditions was as follows: mashing-in was carried out at 52 °C, then the temperature of the mash was held at 52 °C for 20 min, that was followed by a 45 min rest at 63 °C, and finally there was a 15 min rest at 73 °C. The temperature between the enzymatic rests was raised by 1 °C/min. The wort separation was carried out using Whatman MN-615 filter paper (GE Healthcare). The hop boiling lasted for 60 min in Erlenmeyer flask, at the 5th min 0.8 g Magnum hops was added. The wort was cooled to 12 °C prior to fermentation. The main fermentation lasted for 6 days at 12 °C followed by a chill-rest for 15 days at 5 °C.

Beers brewed in pilot brewery

In the 50 L capacity pilot brewery two types of ales—Brown ale and Stout—were produced. Samples were taken at the end of all enzymatic rests during mashing, after first wort separation, after sparging (from the sweet wort), after hop boiling and at every day of the main fermentation.

The recipe of the Brown ale was as follows: the water: malt ratio was 4:1. The malt composition was 60% Maris Otter pale, 15% Vienna, 10% Carapils, 10% Cara hell and 5% Chateau Special B. Hops: Challenger (8% α-acid), yeast: Safale S-04 (Fermentis).

The brewing technology of the Brown ale was as follows: mashing in was carried out at 52 °C, then the temperature was held for 20 min at 52 °C, then it was increased to 63 °C and held for 45 min, then increased to 73 °C and held for 15 min, finally the temperature was raised to 78 °C prior to mashing out. The temperature between the enzymatic rests was increased by 1 °C/min. The mash was pumped into the lauter tun where the hot trub was separated. The hopped wort was cooled to 21 °C prior to fermentation. The main fermentation lasted for 4 days at 21 °C.

The recipe of the Stout was as follows: the water: malt ratio was 4:1. The malt composition was 45% Maris Otter pale, 40% Smoked pale, 10% Cara Bohemian, 2% Carafa III. and 3% Chocolate malt. Hops: Warrior (17% α-acid), yeast: Safale S-04 (Fermentis).

The brewing technology of the Stout was as follows: mashing in was carried out at 45 °C followed by a 15 min rest at 45 °C, then the temperature was increased to 55 °C and held for 15 min, it was followed by a 45 min rest at 63 °C, then the temperature was raised to 73 °C and held for 15 min, finally the mashing out was carried out at 78 °C. The mash was pumped into the lauter tun and was followed by a 20 min sedimentation rest. Then the first
wort was separated and was followed by three times sparging. The hop boiling of the sweet wort lasted for 60 min, at the 5th min 20 g Warrior hops was added. The whirlpool, cooling and fermentation was the same as in case of Brown ale.

**Sample preparation and extraction**

The extraction of barley and malt samples for antioxidant capacity determination was as follows: for 2 g of finely ground sample, ground in an EBC mill, 20 mL of 80:20 Acetone: Distilled water solution was added. It was sonicated for 10 min in an ultrasonic bath, shaken for 60 min at 150 rpm and centrifuged at 2500 g for 10 min. The supernatant was collected and stored at –80 °C until analysis.

**Analyses of antioxidant activity**

The antioxidant activity was determined by five commonly applied assays as there is no standard method which can objectively characterize this parameter. The following assays were applied because these are widely used to determine this parameter so there is a possibility to compare our results with others’, furthermore these assays are easily reproducible, however are not selective to certain components, these methods are determining the reducing ability of the sample. (Huang et al. 2005) All the results were expressed as mg/100 g or mg/100 cm³ ascorbic acid equivalent (AAE), in case of malts related to dry matter (d.m.). All the measurements were carried out in three parallels.

**ABTS radical scavenging activity**

The assay was performed as described by Re et al. (1999). 10 µL degassed sample was pipetted into 96 well plates. 20 µL solution was added, which contained 9% NaCl, 1% glucose, 50 mg/mL myoglobin dissolved in pH 7.4 potassium-phosphate buffer. Then 150 µL 1 mg/mL 2,2’-Azino-bis (3-ethylbenzothiazoline-6-sulfonic acid) dianionium salt (ABTS) solution and 25 µL 3% H₂O₂ dissolved in 0.1 M pH 5 citric buffer was added. It was shaken for 15 min at 37 °C then absorbance was measured at λ = 405 nm.

**Ferric reducing antioxidant power**

The assay was performed according to Benzie and Strain (1996). Samples were added to FRAP reagent that contained 10 mM 2,4,6-Tris(2-pyridyl)-s-triazine (TPTZ) dissolved in 40 mM HCl, 300 mM pH 3.6 acetate buffer and 20 mM FeCl₃*6H₂O. After 5 min of incubation time, absorbance was measured at λ = 593 nm and.

**Total polyphenol content**

The assay was performed based on the description of Singleton and Rossi (1965), which is based on the reduction power of antioxidants rather than on the selective reaction of polyphenols, thus it was evaluated together with the other antioxidant activity assays (Martinez-Perihan et al. 2011). First 1250 µL ten-fold diluted Folin-Cioalteau reagent and 240 µL methanol: water (4:1) solvent were pipetted in the test-tubes. Then 10 µL degassed sample was added. After homogenization and 1 min reaction time 1 cm³ 0.7 M Na₂CO₃ was added, vortexed and before measurement the mixture was allowed to stand for 5 min at 50 °C. The absorbance was measured at λ = 765 nm.

**Cupric reducing antioxidant capacity**

The assay was performed according to Apak et al. (2004). 100 µL sample was added to 1 cm³ 10⁻²M CuCl₂, 1 cm³ 7.5*10⁻³M neocuproine solution (dissolved in 96% ethanol), 1 cm³ pH 7.4 1 M NH₄Ac buffer solution and 0.9 cm³ distilled water. It was incubated in dark for 30 min and the absorbance was measured at λ = 450 nm.

**DPPH radical scavenging activity**

The assay was performed as described by Brand-Williams, Cuvelier and Berset (1995). 6*10⁻⁵M 2,2-Diphenyl-1-picrylhydrazyl (DPPH) solution was prepared with methanol. 100 µL sample was added to 3.9 cm³ DPPH solution and was incubated in dark for 20 min then the absorbance was measured at λ = 517 nm.

**Real extract content**

The real extract content was determined by an Anton-Paar Alcolyzer Plus beer analyzer.

**Moisture content**

The moisture content of the barley and malt samples was determined by an AND MX-50 Moisture Analyzer.

**Results and discussion**

**Antioxidant activity of barleys and malts**

The results of relative antioxidant activity of the six investigated barley varieties, and malts produced from
them are shown in Fig. 1. The 100% values for the different methods were the following in mg/100 g Ascorbic Acid Equivalent (AAE) expressed on dry matter basis (d.m.): DPPH Radical Scavenging Activity (DPPH) = 24.4, Total Polyphenol Content (TPC) = 61.9, Cupric Reducing Antioxidant Capacity (CUPRAC) = 63.3, Ferric Reducing Antioxidant Power (FRAP) = 30.5, ABTS Radical Scavenging Activity (ABTS) = 155.6. As it can be seen in Fig. 1 there are major differences between the antioxidant activities of barley varieties, which is consistent with the results obtained by Zhao et al. (2008). These difference are due to biotic and abiotic factors, which affect plant physiology and secondary metabolites such as antioxidants (Mikkelsen et al. 2015). By regulating these factors, it may be possible to effect on this property of plants. Among barleys Casanova winter cultivar has the highest DPPH (19.8 mg/100 g AAE d.m.), CUPRAC (35.3 mg/100 g AAE d.m.), FRAP (22.2 mg/100 g AAE d.m.) and ABTS (109.4 mg/100 g AAE d.m.) while Kangoo spring cultivar has the lowest values. These results can be related to the generally thicker husk of winter barley varieties. Phenolic compounds, which do contribute to antioxidant activity, are predominantly found in the outer layers of the grain, so a thicker husk can come along with a higher antioxidant activity (Naczk and Shahidi 2004; Fogarasi et al. 2015).

All the antioxidant activity levels measured by the five assays have increased during malting. This result is in accordance with the observation of Fogarasi et al. (2015) who have experienced the same in case of barley, wheat and einkorn wheat. By producing malt from barley a valuable product is made from a nutritional point of view. The highest increase during malting can be observed in TPC values, which was reported by Pejin et al. (2009) as well. The biggest increase of TPC during malting was indicated by Casanova winter cultivar, from 8.1 to 61.6 mg/100 g AAE d.m. As this method is not selective to phenolic substances, rather shows the reducing capacity of the samples (Huang et al. 2005) it cannot be declared that it goes along with the increase of phenolic content. On the other hand, there are studies, which have proven that due to enzymatic release of bound phenolic compounds of barley and easier extractability lead to higher levels of free phenolics in malt compared to barley (Carvalho et al. 2015; Dvorakova et al. 2008). In case of TPC, CUPRAC and FRAP the highest values belong to the malt made from Quench spring barley, while malt made from Kangoo spring barley shows the lowest results in general. It may be due to the malting process as the same technology was applied for all the barleys, and this technology may not have been optimal for all the samples. These conditions favored the germination of Quench barley.

**Antioxidant activity during malting**

In Figs. 2 and 3 the results of antioxidant activity during the malting process of spring and winter barley cultivars can be seen. As we could see on Fig. 1, the antioxidant activity measured by all the assays are higher of malts than barleys, on the other hand it does not increase continuously through the entire process. Our DPPH results are very similar to the results of Pejin et al. (2009), an increase can be observed at the end of steeping, and on the first days of germination, and after that it starts to decrease. On the other hand Lu et al. (2007) have experienced a decrease at the end of steeping by ABTS, TPC and DPPH methods with a very similar sample extraction as ours. They have reported the highest increase of DPPH during kilning, which does not agree with either our or with Pejin et al. (2009) results. This is probably due to the differences in technology and equipment as in both Pejins’s and our study a Schmidt-Seeger micromalting plant was used while the equipment of Lu is unknown. Spring and winter cultivars showed similar tendency during the whole malting process.
as the same biochemical processes occur during their malting. The highest increase is observed during steeping. Antioxidants are more extractable as moisture content of kernels is raised from 11 to 12% (raw barley) to 40%.

During kilning the antioxidant activity assays show different tendencies. It is maybe due to the different sensitivity of the assays to the products being formed or degraded during kilning. According to Bellmer (1978) the kilning step is regarded important for polyphenol solubilization, Leitao et al. (2012) experienced the highest increase of total phenolic content during kilning.

**Antioxidant activity during brewing**

In Figs. 4 and 5 the results of antioxidant activity during the entire brewing process can be seen under laboratory conditions and in the pilot brewery. As can be seen, Pilsner beer has the lowest values through the entire process even though it has almost the highest original extract content among all the investigated beers, 16.51 °B, measured from its hopped wort (Table 1). Vienna lager has slightly higher antioxidant activity except for DPPH results, furthermore it has the highest original extract content, 17.28 °B of hopped wort. In general, Brown ale and Stout, that contain special malts have higher antioxidant activity, even though they have lower original extract content (Brown ale: 14.08 °B, Stout: 14.07 °B, also determined from their hopped wort) than the pale beers. From this it can be concluded that the extract content does not necessarily have an effect on antioxidant activity, much more, the malt composition affects this parameter. This result agrees with the result reported by Ditrych, Kordialik-Bogacka and Czyżowska (2016) who have found that darker beers have higher...
antioxidant potential than pale ones. This is mainly due to the special malts used for their production, as special malts contain more melanoidins, which contribute to the antioxidant activity (Zhao and Zhao 2012).

The TPC, CUPRAC, FRAP, DPPH values are relatively high after the first enzymatic rest at 45 °C (Stout) or 52 °C (Pilsner, Vienna lager, Brown ale). It is consistent with the result of Schwarz, Boitz and Methner (2012), who have reported the same temperature as ideal one for polyphenol release from malt, and with the results of Zhao and Zhao (2012) who have found that antioxidant activity increased the most intensive during the early stage of mashing. Until the end of the first enzymatic rest water-soluble antioxidants can go into solution, furthermore these temperatures are optimal for protease and β-glucanase enzymes of barley malt which can release antioxidants bound to cell walls, polysaccharides or proteins. By holding this temperature longer more antioxidants can be released, on the other hand the quality of the final product can be negatively affected due to excessive degradation of proteins.

The results given by ABTS assay at the early stages of the brewing process both under laboratory circumstances and in the pilot brewery were unexpected. ABTS does not show any relevant antioxidant activity until the end of the enzymatic rest at 73 °C but afterwards increases radically. In case of this assay the above elaborated theory about the importance of the protease, β-glucanase rest at the beginning of mashing is inappropriate. Compounds measured by this method can be either soluble only above 70 °C or they are formed at this higher temperature.

The wort separation using filter paper under laboratory circumstances caused an unexpected radical decrease in
was carried out in Erlenmeyer flasks while in the pilot brewery a steam-heated wort kettle was used for hop boiling and whirlpool for the separation of hot trub.

During fermentation the antioxidant activity decreased in some cases or showed no changes as other researchers have also reported. Fantozzi et al. (1998) observed decrease while Leitao et al. (2011) have found no significant changes. In our study we have not observed difference between the evolution of antioxidant activity during main fermentations performed under different circumstances: one in fermenters of the pilot brewery and the other in a flask with airlock.

The changes during chill rest were determined only in case of beers produced in the laboratory and. no significant changes have been indicated in antioxidant activity.

**Conclusion**

There are numerous studies focusing on the evolution of antioxidants during the entire brewing process but there are still gaps that need to be investigated. Brewing consists of very complex processes. The antioxidant potential of the final product depends already on the growing conditions of the raw materials used. This study revealed the differences of antioxidant activity of spring and winter barley cultivars. Despite the differences between barley varieties and despite similar trends of spring and winter cultivars during steeping and germination, kilning had different effect on antioxidant activity of the varieties. Spring and winter cultivars showed similar trends during steeping and germination, but kilning had different effect on antioxidant activity of the varieties. In the course of brewing differences between the equipment used in the laboratory and in the pilot brewery showed to have effect on antioxidant activity, especially during wort separation and hop boiling. During the mashing process the enzymatic rests at 45 °C or 52 °C are important in the release of antioxidants from malts. Together with adequate sparging and hop boiling mashing can contribute to the antioxidant activity of the final product. Sparging could be a determinative step as we can gain valuable compounds from the spent grain that would not be used in the further steps of beer production. During malting and brewing there are plenty of parameters that can be influenced by the recipe or technology to optimize the conditions for the release and retention of antioxidants. In the light of the complexity of beer production and wide choice of raw materials further research is needed to understand these processes better.

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