Characterization of arabinoxylan degrading enzymes during wheat malting

Bao Chu1*, Yanan Chai2*, Yuhong Jin*, Zhaoran Zhang3, Xiaoyang1, and Junhan Liu

1College of Food Science and Engineering, Shandong Agricultural University, No. 61 Daizong Street, Tai'an, 271018, Shandong Province, PR China.
2Department of Modern Services, Rizhao Technicians College, No. 2 Linyi North Road, Rizhao, 276800, Shandong Province, PR China.
3Tai'an Tai shan Beer Co., No. 99 Keji North Street, Tai'an, 271000, Shandong Province, PR China.

Abstract. In this paper the arabinoxylanase activities and the content of water-extractable arabinoxylans (WEAXs) including their degradation products in wheat Yannong 24 during the melting process were studied. The results showed that the endo-β-1,4-xylanase activity had a large increase during the steeping process, endo-β-1,4-xylanase and α-L-arabinofuranosidase achieved the highest activity on day 4 of germination. During malt kilning, endo-β-1,4-xylanase and α-L-arabinofuranosidase were not heat stable, and the activity decline rate were 65.88% and 69.58%; whereas malt β-D-xylosidase was resistant to heat and it’s activity decline rate was only 18.74%. It is during steeping and the first day of germination that the content of WEAXs and it’s degradation products (WEAXs+DP) dramatically increased, and this corresponded to the greater improvement of endo-β-1,4-xylanase activity in the same phase. The WEAXs+DP content has a very significant positive correlation with the endo-β-1,4-xylanase activity (0.892, p<0.01). It can be indicated that endo-β-1,4-xylanase is the main degrading enzyme in the degradation of WUAXs and the increase of the WEAXs content during steeping and the first day of wheat germination.

1 Introduction

Wheat and wheat malt are known to be high quality raw materials in the beer brewing process [1], and with the rapid development of the brewing industry in China, wheat malt is now widely used in the beer brewing industry.

Arabinoxylans (AXs) are the primary non-starch polysaccharides found in the cell walls of wheat cells [2-4], with molecular weights ranging widely between 10 and 10,000 kDa. AX forms a relatively viscous solution, with both water-unextractable (WUAX) and water-extractable (WEAX) forms present. The basic structure of AX is that of α-L-arabinofuranosyl residues linked by C(O)-2, C(O)-3 or C(O)-2,3 glycosidic bonds to a linear backbone composed of β-(1,4)-D-xylopyranosyl units, with mono- or di-arabinose substitutions found along the xylose backbone [5].

There are three categories of enzymes that play a primary role in the degradation of AXs, namely, endo-β-1,4-xylanases (EC 3.2.1.8), β-D-xylosidases (EC 3.2.1.55) and α-L-arabinofuranosidase (EC 3.2.1.55) [6-7]. Random enzymatic cleavage along the xylose backbone by endo-β-1,4-xylanase results in the release of xylo-oligosaccharide products, allowing exo-β-D-xylosidase to remove xylose units successively from non-reducing ends of the xylo-oligosaccharides [8]. The existence of side-chain substituents may have an inhibitory effect on the activity of endo-β-1,4-xylanase and β-D-xylosidase; thus, α-L-arabinofuranosidase also plays an essential role in xylan degradation by mediating hydrolysis of the glycosidic bonds between the xylose backbone and side-chain substituents.

The overall AXs content in wheat and barley are similar; however, the WEAXs content is higher in wheat [9], resulting in higher WEXs in malt. The endogenous AXs degrading enzymes found in wheat are active during the mashing and malting process, and can convert WUAXs into WEAXs macromolecules [10-11]. Peng and Jin [12] have also isolated and characterized an endo-1,4-β-xylanase from wheat malt.

A significant number of research reports have been published regarding AX degrading enzymes found in barley [13] and barley malt [14-15], but relatively few studies have been reported for the wheat malting process. Wheat malt is a widely used raw material for beer production; however, a large amount of WEAXs remains during brewing, and is found in the final product [6]. Some WEAXs with large molecular weight that can bring problems, including the wort leaching rate, high viscosity [17], filtration difficulties [18] and even lead to the point of causing beer turbidity and other issues. The present study determined the concentration of the WEAXs, as well as the activity changes of the three primary AXs degrading enzymes. The analysis of the relationship between the enzymatic activities and the characteristics of AXs provides a theoretical foundation for future improvements of the wheat malting process.

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2 Materials and Methods

2.1 Experimental materials and reagents

2.1.1 Experimental materials

The wheat variety Yannong 24, which was obtained from Shandong Academy of Agricultural Sciences (Yantai, Shandong) was used as the raw material and malted according to the method described by Jin [19]. The sampling started at the end of steeping process and carried out once every 24 hours. Considering the effect of drying process on enzyme activity, the sample was divided into two portions. One portion of the green malt was used for enzymatic activity assays of endo-β-1,4-xylanase, β-D-xylosidase and α-L-arabinofuranosidase (labeled as E-0, E-1, ... E-5 according to the number of days after germination) whereas the other portion of the green malt was subjected to a normal drying process in order to obtain the dried malt (labeled as M-0, M-1, ... M-5 according to the number of days after germination). The quality of wheat malt was analyzed according to QB/T 1686-2008 Barley malt. Basic indicators of the wheat malt obtained were shown in Table 1.

2.1.2. Reagents

The p-nitrophenyl-α-L-arabinofuranoside, p-nitrophenyl-β-D-xylopyranoside and RBB-xylan were purchased from SIGMA Company.

2.1.3. Instruments

EBC-LF standard malt mill was purchased from Beijing Dezhijie Co., Ltd (Beijing, China). LRH-250-A biochemical incubator was purchased from Shaoguan Thaihung Medical Instrument Co., Ltd. IKAMAG electromagnetic stirrer was purchased from RET-GS JANKE & KUNKEL GAME & CO.KG. LC-2010AT high-performance liquid chromatography (HPLC) system was purchased from Shimadzu Corporation (City, Japan).

2.2. Enzymatic activity assays

2.2.1. Extraction of crude enzymes

The finished product of malt was ground by using a 0.2-mm sieve. A total of 20 g malt powder was added to 70 mL of 0.05 M phosphate buffer (pH 7.0). After stirring extraction in an ice bath for 30 minutes, the solution was topped up to 100 mL. The solution was centrifuged at 5000 rpm at 4 °C for 15 minutes. The supernatant was harvested and filtered through a 0.45-μm membrane filter to obtain the crude enzyme solution. The crude enzyme solution was then stored in a 4 °C refrigerator.

2.2.2. Endo-β-1,4-xylanase activity assay

The endo-β-1,4-xylanase activity assay was performed according to the method of Guo [20].

2.2.3. α-L-arabinofuranosidase activity assay

A total of 0.5 mL crude enzyme solution was added to 2.5 mL of 0.05 M acetic acid-sodium acetate buffer solution (pH 4.5) and equilibrated in a circulating water bath at 45 °C for 10 minutes, followed by the addition of 1 mL substrate (1 mmol/L p-nitrophenyl-α-L-arabinofuranoside). The blank control was prepared via the addition of 1 mL of 2% Tris-base solution to inactivate the enzymatic activity before the addition of substrate. After 10 minutes of reaction, 1 mL of 2% Tris-base was used to terminate the reaction. Then, the solution was centrifuged at 10000 r/min for 5 minutes and the supernatant was harvested for the measurement of absorbance at 410 nm. The enzyme unit of α-L-arabinofuranosidase was represented as 0.01 change of the absorbance at 410 nm per hour in each g of bone-dry raw material at 45 °C relative to the absorbance value of control.

2.2.4. β-D-xylosidase activity assay

The enzymatic activity assay was performed by using the method described in section 2.2.3 with slight modifications. A total of 0.2 mL crude enzyme solution was added to 2.8 mL of 0.05 M NaAC-HAC buffer solution (pH 4.5) and equilibrated in a circulating water bath at 70 °C for 10 minutes in a circulating water bath, followed by the addition of 1 mL substrate solution (1 mmol/L p-nitrophenyl-β-D-xylopyranoside).

The enzyme unit of β-D-xylosidase was represented as 0.01 change of the absorbance at 410 nm per hour in each g of bone-dry raw material at 70 °C relative to the absorbance value of control.

Table 1 Characteristics of the wheat malts during the malting process

| Wheat malts | M-O | M-1 | M-2 | M-3 | M-4 | M-5 |
|-------------|-----|-----|-----|-----|-----|-----|
| Water content (%) | 5.5±0.0b | 5.8±0.0a | 5.1±0.0c | 5.0±0.0c | 4.9±0.0d | 4.3±0.0e |
| Chromaticity (EBC) | 3.33±0.00f | 4.47±0.00d | 4.09±0.00e | 4.71±0.00c | 5.16±0.00b | 6.17±0.00a |
| α-amino nitrogen (mg/100g) | 51.4±0.6c | 84.1±0.0b | 120.8±0.6a | 124.3±0.3a | 120.2±0.6a | 120.3±0.6a |
| Extract (%) | 72.8±0.0f | 79.6±0.0e | 81.7±0.0c | 82.2±0.0a | 81.9±0.0b | 81.3±0.0d |
2.3. Determination of the content of WEAX and its degradation products

The phloroglucinol-based colorimetry was carried out by using xylose as a standard. A standard curve was constructed by plotting the xylose concentrations (mg) on the x-axis and the absorbance at 552-nm on the y-axis. The data were fitted into the equation \( y = 2.6354x + 0.0134 \) with \( r^2 = 0.9981 \).

The finished product of the malt was ground by using a 0.2-mm sieve. A total of 20 g malt powder was added to 100 mL water and after the stirring extraction in an ice bath for 60 minutes, the solution was centrifuged at 5000 \( r/min \) for 15 minutes. The supernatant was harvested for measurement.

During the measurement, the sample solution was appropriately diluted by adding 1 mL diluent to 5 mL phloroglucinol-glacial acetic acid reaction solution. After incubation in a boiling water bath for 25 minutes, the reaction was terminated in an ice bath, followed by the A552 measurement. The content of WEAX and its degradation products (WEAXs+DP) in the sample was calculated based on the standard curve.

2.4. Data processing

The data were averages of three replicates. Correlation and differential analyses were carried out by using the SPSS 17.0 and DPS 9.50 analysis software. Correlation coefficients with a very significant correlation at the 0.01 level in the correlation analysis were indicated by "***", whereas "**" indicates a significant correlation at the 0.05 level. Different lower case alphabets in the differential analysis indicate a significant difference at the 0.05 level.

Note: Different alphabets in each row indicate a significant difference (\( p<0.05 \)). M-0, M-1, ... M-5 refer to the dried malt at the end of steeping process, the first day, the second day ....... the fifth day of germination, respectively, and the same hereinafter.

3 Results and Discussion

3.1. Arabinoxylanase activities in wheat during the malting process

Activity changes of endo-\( \beta-1,4 \)-xylanase, \( \beta \)-D-xylosidases and \( \alpha \)-L-arabinofuranosidase in the malting process were shown in Figure 1. It could be seen from Figure 1 that the enzymatic activity of endo-\( \beta-1,4 \)-xylanase was low in the raw wheat at 1.27 u. At the end of the steeping process, the enzymatic activity of endo-\( \beta-1,4 \)-xylanase increased significantly up to 7.71 u, but slightly decreased on the first and second day of germination to 6.18 u and 5.74 u, respectively. The enzymatic activity gradually increased on the third and fourth day of germination to 8.21 u and 10.10 u, respectively, where it reached a plateau on the fourth day. The enzymatic activity slightly decreased on the fifth day of germination, but it was not significantly different from that on the fourth day, indicating that the enzymatic activity of endo-\( \beta-1,4 \)-xylanase was stable after 4 days of germination. The reason for the activity declined of endo-\( \beta-1,4 \)-xylanase at the first and second day of germination might be that extraction of enzyme contained some WEAXs derived from wheat malt. And in the process of enzyme activity determination, the WEAXs in the enzyme extraction occupied some endo-\( \beta-1,4 \)-xylanase, so that the enzyme activity became smaller. While in late germination the WEAXs molecules in the wheat malt became smaller, no longer suitable as substrates for endo-\( \beta-1,4 \)-xylanase, no longer take up endo-\( \beta-1,4 \)-xylanase. So that the activity of endo-\( \beta-1,4 \)-xylanase rised.

Changes in \( \beta \)-D-xylosidase and \( \alpha \)-L-arabinofuranosidase activities were basically consistent with one another during the malting process. Their enzymatic activities were the lowest in the raw wheat, slightly increased at the end of the steeping process, and then significantly increased during the germination in a time-dependent manner. The \( \alpha \)-L-arabinofuranosidase activity peaked on the fourth day of germination and did

| Kolbach Index (%) | 17.6±0.1e | 25.4±0.0d | 33.7±0.6c | 35.0±0.2b | 35.8±0.3ab | 36.8±0.2a |
|------------------|-----------|-----------|-----------|-----------|-----------|-----------|
| Viscosity (mPa s) | 2.23±0.00a | 2.13±0.00b | 1.67±0.00c | 1.61±0.00d | 1.52±0.00f | 1.56±0.00e |
| Diastatic power (WK) | 179±0f | 220±1e | 304±4d | 323±0c | 366±0a | 332±1b |
| Total nitrogen (%) | 2.3±0.0a | 2.3±0.0a | 2.3±0.0a | 2.3±0.0a | 2.3±0.0a | 2.4±0.0a |

Fig. 1 Enzyme activity change of xylanase in the process of malting. E-1, ... E-5 refer to the green malt of the first day, the second day ....... the fifth day of germination, respectively, and the same hereinafter. Y-axis on the right is for the activity of \( \beta \)-D-xylosidase. Y-axis on the left is for the activity of endo-\( \beta-1,4 \)-xylanas and \( \alpha \)-L-arabinofuranosidas
not change significantly on the fifth day relative to the fourth day. The β-D-xylosidase activity peaked on the fifth day of germination. However, the β-D-xylosidase activity was greatly higher than the α-L-arabinofuranosidase activity (Table 2). The β-D-xylosidase activities in the raw wheat, green malt, and final product of the malt (7.36 u, 170.31 u and 138.39 u, respectively) were 2, 4, and 11 times greater than the α-L-arabinofuranosidase activities (3.50 u, 40.07 u and 12.19 u, respectively).

As can be seen from Table 2, the enzymatic activities of the xylanases in the malt decreased before and after the drying process, whereas the activities of endo-β-1,4-xylanase, α-L-arabinofuranosidase and β-D-xylosidase decreased by 65.88%, 69.58%, and 18.74%, respectively. The results showed that endo-β-1,4-xylanase and α-L-arabinofuranosidase in the wheat malt were not heat-resistant, whereas β-D-xylosidase had a higher heat resistance.

| Table 2 | Comparison of the xylanase activity in wheat, green wheat malt and dry wheat malt |
|-----------------------------------------------|
| Enzyme                                      | Enzyme activity (u) | Decline rate in the drying process (%) |
|-----------------------------------------------|---------------------|-----------------------------------------|
| endo-β-1,4-xylanase                          | 1.27                | 9.70                                    |
|                                             | 3.31                | 65.88                                   |
| α-L-arabinofuranosidase                      | 3.50                | 40.07                                   |
|                                             | 12.19               | 69.58                                   |
| β-D-xylosidase                               | 7.36                | 170.31                                  |
|                                             | 138.39              | 18.74                                   |

Note: The results suggested the best germination time was between four to five days regarding the enzymatic activities of xylanases in the wheat malt.

3.2. Changes in the content WEAXs+DP during the malting process

Changes in content of WEAXs+DP during wheat malting were shown in Table 3. Table 3 showed that the WEAXs+DP content was 5.25 mg/g in the raw wheat, 6.77 mg/g at the end of steeping process, 12.09 mg/g on the first day of germination, and 12.94 mg/g on the second day of germination, then 13.23 mg/g, 13.72 mg/g and 13.80 mg/g on the third, fourth and fifth day. The increment of the WEAXs content was due to the degradation of arabinoxylan by the endogenous xylanase that leads to the conversion of WUAXs into WEAXs. In the malting process of wheat malt, the WEAXs+DP content mainly increased during the steeping process and the first day of germination. indicating that the conversion of WUAXs into WEAXs in wheat was more intense during the steeping process and the first day of germination. The WEAXs+DP content in the malt changed slightly after the second day of germination, indicating that the conversion of WUAXs into WEAXs in wheat was slower. But may be the conversion that WEAXs from large molecules to small molecules were more intense, this speculation needed to be confirmed by further study.

| Table 3 | Changes of the content of water-extractable arabinoxylan during the malting process |
|-----------------------------------------------|
| Germination | WEAXs+DP content (mg/g) | Increasing rate of WEAXs+DP content (%) |
|-----------------------------------------------|--------------------------|-----------------------------------------|
| wheat                            | 5.25±0.05f               | 0.00                                    |
| M-0                              | 6.77±0.10e               | 28.88                                   |
| M-1                              | 12.09±0.09d              | 130.29                                  |
| M-2                              | 12.94±0.18c              | 146.17                                  |
| M-3                              | 13.23±0.08b              | 151.94                                  |
| M-4                              | 13.72±0.07a              | 161.33                                  |
| M-5                              | 13.80±0.15a              | 162.80                                  |

Note: Increasing rate of WEAXs+DP content (%)=(WEAXs+DP content in malt - WEAXs+DP content in wheat)/ WEAXs+DP content in wheat×100

3.3. Correlation between the arabinoxylanase activities and the WEAXs+DP content

The correlation between the activities of the three arabinoxylanases and the WEAXs+DP content was presented in Table 4. The WEAXs+DP content has a very significant positive correlation with the endo-β-1,4-xylanase activity (0.892, p<0.01). However, it does not have a significant correlation with the activities of α-L-arabinofuranosidase and β-D-xylosidase, indicating that endo-β-1,4-xylanase is the main degrading enzyme in the degradation of AXs and the increase of the WEAXs content in the malting process. There was a significant positive correlation between the activity of endo-β-1,4-xylanase and β-D-xylosidase (0.765, p<0.05) and a very significant positive correlation between the activity of α-L-arabinofuranosidase and β-D-xylosidase (0.976, p<0.01), indicating that there is a certain synergy between these two types of enzymes.

| Table 4 | Correlation between the activities of the arabinoxylan degrading enzymes and the content of WEAXs |
|-----------------------------------------------|
| WEA Xs | endo-β-1,4-xylanase | α-L-arabinofuranosidase | β-D-xylosidase |
|-----------------------------------------------|------------------------|------------------------|----------------|
| WEAXs | 1                      | 0.892                  | 1                      |
| endo-β-1,4-xylanase | 0.892                  | 1                      | 0.892                  |
| α-L-arabinofuranosidase | 0.711                  | 0.747                  | 0.976                  |
| β-D-xylosidase | 0.650                  | 0.765                  | 1                      |

Note: Increasing rate of WEAXs+DP content (%)=(WEAXs+DP content in malt - WEAXs+DP content in wheat)/ WEAXs+DP content in wheat×100

As can be seen from Table 2, the enzymatic activities of the xylanases in the malt decreased before and after the drying process, whereas the activities of endo-β-1,4-xylanase, α-L-arabinofuranosidase and β-D-xylosidase decreased by 65.88%, 69.58%, and 18.74%, respectively.
4 Conclusion

The enzymatic activity of endo-β-1,4-xylanases increased at the highest magnitude during the steeping process and increased at a lower magnitude during the germination, where its enzymatic activity peaked on the fourth day of germination. The best germination time is between four to five days regarding the enzymatic activities of xylanases in the wheat malt. During the drying process, both endo-β-1,4-xylanase and α-L-arabinofuranosidase in the wheat malt were not heat-resistant whereas β-D-xylosidase was heat resistance. In the malting process, the conversion of WUAXs into WEAXs in the wheat by the endogenous endo-β-1,4-xylanase was more intense during the steeping process and the first day of germination. Endo-β-1,4-xylanase was the main degrading enzyme in the degradation of AXs and the increase of the WEAXs content in the malting process.

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