Impact of exogenous maltogenic α-amylase and maltotetraogenic amylase on sugar release in wheat bread

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Abstract

The use of exogenous maltogenic α-amylases or maltotetraogenic amylases of bacterial origin is common in wheat bread production, mainly as antistaling agents to retard crumb firming. To study the impact of maltogenic α-amylase and maltotetraogenic amylase on straight dough wheat bread, we performed a discovery-driven proteomics approach with commercial enzyme preparations and identified the maltotetraogenic amylase P22963 from Pelomonas saccharophila and the maltogenic α-amylase P19531 from Geobacillus stearothermophilus, respectively, as being responsible for the amylolytic activity. Quantitation of mono-, di- and oligosaccharides and residual amylase activity in bread crumb during storage for up to 96 h clarified the different effects of residual amylase activity on the sugar composition. Compared to the control, the application of maltogenic α-amylase led to an increased content of maltose and especially higher maltooligosaccharides during storage. Residual amylase activity was detectable in the breads containing maltogenic α-amylase, whereas maltotetraogenic amylase only had a very low residual activity. Despite the residual amylase activities and changes in sugar composition detected in bread crumb, our results do not allow a definite evaluation of a potential technological function in the final product. Rather, our study contributes to a fundamental understanding of the relation between the specific amylases applied, their residual activity and the resulting changes in the saccharide composition of wheat bread during storage.

Keywords Bread · Discovery-driven proteomics · Enzyme activity · Maltogenic α-amylase · Maltotetraogenic amylase · Wheat

Introduction

Wheat bread can be considered as an unstable multiphase food matrix subject to changes during storage: the flavor of fresh bread and the crispiness of the bread crust decrease, whereas crumb firmness and crumbliness increase. The changes in the bread crumb, also known as staling, lead to a loss of resilience and to firming [1], caused by water immobilization and redistribution processes. The retrogradation of amylopectin during storage immobilizes water, which leads to a decrease in freezable water in the bread crumb [2]. Furthermore, water redistribution occurs via water migration from gluten to starch and from crumb to crust. These migration patterns lead to an additional decrease in freezable water and to an increase in crumb firmness [3, 4]. Wheat starch rearrangements play the leading role in crumb firming. Amylose gelation occurs within hours, whereas that of amylopectin takes several days and involves reorganization of amylopectin side chains to form tightly
packed double-helical structures. During storage, the amorphous gelatinized starch network present in fresh bread is transformed first into a partly crystalline state and then to a mostly continuous rigid crystalline starch network [3, 5, 6].

Amylases can be added during bread production as antistaling agents [7]. Maltotetraogenic amylases and maltogenic α-amylases are most commonly described as efficient antistaling agents in bread making [6, 8]. Maltotetraogenic amylase from *Pelomonas saccharophila* hydrolyzes α-(1 → 4)-d-glucosidic bonds to remove successive maltotetraose residues from the non-reducing ends. It mainly acts on amylopectin, thus reducing amylopectin side chain rearrangement and crystallization [9, 10]. Maltogenic α-amylase from *Geobacillus stearothermophilus* successively cleaves α-maltose units from the non-reducing ends of amylose and amylopectin. In comparison to maltotetraogenic amylase, it has a smaller effect on the molecular weight of the starch polymers and degrades the amylopectin molecules relatively slowly [8, 10]. However, the enzyme reduces the molecular weight of amylose and thus, enhances amylase mobility and network formation [5]. This results in a greater initial firmness of breads manufactured with maltogenic α-amylase [11, 12]. The main influence on bread firming caused by maltogenic α-amylase is due to its subsequent cleavage along the side chains of the amylopectin molecules. They become too short to crystallize and the formation of a permanent amylopectin network during storage is inhibited [8, 10]. Despite the initial increase in crumb firmness, this leads to a substantial delay in crumb firming [12].

An additional antistaling effect of maltotetraogenic and maltogenic α-amylases may result from their ability to increase dextrin levels in wheat bread. Dextrins hinder the crystallization of amylopectin helices and may increase relative crumb humidity. However, compared to the hydrolytic effect on the starch polymers, the antistaling effect of dextrins is described to be of minor importance [13]. Maltotetraogenic amylase from *P. saccharophila* and maltogenic α-amylase from *G. stearothermophilus* have a temperature optimum at approximately 60 °C [14]. Several studies showed that both enzymes have a high temperature stability, whereby it is accepted that *G. stearothermophilus* maltogenic α-amylase has a higher temperature stability and can be still active in wheat bread crumb after baking [8, 15, 16]. It is unknown whether this activity is associated with an additional antistaling effect during storage of bread. In the following, residual activity means the amylolytic activity measured using enzymatic assays after extraction from baked bread crumb. To gain a better understanding of the effects of maltogenic α-amylase and maltotetraogenic amylase in wheat bread making, it is necessary to study the formulation of the specific preparations as well as the impact of the preparations on sugar release during bread storage. Thus, a combined analytical approach has to be implemented to focus on the different relevant parameters in amylase preparations and wheat bread.

The use of enzymes in food production is defined in the European Union by the Regulation (EC) no. 1332/2008 and has to be in line with the Regulation (EU) no. 1169/2011 on the provision of food information to consumers [17, 18]. According to these regulations, enzymes used as processing aids do not have to be labeled, because they are not classified as food ingredients of technological relevance in the final product. However, only in case this potential residual activity would have a functional effect in the final product, it would have to be labeled [17, 18].

The aim of our work was to characterize the protein composition of commercial amylase preparations commonly used in bread making and to elucidate their impact on the formation of sugars in the bread crumb during storage. In the first step, we identified and quantitated the amylase compared to other proteins in the preparation. Then we monitored sugar concentrations in wheat bread crumb during storage for up to 96 h and combined this with the quantitation of residual amylase activity. This approach allowed us to detect changes caused by a residual amylolytic activity in straight dough wheat bread.

**Materials and methods**

**Chemicals and flours**

All chemicals and solvents were HPLC-grade or higher. Water was deionized and purified using an Arium 611VF water purification system (Sartorius, Goettingen, Germany). Trypsin (from bovine pancreas, TPCK-treated, ≥ 10,000 BAEE U/mg protein), sodium hydroxide solution (50–52%; eluent for IC) and anhydrous sodium acetate (≥ 99%) were from Sigma-Aldrich (Steinheim, Germany). α-Amylase assay reagent (4-nitrophenyl α-d-maltotrioside, plus excess α-glucosidase and glucoamylase) and β-amylase assay reagent (4-nitrophenyl β-d-maltotriose, plus excess thermostable β-glucosidase) were purchased from Megazyme (Bray, Ireland). d-(+)-Glucose (≥ 99.5%), d-(−)-fructose (≥ 99%), sucrose (≥ 99.5%), maltose monohydrate (≥ 99%), maltotriose (≥ 90%) and maltotetraose (≥ 70%) were from Sigma-Aldrich (Steinheim, Germany).

All wheats and flours were HPLC-grade or higher. Water was deionized and purified using an Arium 611VF water purification system (Sartorius, Goettingen, Germany). Trypsin (from bovine pancreas, TPCK-treated, ≥ 10,000 BAEE U/mg protein), sodium hydroxide solution (50–52%; eluent for IC) and anhydrous sodium acetate (≥ 99%) were from Sigma-Aldrich (Steinheim, Germany). α-Amylase assay reagent (4-nitrophenyl α-d-maltotrioside, plus excess α-glucosidase and glucoamylase) and β-amylase assay reagent (4-nitrophenyl β-d-maltotriose, plus excess thermostable β-glucosidase) were purchased from Megazyme (Bray, Ireland). d-(+)-Glucose (≥ 99.5%), d-(−)-fructose (≥ 99%), sucrose (≥ 99.5%), maltose monohydrate (≥ 99%), maltotriose (≥ 90%) and maltotetraose (≥ 70%) were from Sigma-Aldrich (Steinheim, Germany). According to these regulations, enzymes used as processing aids do not have to be labeled, because they are not classified as food ingredients of technological relevance in the final product. However, only in case this potential residual activity would have a functional effect in the final product, it would have to be labeled [17, 18].

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Amylase preparations

Seven different commercial amylase preparations commonly applied in wheat bread making were used (Table 1). Two of them, (I) and (II), contained maltotetraogenic amylase (EC 3.2.1.60) from *P. saccharophila*. The maltogenic α-amylase (EC 3.2.1.133) in the preparations (III), (IV), (V), (VI), and (VII) was from *G. stearothermophilus*.

Determination of maltotetraogenic amylase activity

The extraction of amylases for the activity assays was performed according to Cornaggia et al. [19]. In total, 1.0 g of dry matter of amylase preparation or bread was extracted with 10 mL of extraction buffer (0.1 mol/L sodium maleate, pH 5.5, 1.0 mg/mL bovine serum albumin, 0.025 mol/L dithiothreitol (DTT)) for 16 h at 22 °C. Then, centrifugation (2500×g; 20 °C; 10 min) and filtration (0.45 µm; cellulose acetate filters) were performed.

The activity of maltotetraogenic amylase was determined using the Ceralpha method (Megazyme). The assay procedure was carried out as described by the manufacturer, except that the incubation time was extended to 60 min at 40 °C for the assay reaction. Quantitation of 4-nitrophenol was performed by means of external calibration with a standard solution containing 4-nitrophenol dissolved and diluted with 1.5 mL of 20% (w/v) tri-sodium phosphate buffer (pH 11). Maltotetraogenic amylase activity was reported in nkat/g dm. One kat is defined as the amount of 4-nitrophenol in mol released from the substrate non reducing-end blocked 4-nitrophenylmaltoheptaoside in one second under the assay conditions. All determinations were performed with three technical replicates.

Determination of maltogenic α-amylase activity

The extraction of maltogenic α-amylase was carried out as described above according to Cornaggia et al. [19]. Then the betamyl-3 method (Megazyme) was used to determine maltogenic α-amylase activity. The amylase extract was diluted in an appropriate ratio with 0.1 mol/L 2-(N-morpholino)ethanesulfonic acid buffer (pH 6.2) containing 0.01 mol/L ethylenediaminetetraacetic acid and 1.0 mg/mL bovine serum albumin. For the assay procedure, the incubation time of the extract and the substrate was set to 20 min at 40 °C. The following procedure was performed exactly as described by the kit manufacturer. Quantitation of 4-nitrophenol was performed by means of external calibration with a standard solution containing 4-nitrophenol dissolved and diluted in 1% (w/v) tris(hydroxymethyl)aminomethane (Tris)-buffer (pH 8.5). Amylase activity was expressed in nkat/g dm. One kat is defined as the amount of 4-nitrophenol in mol released from the substrate non reducing-end blocked 4-nitrophenyl-β-α-maltotrioside in 1 s under the assay conditions. All determinations were performed with three technical replicates.

Extraction of enzymes from the amylase preparations and determination of the protein content

For high performance liquid chromatography time-of-flight mass spectrometry (UHPLC–TOF–MS) analysis, the amylases were extracted from the preparations as described by Uhr et al. [20]. Each sample (0.5 g of dry matter) was extracted using 5.0 mL of extraction buffer (0.1 mol/L ammonium bicarbonate, 4.0 mol/L urea and 0.005 mol/L DTT, pH 8.0, in water). The extraction was carried out at activity in the bread crumb and relative residual activity based on the activity added to the dough.

Table 1 Composition of the bacterial amylase preparations and dosage recommendation as indicated by the manufacturer, activity of the amylase added to the dough in the baking experiments, residual activity in the bread crumb and relative residual activity based on the activity added to the dough.

| Preparation | Enzyme                          | EC number | Other ingredients                        | Dosage (mg/kg of flour) | Activity (nkat/g dm) | Residual activity (nkat/g dm) Mean (%) | Range (%) |
|-------------|--------------------------------|-----------|------------------------------------------|-------------------------|---------------------|----------------------------------------|----------|
| (I)         | Maltotetraogenic amylase       | 3.2.1.60  | Sodium chloride, wheat starch            | 850                     | 85.8                | 0.9±0.3                               | 1.1      |
| (II)        | Maltotetraogenic amylase       | 3.2.1.60  | Sodium chloride, wheat starch, microcrystalline cellulose | 300                     | 31.7                | 0.6±0.2                               | 1.9      |
| (III)       | Maltogenic α-amylase           | 3.2.1.133 | Wheat flour, sodium chloride             | 100                     | 22.0                | 9.3±1.4                               | 42.3     |
| (IV)        | Maltogenic α-amylase           | 3.2.1.133 | Maltodextrin, sodium chloride            | 100                     | 43.0                | 11.0±5.3                              | 25.7     |
| (V)         | Maltogenic α-amylase           | 3.2.1.133 | Maltodextrin, sodium chloride, sunflower oil | 100                     | 38.6                | 11.0±5.0                              | 28.6     |
| (VI)        | Maltogenic α-amylase           | 3.2.1.133 | Wheat flour, sodium chloride             | 100                     | 31.5                | 14.3±3.8                              | 45.5     |
| (VII)       | Maltogenic α-amylase           | 3.2.1.133 | Maltodextrin, sodium chloride, sunflower oil | 50                      | 41.0                | 5.7±1.2                               | 13.9     |

dm, based on dry matter, values are given as means (n=3)± standard deviation
room temperature by stirring for 30 min. The extract was centrifuged (20 min; 20 °C; 3750xg) and the supernatant was collected for further analysis. To optimize the conditions for the tryptic digest of extracts, the protein contents were quantified via a RP-HPLC method with UV detection at 210 nm as described earlier [21].

**Sample preparation, tryptic digest, and peptide clean-up**

The supernatant of the enzyme extracts was diluted in an appropriate volume to a protein concentration of 0.5 mg/mL in 0.3 mL and 0.1 mL Tris–HCl (1.5 mol/L) and 0.1 mL 1-propanol were added. Reduction was performed by adding tris(2-carboxyethyl)phosphine solution (0.05 mol/L tris(2-carboxyethyl)phosphine in 0.5 mol/L Tris–HCl, pH 8.5) and incubation for 30 min at 60 °C. Alkylation of cysteine residues was performed for 45 min at 37 °C in the dark using 0.1 mL chloroacetamide solution (0.5 mol/L chloroacetamide in 0.5 mol/L Tris–HCl, pH 8.5). Then the samples were dried using a rotary vacuum concentrator (37 °C, 3 h, 800 Pa). For protein hydrolysis, the samples were reconstituted in 0.5 mL of trypsin stock solution with a trypsin-to-substrate ratio of 1:20 (w/w) (0.04 mol/L urea in 0.1 mol/L Tris–HCl, pH 7.8) and incubated at 37 °C for 16 h. The hydrolysis was stopped by heating the samples at 95 °C for 8 min. The peptide mixture was cleaned by solid phase extraction (SPE) using 100 mg Discovery DSC-18 tubes (Merck, Darmstadt, Germany). The tubes were activated using 0.1 mL chloroacetamide solution (0.5 mol/L chloroacetamide in 0.5 mol/L Tris–HCl, pH 8.5). Then the samples were reconstituted in 500 µL FA and filtered (0.45 µm; wwPTFE).

**Discovery driven LC–MS/MS analysis**

A Sciex TripleTOF 6600 mass spectrometer (Sciex, Darmstadt, Germany) connected to a Shimadzu Nexera X2 system (Shimadzu, Kyoto, Japan) operating in the positive electrospray ionization (ESI) mode was used for UHPLC–TOF–MS analysis. Control of the instrument and data acquisition were performed using AnalystTF software (version 1.7.1, Sciex). The chromatography system consisted of a 150 × 2.1 mm, 1.6 µm BioZen Peptide PS-C-18 LC column (Phenomenex, Aschaffenburg, Germany) equipped with a security guard ultra-column (2 × 2.1 mm) of the same type. The injection volume was 10 μL and the temperature of the column oven was set to 40 °C. The elution solvents were (A) 1% (v/v) aqueous FA and (B) acetonitrile containing 1% (v/v) FA. A 75 min gradient was used to separate the peptides with a flow rate of 0.35 mL/min starting at 0 min, 5% B; 5 min, 5% B; 60 min, 40% B; 65 min, 100% B; 69 min, 100% B; 70 min, 5% B; and 75 min, 5% B. The following parameters were set: ion spray voltage: 5500 eV, source temperature: 550 °C, nebulizing gas: 55 psi, heating gas: 65 psi and nitrogen as curtain gas: 35 psi.

A TOF–MS survey scan was acquired in the information-dependent acquisition mode (IDA), from mlz 400 to 1000 using 250 ms as accumulation time, 10 V as collision energy (CE) and 80 V as declustering potential (DP). Product ion spectra were recorded in the high-resolution mode for 70 ms for the 12 most abundant compounds. The mlz range was mlz 100–2000 using a DP of 80 V and CE of 40 V, with CE spread of 15 V. IDA criteria were set for the precursor ion intensity at > 100 counts/s. Q1 resolution was set to 0.7 Da.

**Identification of amylases using MaxQuant**

TOF raw files were used for the identification and relative quantitation of proteins using MaxQuant (version 1.6.10.43) [22]. A workflow with different searches against databases derived from UniProtKB using the search engine Andromeda was developed [23]. The first step of data evaluation comprised the identification of enzymes from the class EC 3.2.-.- (glycosylases). A database with all proteins belonging to the class EC 3.2.-.- from *Triticum aestivum* and from 14 microorganisms relevant for the production of food grade amylases according to EFSA and FDA was built [24, 25]. This database contained 5550 proteins (downloaded from UniProtKB on July 01, 2019). In the second step, all proteins present in the sample were identified, using databases containing the proteome of the identified organisms, respectively. In the final step, the relative quantitation of the proteins was performed using the intensity based absolute quantitation (iBAQ) algorithm implemented within MaxQuant [26]. For each sample, an individual database was built containing all proteins identified in the second step. A total sum normalization of iBAQ protein intensities between samples was performed to correct for the different total protein injection amounts.

The MaxQuant parameters were set as follows for all searches: Variable modifications: oxidation of methionine, N-terminal protein acetylation; fixed modification: carboxymethylation on cysteine; proteolytic enzyme: trypsin; missed cleavage sites: up to two. Match-between-runs was enabled with a matching time window of 0.7 min and an alignment time window of 20 min. Results were filtered for a minimal length of five, a maximal length of 50 amino acids and 1% peptide and protein false discovery rate.

The content of amylase [mg/g] present in the preparations added in the baking experiments, respectively, was calculated based on the protein contents of the amylase extracts.
for UHPLC–TOF–MS analysis considering the percentage of amylase determined via iBAQ. Thus, the absolute protein content in the sample was multiplied with the relative iBAQ value for each amylase.

**Baking experiments**

The recipe for the baking experiments was: Wheat flour type 550 (30 g, based on 14% moisture), 2.0 g baking soda, 0.25 g sodium chloride and bulk liquid as determined in the Farinograph-E [27]. Water as bulk liquid was replaced by a buffer solution with 0.1 mol/L citric acid monohydrate and 0.1 mol/L trisodium citrate dihydrate (pH 2.75) to obtain a pH comparable to yeast-leavened wheat bread (pH 6). Baking soda was added 1 min before the end of the kneading time. The dough was kneaded at 60 rpm to optimum consistency in the Farinograph-E [27]. After kneading, the dough was rounded for eight cycles and placed in the proofing cabinet for 20 min at 30 °C in a water-saturated atmosphere. Then the dough piece was rolled (roll gap: 0.8 cm), folded in half, rounded for five cycles, and put into an aluminium baking pan for the second dough rest for 40 min under the same conditions as before. Baking took place in a deck oven Type-9110-0082 (Binder, Tuttlingen, Germany) at 230 °C for 18 min.

The breads were either analysed after cooling (2 h) or they were stored at 22 °C under vacuum until further analysis. Amylase activity was determined after a storage time of 30 h. To identify and quantitate sugars in the samples they were either extracted directly after cooling (2 h) or after storage for 22 h, 48 h and 96 h at 22 °C. The baking experiments were carried out with four replicates.

**Quantitation of mono-, di-, and oligosaccharides in wheat bread crumb**

Mono-, di- and oligosaccharides were extracted from a sample taken from the centre of the bread crumb (0.5 g of dry matter) using 4.0 mL of 50% (v/v) methanol while stirring for 30 min at 22 °C followed by centrifugation at 3500×g and 20 °C for 20 min. The solvent was removed using a rotary vacuum concentrator (37 °C, 9 h, 800 Pa). The dry samples were reconstituted in 1.0 mL deionized water and cleanup was performed using strong cation exchange columns (StrataX-C 33 µm, 200 mg, 3 mL, Phenomenex). The columns were activated with methanol (1 × 3.0 mL) and washed with 0.1% (v/v) FA (3 × 3.0 mL). After loading the sample, the sugars were eluted with 0.1% (v/v) FA (2 × 2.0 mL). The eluent was removed using a rotary vacuum concentrator (37 °C, 9 h, 800 Pa) and the sugars were redissolved in deionized water for further analysis.

Mono-, di- and oligosaccharides were analysed by high-performance anion exchange chromatography with pulsed amperometric detection (HPAEC-PAD) on a Dionex ICS-5000 system (Thermo Fisher Scientific). The HPAEC-PAD system was equipped with a CarboPac PA 100 column (250 × 4 mm) connected to a CarboPac PA 100 guard column (50 × 4 mm). Elution was performed with 1.0 mol/L sodium acetate solution (solvent A), 1.0 mol/L sodium hydroxide solution (solvent B) and deionized water (solvent C) at a flow rate of 1.0 mL/min. Solvent B was set to 15% for the total run time. The gradient was 0 min, 0.5% A; 2 min, 0.5% A; 8 min, 10% A; 18 min, 20% A; 23 min, 20% A; 24 min, 0.5% A; 39 min, 0.5% A. PAD detection was carried out with a gold working electrode operating with a standard quadrupole waveform. Chromeleon software 7.2 (Thermo Fisher Scientific) was used for system control and data analysis. Quantitation of the sugars was performed by means of external calibration with standard solutions.

**Statistical analysis**

The sugar contents of the breads containing one amylase preparation each are presented as mean value ± standard deviation. Linear mixed models were used to explore possible differences of the sugar contents between the time points (2 h, 22 h, 48 h and 96 h after baking). The Bonferroni correction was applied to account for multiple testing. Statistical analyses were performed with R statistical software, version 3.6.1. (R Core Team, Vienna, Austria).

**Results**

**Identification of amylases and the protein composition of the preparations**

One maltotetraogenic amylase from *P. saccharophila* (UniProtKB accession P22963) was identified in preparations (I) and (II), respectively. In addition, an α-amylase (D9J2M5) was identified in preparation (II). All maltogenic α-amylase preparations (III)–(VII) contained P19531, a maltogenic α-amylase from *G. stearothermophilus* and no additional amylolytic enzyme was detected (Table 2).

The composition of the preparations is shown in Figs. 1 and 2. Comparing the relative iBAQ values for preparation (I), the amylase P22963 accounted for the largest share (87.8%) of all proteins, followed by 7.5% of proteins from *Bacillus licheniformis* and 2.1% of other proteins from *P. saccharophila*. In preparation (II), the major proportion of 71.1% was P22963 followed by other *P. saccharophila* proteins with 12.4%. The α-amylase D9J2M5 from *Aspergillus oryzae* accounted for 1.6% whereas other *Aspergillus oryzae* proteins were present with 4.4%. Preparations (I) and (II) additionally contained other wheat proteins [2.6% for (I) and 10.5% for (II)].
All maltogenic α-amylase preparations (Fig. 2) contained P19531 with a share of ≥ 90% [(IV) and (VII)), about 50% ((V) and (VI)] or comparably low with 5.4% (III). Preparations (IV) and (VII) contained exclusively proteins from *G. stearothermophilus*. Only preparations (III) and (VI) contained wheat proteins besides P19531 and other *G. stearothermophilus* proteins. Preparation (III) contained 53.4% gluten and 41.2% other wheat proteins and preparation (VI) 22.8% gluten proteins and 30.8% other wheat proteins. Preparation (V) contained 34.9% proteins from *Bacillus atrophaeus*.

Based on the iBAQ values, the quantity of P22963 from *P. saccharophila* added in the baking experiments was 0.32 mg for preparation (I) and 0.05 mg for (II). Accordingly, the absolute amount of P19531 from the maltogenic α-amylase preparations in the baking recipe ranged from 0.01 mg (III) to 1.21 mg (VII).

### Composition of mono-, di-, and oligosaccharides in bread without amylase addition

The composition of mono-, di-, and oligosaccharides in bread crumb without addition of exogenous amylases (control) was already reported by Rebholz et al. [21]. The contents were as follows, sorted in descending order: maltose: 14.6 mg/g dm (± 1.1), sucrose: 5.8 mg/g dm (± 0.8), glucose: 2.9 mg/g dm (± 0.6), fructose: 2.8 mg/g dm (± 2.0), maltotriose: 1.5 mg/g dm (± 0.1) and maltotetraose: 1.1 mg/g dm (± 0.3). No significant changes in the content of the detected sugars were detected during 48 h of storage.

### Composition of mono-, di-, and oligosaccharides in breads with amylase addition and changes during storage

The total sugar content was 39.1 mg/g dm (± 3.8) for preparation (I) (Fig. 3) and 58.9 mg/g dm (± 5.4) for preparation (II) 2 h after baking (Fig. 4). The glucose content did not increase significantly from 2 to 22 h, but increased significantly from 22 to 48 h [3.8 mg/g dm (± 0.3) to 4.8 mg/g dm (± 0.4)] in the bread containing preparation (I). The content of maltotriose increased significantly from 22 h [0.2 mg/g dm (± 0.2)] to 48 h [1.6 mg/g dm (± 0.5)], while the content of fructose, maltose and maltotetraose did not show significant changes from 2 to 48 h of storage. Glucose, fructose, sucrose, maltose, maltotriose and maltotetraose were detected 2 h after baking in the bread containing preparation (II). In contrast to preparation (I), maltopentaose, maltohexaose, maltoheptaose and maltooctaose were additionally detected in the bread crumb after 48 h. The maltotetraose content increased significantly from 2.1 mg/g dm (± 1.7) to 29.5 mg/g dm (± 2.0), so that it was the main sugar in the bread manufactured with (II) 96 h after baking.
Glucose, fructose, sucrose, maltose, maltotriose and maltotetraose were identified 2 h after baking in all breads manufactured with exogenous maltogenic α-amylase (Figs. 5, 6), except for breads with (IV) where maltotetraose was identified for the first time at 22 h after baking. Additionally, maltopentaose was detected after 48 h and maltohexaose after 96 h of storage in the samples containing preparation (VI). The total sugar content ranged from 55.3 mg/g$_{dm}$ (± 4.4) (V) to 108.4 mg/g$_{dm}$ (± 1.6) (VI) after 2 h. Maltose contents ranged from 25.3 mg/g$_{dm}$ (± 1.9) for (V) to 50.0 mg/g$_{dm}$ (± 4.4) for (VII) 2 h after baking. No significant increase in maltose content was detected between 2 and 96 h in the breads manufactured with (III) and (VII). For preparation (IV), the maltose content increased significantly
from 27.7 mg/g dm (± 0.8) up to 38.8 mg/g dm (± 8.6), for (V) from 25.3 mg/g dm (± 1.9) to 40.4 mg/g dm (± 0.7) and for (VI) from 37.0 mg/g dm (± 3.2) to 68.6 mg/g dm (± 4.9) considering 2 h and 96 h of storage. The maltotetraose content of the bread crumbs containing preparations (V), (VI) and (VII) increased significantly between 2 and 96 h of storage. The increase was from 5.9 mg/g dm (± 3.9) to 11.9 mg/g dm (± 2.2) for (V), from 9.7 mg/g dm (± 1.7) to 44.1 mg/g dm (± 6.6) for (VI) and from 0.6 mg/g dm (± 0.2) to 22.9 mg/g dm (± 9.3) for (VII). No significant changes in the content of sucrose were observed during storage of the breads containing either of the maltogenic α-amylase preparations.

Residual activity of amylases in model wheat bread

The endogenous activity of the wheat flour used for the baking experiments was 4.1 nkat/g dm (± 0.7) for α-amylase and 378.4 nkat/g dm (± 6.1) for β-amylase. As we have shown before, no cereal amylase activity was detected in the crumb of the control breads after baking [21].

For the maltotetraogenic amylase preparation, (I) we detected 85.8 nkat/g dm (± 11.6) and for preparation (II) 31.7 nkat/g dm (± 6.6). After baking, the residual amylase activity was 0.9 nkat/g dm (± 0.3) in (I) and 0.6 nkat/g dm (± 0.2) in (II). This corresponds to a relative residual activity of 1.1% for (I) and 1.9% for (II) compared to the amylase activity originally used in the dough (Table 1). The activity of maltogenic α-amylase preparations applied to the dough was between 22.0 nkat/g dm (± 0.7) for preparation (III) and 42.9 nkat/g dm (± 2.0). All tested maltogenic α-amylase preparations showed a residual amylase activity (Table 1). Based on the absolute values, the residual activity was between 5.7 nkat/g dm (± 1.2) for (VII) and 14.3 nkat/g dm (± 3.8) for (VI). Compared to the activities applied to the doughs, the relative residual amylase activity in the breads ranged from 13.9% in the breads with (VII) to 45.5% for (VI).

Discussion

In this study, we combined discovery-driven proteomics with the quantitation of mono-, di- and oligosaccharides in bread crumb and the determination of residual amylase activity to get insights into changes in sugar composition caused by the action of exogenous maltotetraogenic amylases and maltogenic α-amylases during straight dough wheat bread making and storage.

For the group of maltotetraogenic amylase and the group of maltogenic α-amylase preparations, one amylase each was identified as being responsible for the amylolytic activity. Our results are in accordance with Prandi et al. who also identified P19531, a maltogenic α-amylase from G. stearothermophilus, as being responsible for the amylolytic activity in maltogenic α-amylase preparations applied in baking [28]. To the best of our knowledge, we are the first to analyze maltotetraogenic amylase preparations with proteomics tools and we identified a single amylase P22963 as cause of the amylolytic activity. The iBAQ evaluation was
shown to be suitable for the relative quantitation of several different proteins in one sample [26, 29]. The proportion of maltotetraogenic amylase in the preparations (I) and (II) was 87.8% and 71.0%, respectively. Both preparations also contained 4–7% of proteins from other microorganisms and non-gluten proteins, such as the starch synthases P27736 and Q8W2G8, indicating that wheat starch was used as a filler material [30]. The maltogenic α-amylase preparations showed large differences in composition considering that the percentage of maltogenic α-amylase ranged from 5.4% (III) up to 96.9% (VII).

Although both groups of preparations, maltotetraogenic amylase and maltogenic α-amylase, contained the same active enzyme, respectively, the effects on the sugar composition of the bread crumb were quite different. A higher initial activity applied to the dough resulted in a higher maltotetrose content in the bread crumb using the maltotetraogenic amylase preparations (I) and (II). The activity applied to the dough of (I) was 85.8 nkat/g dm and 171% higher than the activity of (II) with 31.7 nkat/g dm. Corresponding to that, 

Fig. 5 Sugar content of model breads manufactured with addition of maltogenic α-amylase preparations from Geobacillus stearothermophilus (III), (IV) and (V). Samples were taken 2 h, 22 h, 48 h, and 96 h after baking. Values are displayed as means ± standard deviation (n = 4). Different letters represent significant differences in content for each sugar, respectively (p < 0.05)

Fig. 6 Sugar content of model breads manufactured with addition of maltogenic α-amylase preparations from Geobacillus stearothermophilus (VI) and (VII). Samples were taken 2 h, 22 h, 48 h, and 96 h after baking. Values are displayed as means ± standard deviation (n = 4). Different letters represent significant differences in content for each sugar, respectively (p < 0.05)
the content of the main reaction product, maltotetraose, was about 219% higher with a mean of 6.7 mg/g \text{dm} (I) compared to 2.2 mg/g \text{dm} (II) 2 h after baking. A similar result was described by Bae et al. when using *Pseudomonas saccharophila* maltotetraogenic amylase. An increase of 200% of maltotetraogenic amylase activity in whole wheat bread making resulted in an increase of 145% of maltotetraose in their study [31].

Overall, only minor changes in the sugar composition were detected during the 48 h of storage of the breads containing preparation (I) and especially no changes in maltotetraose content. This agrees well with the results of Bosmans et al. [8] who also observed no significant changes in the content of maltooligosaccharides in wheat bread during storage using *Pseudomonas saccharophila* maltotetraogenic amylase. Bosmans et al. [8] and Derde et al. [9, 14] assumed inactivation of the maltotetraogenic amylase during baking. We detected a very low residual activity (1.1% of the applied activity) for (I), but apparently this did not cause major changes in the sugar composition of the bread during storage. Opposed to that, the residual activity of (II) was also very low (1.9%), but the absolute maltotetraose content increased from 2.1 mg/g \text{dm} 2 h after baking up to 29.5 mg/g \text{dm} after 96 h. Thus, maltotetraose increased by 1112% during 48 h and 1300% during 96 h of storage. An additional effect was the formation of maltopentaose, maltohexaose, maltoheptaose and maltooctaose after 48 h and 96 h. These sugars were not present up until 22 h of storage and could be products caused by endohydrolysis of starch chains [9].

Besides its exo-action on starch molecules, *G. stearothermophilus* maltogenic α-amylase performs various endo-attacks on starch [9], with maltose and higher maltooligosaccharides as main products [8, 32]. Our study also confirmed these findings, because the content of maltose and maltooligosaccharides (40.0 mg/g \text{dm} for (V) to 66.0 mg/g \text{dm} for (VII)) was higher in all breads with maltogenic α-amylase preparations 2 h after baking compared to the control bread (17.3 mg/g \text{dm}). In addition, preparation (VI) caused the formation of maltopentaose and maltohexaose, which were not detected in the control. It is also known that maltogenic α-amylase can release glucose from amylose and amylpectin [32]. Therefore, the significant increase of glucose in breads made with the preparations (IV), (V), and (VI) could be a possible consequence of the residual activity.

The main action of maltogenic α-amylase on starch takes place during bread baking, but changes in maltose content during cooling and storage have also been reported for wheat bread [8, 15]. We observed significant changes for maltose and higher maltooligosaccharides for all maltogenic α-amylase preparations during storage (Figs. 5, 6), but these changes were not always the same. There was no direct correlation between the level of residual activity and the amount of sugars formed during storage for the breads made with of maltogenic α-amylase. Using preparation (IV), maltotetraose was identified for the first time at 22 h, whereas (VI) showed the largest impact on the sugar spectrum of wheat bread compared to the control, including the release of maltooligosaccharides after 96 h. In contrast, no significant increase in the sugar concentrations was determined for the breads made with preparation (III), although a residual activity of 42.3% was detected. The absolute contents of maltogenic α-amylase added in the baking experiments ranged from 0.01 mg (III) to 1.21 mg (VII), but there was no direct correlation between the content and the residual activity. This might be due to differences inherent in the preparations we used and could be studied using various dosages of each preparation. However, the intent of our study was not to provide a detailed comparison of the different preparations, but rather show how sugar concentrations are affected when using each preparation at the recommended dosage.

Considering the applied activities in bread dough (Table 1), the relative residual maltogenic α-amylase activities were between 13.9% for (VII) and 45.5% for (VI) and are likely to be responsible for the changes in mono-, di- and oligosaccharide content observed during 96 h of storage. Reichenberger et al. found a comparable relative residual activity of 17.8% for maltogenic α-amylase in their wheat baking experiment [16]. Whether the detected residual activity is related to the characteristics of bread staling or bread crumb properties must be clarified in further studies. We showed that measurements of residual enzymatic activities alone cannot provide sufficient information to evaluate a potential technological function in the final product.

**Conclusion**

In this study, we identified the specific amylases in two maltotetraogenic amylase and five maltogenic α-amylase preparations commonly applied in wheat bread making. Due to their high temperature stability, a low to medium residual amylolytic activity was found in the bread crumbs containing each of the preparations. Significant changes for glucose, maltose, and higher maltooligosaccharides were observed during storage for up to 96 h and these were most likely caused by the residual amylolytic activity. By applying a methodological approach which targets various amylases and enables the identification of the link between amylases, their residual activity and the resulting changes in the saccharide content of straight dough wheat bread, we provide the foundation for a better understanding of the residual activity of exogenous amylases in wheat bread making.

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**Declarations**

**Conflict of interest** The authors declare no competing financial interest.

**Compliance with ethics requirements** This article does not contain any studies with human or animal subjects.

**Informed consent** Not applicable.

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