Characterization of the recombinant Brettanomyces anomalus β-glucosidase and its potential for bioflavouring

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Abstract
Aim: Plant materials used in the food industry contain up to five times more aromas bound to glucose (glucosides) than free, unbound aromas, making these bound aromas an unused flavouring potential. The aim of this study was to identify and purify a novel β-glucosidase from Brettanomyces yeasts that are capable of releasing bound aromas present in various food products.

Methods and Results: We screened 428 different yeast strains for β-glucosidase activity and are the first to sequence the whole genome of two Brettanomyces yeasts (Brettanomyces anomalus and Brettanomyces bruxellensis) with exceptionally high β-glucosidase activity. Heterologous expression and purification of the identified B. anomalus β-glucosidase showed that it has an optimal activity at a higher pH (5.75) and lower temperature (37°C) than commercial β-glucosidases. Adding this B. anomalus β-glucosidase to cherry beers and forest fruit milks resulted in increased amounts of benzyl alcohol, eugenol, linalool and methyl salicylate compared to Aspergillus niger and Almond glucosidase.

Conclusions: The newly identified B. anomalus β-glucosidase offers new possibilities for food bioflavouring.

Significance and Impact of the Study: This study is the first to sequence the B. anomalus genome and to identify the β-glucosidase-encoding genes of two Brettanomyces species, and reports a new bioflavouring enzyme.

Introduction
Flavour is considered to be one of the major quality attributes of a food product and flavour compounds represent over a quarter of the world market for food additives (Mouret et al. 2015). Due to an increased environmental awareness, nowadays consumers prefer naturally produced flavours, obtained from plant materials or by fermentation, where micro-organisms add specific flavours, over chemically synthesized aroma compounds (Vanderhaegen et al. 2003). Although plant material contains volatile flavours, it also contains two- to five times as many nonvolatile aroma molecules that are bound to monosaccharides like β-glucose (i.e. β-glucoside) or more rarely disaccharides (Sarry and Günata 2004). These nonvolatile aromas or ‘aglycones’ can contribute to the food aroma when they are released from the sugar molecule (glycone). This release can occur through acidic or enzymatic hydrolysis (Li et al. 2013). Since acidic hydrolysis occurs at very low pH and induces aglycone rearrangements (Gunata et al. 1988), enzymatic hydrolysis by β-glucosidases is the preferred industrial strategy to release aglycones. β-Glucosidases are often added during food production because (i) the ingredients or micro-organisms present lack or show little β-glucosidase activity, (ii) because of glucose or ethanol inhibition of present glucosidases or (iii) because enzymatic activity of present glucosidases is inhibited by the pH or temperature of the application (Gil et al. 2005). Various β-glucosidases from moulds (Thongpoo et al. 2014), bacteria...
(Michlmayr et al. 2009) and yeasts (Wang et al. 2012) already have been characterized. Additionally, heterologous expression systems allow producing higher levels of β-glucosidases (Zietsman et al. 2010). However, some current commercial β-glucosidases show unwanted side activities. This is for example the case for AR2000, a cell extract from Aspergillus niger that is used to enhance the aroma of wines. Apart from releasing aldehydes, AR2000 also hydrolyses anthocyanins, causing decolourization of red wines (Sarry and Günata 2004). Hence, there is a need for β-glucosidases that perform well under the conditions of the production process.

Saccharomyces yeasts are often used in food production and generally release desirable aroma compounds, such as esters (Steensels et al. 2014), but show only weak and variable β-glucosidase activity (Rosi et al. 1994). More pronounced β-glucosidase activities are mainly found in non-Saccharomyces yeasts such as Brettanomyces, Candida, Debaromyces and Kloekera (Wu et al. 2014). While commonly known as spoilage organisms in wine, Brettanomyces are also part of the natural, desirable microbiome in the production of lambic and gueuze beers and wines like Château de Beaucastel (Steensels et al. 2015). Although Brettanomyces yeasts have higher β-glucosidase activity than Saccharomyces yeasts, no genetic information concerning the genes encoding β-glucosidases in these species is available (Woolfit et al. 2007; Crauwels et al. 2014; Steensels et al. 2015).

In this study, we identified the β-glucosidase-encoding genes in Brettanomyces anomalus by high-quality whole-genome sequencing. The β-glucosidase was heterologously expressed, purified and thoroughly characterized. We found that the B. anomalus β-glucosidase performs optimal at pH 5.75 and 37°C, showing higher activity than commercial enzymes. Moreover, we found significant aroma differences in cherry beers and fruit milk beverages treated with the B. anomalus or commercial β-glucosidases.

Materials and methods

Reagents and enzymes

The following products were purchased from Sigma Aldrich (Diegem, Belgium): 2-mercaptoethanol (99%), almond β-glucosidase (6-6 U mg⁻¹), amygdalin (>99%), arbutin (>98%), citric acid monohydrate (>98%), D-glucose monohydrate (>98%), imidazole (>99%), Luria-Bertani (LB) medium, magnesium acetate tetrahydrate (>99%), PBS (10× concentrate), salicin (>99%), SDS and sorbitol (>98%). Ammonium acetate (>98%), EDTA, ethanol, HCl (37%), sodium acetate trihydrate, sodium chloride, disodium hydrogen phosphate dihydrate (>99%) and sodium hydroxide (99.5%) were purchased from VWR (Herelle, Belgium). Other products used in this study include 4-nitrophenol (Fluka, Diegem, Belgium), agar (Invitrogen, Ghent, Belgium), amino acid mix with (NH₄)₂SO₄ (MP Biomedicals, Brussels, Belgium), AR2000 (A. niger cell extract showing glucosidase activity; DSM, Brussels, Belgium), bactopeptone (BD Bioscience, Erembodegem, Belgium), cellulbiose (>98%; Fluka, Diegem, Belgium), DNasel (Roche, Vilvoorde, Belgium), glycerol (99%; Biosolve BV, Valkenswaard, Netherlands), Glucose Oxidase-Phenol 4-Aminoantipyrine Peroxidase (GOD-PAP) (Dialab, Belsele, Belgium), hop pellets (Saaaz Saaaz), Isopropyl β-D-1-thiogalactopyranoside (IPTG; >99%; Biosolve BV, Valkenswaard, Netherlands) isomerized hop extract (Brewferm, Beverlo, Belgium), isopropanol (>99.8%, Labscan, Zedelgem, Belgium), kanamycin sulphate (Gilco, Ghent, Belgium), malt extract light (7–12 EBC; Brouwland, Beverlo, Belgium), protease inhibitor cocktail (Roche), RNase (Westburg), sour cherry extract (Alcoferm, Beverlo, Belgium), Tris Base (Formedium, Hunstanton, UK), yeast extract (LabM, Brussels, Belgium), yeast nitrogen base (YNB; MP Biomedicals, Brussels, Belgium) and zymolyase (AMS BIO, Abingdon, UK).

Screening of yeast collection for β-glucosidase activity and ethanol sensitivity

The β-glucosidase activity of 428 yeast strains from different fermentation industries (Table S1) was assessed by spotting assays on agar plates with a β-glucosidase (cellobiose, salicin or arbutin; Table S2) as carbon source. Spottings on 2% glucose and 2% ethanol plates were added as growth control. Lab strain BY4741 (Brachmann et al. 1998) was used as negative control as it does not show any β-glucosidase activity. Single colonies fromYPD plates were grown overnight in 600 μl yeast extract peptone dextrose (YPD) at 30°C and 200 rev min⁻¹. OD₆₀₀ was measured, cultures were diluted to OD 1 and strains were spotted on the agar plates. Growth was evaluated by checking colony size after 48 h of incubation (72 h for nonSaccharomyces) at 30°C.

Whole-genome sequencing of YV396 and YV397 and genome comparison

Genomic DNA of YV396 (B. anomalus) and YV397 (Brettanomyces bruxellensis) was sequenced by the Beijing Genome Institute (BGI) using paired-end Illumina HiSeq sequencing. One paired-end library with 86 bp read length and 500 bp insert size and two mate pair libraries with 49 bp read length and 2000 and 5000 bp insert sizes were developed for each sample. Initial average coverages for each library were 127×, 40× and 40× respectively.
Before de novo sequencing, clean reads were obtained using the following filters: (i) Remove reads with more than 10% ‘N’ basepairs; (ii) remove reads with more than 20 bp with low quality (<20); (iii) Remove reads showing an overlap of at least 15 bp with the adapter sequence; (iv) Remove the first 4 bp of each read. Because the 2000 bp and 5000 bp libraries are obtained by DNA circularization, the following additional filters were applied for these libraries: (i) Remove reads with significant poly-A structure and (ii) Remove reads with k-mer frequency equal to 1. De novo assembly was performed using SOAPDENovo software. YV396 was assembled in 30 scaffolds with an N50 value of 1-55 Mbp and a total assembly size 12-88 Mbp. YV397 was assembled in 85 scaffolds with an N50 of 732 Kbp and a total assembly size of 13-06 Mbp (Crauwels et al. 2014). The software GENMARK-ES (Ter-Hovhannisyan et al. 2008) was used to perform gene prediction on the scaffolds obtained with SOAPDENovo. BLASTP searches were performed on the predicted genes to identify potential orthologs of β-glucosidase genes in the genes annotated using GENMARK-ES. Sequences were compared with known β-glucosidase genes from Aspergillus aculeatus (Genbank Acc. No. JN121996.1), Kluyveromyces marxianus (Genbank Acc. No. X05918.1) and Trichoderma reesei (Genbank Acc. No. XP_006969215.1) using CLUSTALW to identify the β-glucosidase-encoding genes of YV396 and YV397.

Construction of synthetic DNA fragments

Synthetic DNA fragments containing the β-glucosidase gene of YV396 (B. anomalus), YV397 (B. bruxellensis) or YV404 (K. marxianus), a purification (Lys6) and immobilization tag (His6), the Shine-Dalgarno sequence, a XbaI and XhoI recognition sequence and two linkers that allow using the enzyme without His6 and/or Lys6 tag (Fig. S1) were synthesized by Geneart. The constructs were codon optimized (Fig. S2) for expression in Escherichia coli.

Production of the Brettanomyces anomalus (YV396) β-glucosidase

Escherichia coli strain BL21 (DE3) was transformed with pET28-YV396-His6. Bacteria were grown overnight at 37°C in LB-kanamycin (50 µg ml⁻¹) before 100× dilution in a 20 l fermenter with LB-kanamycin (50 µg ml⁻¹) and 1% glycerol. The initial stirring and airflow were respectively 200 rev min⁻¹ and 1·5 l min⁻¹. Further, flow was automatically adapted to keep pO₂ at 30% and temperature at 37°C until cells were grown to OD 0·8. Expression was induced by adding 0·5 mmol l⁻¹ IPTG overnight at 20°C and mostly occurred in the cytoplasm and, to lesser extent, in inclusion bodies (see Fig. S3). Cells were harvested and frozen at −20°C. Two independent fermentations yielded 240 and 229 g of cells. After thawing, cells were resuspended at 3 g ml⁻¹ in 20 mmol l⁻¹ NaH₂PO₄ pH 7·4, 500 mmol l⁻¹ NaCl, 20 mmol l⁻¹ imidazole, one tablet 100 ml⁻¹ protease inhibitor cocktail and 1 mg 100 ml⁻¹ DNasel. Cells were sonicated (Sonic Ultra Cell with 13 mm probe, Sonics & Materials, Newtown, CT, USA) on ice at 70% amplitude for 24 cycles of 5 s with 9 s between different cycles. The cytoplasmic suspension was centrifuged at 18 000 g for 30 min at 4°C. Next, the supernatant was applied to the Äkta purification system equipped with a 1 ml Ni-Sepharose 6 FF chromatography column (GE Healthcare, Diegem, Belgium). In this column, the histidine tag of the protein will interact with Ni²⁺ ions as described by Ueda et al. (2003). The column was equilibrated with 20 mmol l⁻¹ NaH₂PO₄ pH 7·4, 500 mmol l⁻¹ NaCl, 20 mmol l⁻¹ imidazole to base line, washed with 10 column volumes of 20 mmol l⁻¹ NaH₂PO₄ pH 7·4, 20 mmol l⁻¹ NaCl, 400 mmol l⁻¹, 50 mmol l⁻¹ imidazole and eluted in 10 column volumes of 400 mmol l⁻¹ imidazole in the same buffer. All flow rates were adjusted to 3·5 ml min⁻¹ and 2·2 ml min⁻¹ for DE3 and BL21 (DE3), respectively 200 rev min⁻¹.
were 1 ml min$^{-1}$. Detection was done at 280 nm. The protein of interest eluted in the 400 mmol l$^{-1}$ imidazol fraction in a rather broad peak spread over the 10 column volumes. The elution fraction from the Ni-Sepharose 6FF column was injected on a Superdex 200 XK26 × 65 column with PBS as running solution (4 ml min$^{-1}$) for formulation and to remove minor contaminants. The obtained purified fractions were analysed by SDS-PAGE in reducing conditions (Fig. 1) and protein concentration was measured using the Micro-BCA assay with BSA standard. The two independent purifications yielded 47 and 48 mg of protein in 28 (1.68 mg ml$^{-1}$) and 30 ml (1.6 mg ml$^{-1}$).

Optimal pH and temperature, thermostability and apparent $K_M$ and $k_{cat}$ values of the different enzymes

Optimal pH and temperature, thermostability and apparent $K_M$ and $k_{cat}$ values were determined using an enzymatic assay that was performed in the linear range of product formation (Fig. S4) in 96 well PCR blocks (Bio-Rad, Nazareth, Belgium; C1000 Thermal Cycler). Twenty-five microlitres of McIlvaine buffer (0.1 mol l$^{-1}$ citric acid, 0.2 mol l$^{-1}$ Na$_2$HPO$_4$, pH specific), 15 μl substrate solution [25 mmol l$^{-1}$ cellobiose, amygdalin or salicin in McIlvaine buffer (pH specific and various concentrations)] and 10 μl enzyme solution (3.78*10$^{-2}$ and 1.89*10$^{-2}$ μmol l$^{-1}$) or 10 μl McIlvaine buffer (blank) were mixed and incubated for 1 h at specific temperature, after which a 10 min denaturation step at 95°C was applied. Next, 30 μl McIlvaine buffer was added. Fifty microlitres of sample was added to 100 μl of GOD–PAP, incubated for 15 min at 30°C and 200 rev min$^{-1}$ and absorbance was measured at 505 nm.

Optimal pH and temperature were determined by performing this assay for cellobiose at pH ranging from 3 to 7 at 30°C and temperatures ranging from 15 to 70°C at pH 5 respectively. We determined the amount of cellobiose that was hydrolysed per minute, which was normalized to the condition with the highest activity value.

Thermostability of the $\beta$-glucosidases was measured by incubating the enzymes 4 h in PBS at 4, 30, 40, 50, 60, 70 or 80°C after which the enzymatic assay was performed for cellobiose at 30°C and pH 5. Enzymatic activity of the heat-treated enzymes was normalized to the untreated samples (incubation at 4°C).

$K_M$ and $k_{cat}$ values were measured for cellobiose, amygdalin and salicin (0.15–60 mmol l$^{-1}$) at pH 4.5 and 17°C and pH 5.75 and 37°C. Equimolar amounts of B. anomalus and almond $\beta$-glucosidase and the average mass amount of B. anomalus and almond $\beta$-glucosidase for AR2000 were used, considering this is a crude cell extract. Enzymatic activity was calculated as the amount of glycoside hydrolysed per minute, $k_{cat}$ values as enzymatic activity per enzyme concentration. Data were analysed using GRAPHPAD PRISM 6.01 (La Jolla, CA, USA).

Industrial applications

The different $\beta$-glucosidases were tested in beer fermentations and fruit milk beverages. For beer samples, 600 ml 12°P malt with 4 ml isomerized hop extract or cherry extract or 400 mg hop pellets were pitched with $10^7$ cells YV15 ($\beta$-glucosidase negative ale strain, Table S3), per ml and 10-5 nmol B. anomalus or almond $\beta$-glucosidase or the average weight amount of protein of AR2000. Beers were fermented for 7 days at 18°C. Fruit milk beverages were prepared by defrosting, mixing and centrifuging frozen forest or tropical fruit. Next, 180 ml of ultra-high temperature processed (UHT) milk was mixed with 20 ml pasteurized juice (12 min at 80°C) and 3-5 nmol B. anomalus or almond $\beta$-glucosidase or the average weight amount of protein of AR2000 and incubated for 24 h at 37°C.

Sensory analysis

The aroma of beers and milk beverages was evaluated by a test panel consisting of 20 untrained persons. Samples with B. anomalus $\beta$-glucosidase were compared to samples with AR2000, almond or without enzyme. Differences were significant when 55% of the panelists indicated the same sample as being different ($\alpha=0.05$, $\beta=0.10$ and $P_d=50%$), and next, these samples were subjected to headspace-solid phase microextraction-gas chromatography-mass spectrometry (HS-SPME-GC-MS) and preference tests. In preference tests, panelists had to indicate their preference for the aroma of samples with B. anomalus $\beta$-glucosidase or samples with AR2000, almond or without enzyme. A significant preference was noted when at least 75% of the panelists preferred the same sample ($\alpha=0.05$, $\beta=0.40$ and $p$(preference for a specific aroma) = 75%) (Meilgaard et al. 2007).

Identification and quantification of volatile compounds in beverages

Aroma compounds of beverages with significant differences were identified and quantified by HS-SPME GC-MS. A 20 ml vial with 5 ml of sample, 1.75 g NaCl and 5 μl of internal standard (IS; 2-heptanol, 250 μg ml$^{-1}$) was immersed in a 40°C water bath. After 5 min of equilibration, a triphase DVB/Carboxen/PDMS 50/30 μm SPME fibre (Supelco Co., Bellefonte, PA) was exposed to the headspace for 30 min. Next, volatiles were desorbed in the GC-MS (Shimadzu, Brussels, Belgium; QP2010

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Ultra Plus) by heating for 5 min at 250°C. The GC-MS was equipped with a HP-5 ms nonpolar column (Agilent, Diegem, Belgium; 30 m × 0.25 mm i.d., 0.25 μm thin layer), helium was used as carrier gas at a pressure of 100 kPa and samples were injected in split/splitless mode. The temperature program is displayed in Fig. S5. The mass detector was operated in scan mode (35–500 amu) using electronic impact ionization (70 eV). The interface and detector were kept at 250°C.

Linear n-alkanes (C8 to C19) were injected in the GC-MS as external retention index (RI) markers. RI of aglycones were calculated using cubic spline interpolation. Compounds were identified using AMDIS ver. 2.71, followed by matching deconvoluted spectra to commercial GC/MS libraries described in Robert and Adams (2007) and Mondello (2011). RI of aglycone standards were used to confirm the identification. Since some compounds coeluted, analysis was performed by integrating over characteristic ions using OPENCHROM (ver. 0.9.0). Next, integrated areas were converted back to a total ion scale based on the relative abundance of each characteristic ion in the spectrum of each compound. Relative concentrations were calculated by normalizing to the 2-heptanol IS peak and comparing the log-transformed relative peak area of each compound across treatments using a linear mixed model analysis, in which treatment and biological replicate nested within treatment were coded as fixed and random factors. Treatment averages were compared in a pairwise fashion using Tukey’s post hoc tests and final significance levels were corrected for multiple testing across all compounds using the Bonferroni procedure with R package LME4. In addition, we calculated the absolute concentration (in ppb) of aglycones based on their measured response factor relative to the IS.

Results

Screening of yeast strains

In order to identify yeast strains with high β-glucosidase activity, a high-throughput screen of 428 strains (Table S1) was performed doing spotting assays. Forty-five percentage of the Saccharomyces strains and 47% of the non-Saccharomyces yeasts grew on all β-glucoside media. Two Brettanomyces strains that were isolated from spontaneous beer fermentations (YV396; B. anomalus and YV397; B. bruxellensis) had pronounced growth on β-glucoside and ethanol media (Table S3). Kluyveromyces marxianus (YV404), a
Table 1 Kinetic parameters of different β-glucosidases measured on different substrates, pHs and temperatures. Apparent $K_M$ and $k_{cat}$ values ($n = 2$) were determined excluding data points that showed lower enzymatic activity values because of product inhibition, caused by glucose accumulation (Fig. S8)

| Substrate | Enzyme          | $K_M$ (mmol l$^{-1}$) | $k_{cat}$ (s$^{-1}$)* | $K_M$ (mmol l$^{-1}$) | $k_{cat}$ (s$^{-1}$)* |
|-----------|-----------------|-----------------------|------------------------|-----------------------|------------------------|
| Cellobiose| AR2000          | 0.60 ± 0.05           | 1.09 ± 0.04            | 0.52 ± 0.09           | 1.01 ± 0.09            |
| Almond    | 0.18 ± 0.06     |                       | 0.05 ± 0.01            | 0.34 ± 0.08           | 0.06 ± 0.02            |
| Brettanomyces | anomalus         | 1.15 ± 0.16           | 0.07 ± 0.01            | 15.06 ± 0.73          | 2.73 ± 0.32            |
| Amygdalin | AR2000          | 0.40 ± 0.05           | 0.93 ± 0.04            | 0.12 ± 0.03           | 0.93 ± 0.16            |
| Almond    | 10.30 ± 0.09    |                       | 1.11 ± 0.11            | 10.08 ± 0.13          | 3.45 ± 0.56            |
| B. anomalus |                 | 0.09 ± 0.03           | 0.07 ± 0.02            | 1.00 ± 0.05           | 0.31 ± 0.002           |
| Salicin   | AR2000          | 0.19 ± 0.06           | 1.08 ± 0.14            | 0.19 ± 0.06           | 0.86 ± 0.08            |
| Almond    | 10.95 ± 2.00    |                       | 0.43 ± 0.01            | 19.65 ± 2.06          | 0.68 ± 0.01            |
| B. anomalus |                 | 6.49 ± 1.11           | 0.72 ± 0.02            | 1.01 ± 0.18           | 4.21 ± 0.34            |

* $k_{cat}$ values for AR2000 are expressed in (μmol substrate s$^{-1}$ genzyme$^{-1}$) as AR2000 is a crude cell extract.

Table 2 Triangle tests of different beers and fruit milk beverages. Beers with isomerized hop extract (B + HE), cherry extract (B + CE) or hop pellets (B + HP) and milk beverages with tropical (M + T) – and forest fruit juice (M + F) were subjected to triangle tests by 20 untrained panelists. Cherry beers and milk beverages with forest fruit juice showed significant differences across the enzyme treatments. Beer with hop extract and Brettanomyces anomalus β-glucosidase was also significant different from the untreated beer.

| Comparison                           | % correct identifications |
|--------------------------------------|---------------------------|
|                                      | Beers         | Milk beverages |
|                                      | B + HE | B + CE | B + HP | M + T | M + F  |
| Brettanomyces anomalus enzyme vs no enzyme | 55*  | 55*   | 40    | 50    | 75*    |
| B. anomalus vs almond β-glucosidase   | 35    | 40    | 30    | 30    | 75*    |
| B. anomalus β-glucosidase vs AR2000   | 20    | 60*   | 40    | 50    | 75*    |

*Significant differences were observed between treatments at $\alpha = 0.05$, $\beta = 0.10$ and $\rho_0$ (the ability of a person to smell a certain difference) = 50% (Meilgaard et al. 2007).

Whole-genome sequencing and construction of synthetic DNA fragments

The assembled high-coverage genome sequences of B. bruxellensis (YV397) and B. anomalus (YV396) and their β-glucosidase-encoding genes (with a gene size of 2526 and 2523 bp respectively) are freely available from Genbank under accession numbers PRJNA244003, PRJNA281311, KR181959 and KR181960 respectively. The Brettanomyces β-glucosidases showed most similarity to the K. marxianus GH3 β-glucosidase (Fig. S6). We identified several conserved amino acid regions, including a typical FGYGLSY domain, and predicted catalytic residues (aspartic acid and glutamic acid; Henrissat 1991; Sarry and Günata 2004; Quatrich et al. 2008). Based on several Swiss-Models (ExPASy, Swiss Institute of Bioinformatics, Lausanne, Switzerland), the isolated Brettanomyces β-glucosidases are probably monomeric.

The codon-optimized DNA constructs containing the YV396, YV397 or K. marxianus GH3 β-glucosidase (Fig. S1) were expressed in E. coli strain BL21(DE3) and purified on a Talon Metal Affinity Resin column (Clontech, Leusden, Netherlands). Enzymatic activity was assessed on arbutin, salicin and cellobiose. Since the Brettanomyces anomalus GH3 β-glucosidase (Fig. S6) showed highest specific activity (data not shown), large scale heterologous production and further experiments focused on this enzyme.

Figure 4 Volatile compounds in cherry beers with different enzyme treatments (n = 6). The aroma of cherry beers treated with the Brettanomyces anomalus β-glucosidase differed significantly from beers without enzyme (a), with Almond β-glucosidase (b) or AR2000 (c). Colour codes indicate relative concentrations, which were calculated as z-scores of log-transformed peak areas normalized relative to the internal standard (IS) peak. Significance levels were based on Tukey’s post hoc tests and linear mixed model analysis on log-transformed relative peak areas, in which biological replicate and treatment were coded as random and fixed factors. All $p$-values are Bonferroni adjusted to correct for multiple testing across all 44 integrated compounds.
| Compound                      | P-value |
|-------------------------------|---------|
| Eugenol                      | 1E-13   |
| Phenethyl hexanoate          | 3E-8    |
| Ethyl octanoate              | 3E-7    |
| Benzyl alcohol               | 0·01    |
| Ethylhexanoate               | 3E-12   |
| 2-phenethyl acetate          | 1E-13   |
| Isoamyl octanoate            | 1E-13   |
| Ethyl decanoate              | 1E-10   |
| Isomaltoe                    | 2E-10   |
| Dodecanoic acid              | 1E-13   |
| Phenethyl hexanoate          | 1E-13   |
| Ethyl octanoate              | 1E-13   |
| 2-methylbutyl acetate        | 0·03    |
| Isoamyl octanoate            | 1E-7    |
| Ethyl decanoate              | 2E-8    |
| Ethyl hexanoate              | 1E-13   |
| 2-phenethyl acetate          | 1E-6    |
| 2-acetyl-furan               | 0·005   |
| Para-vinyl-guaiacol          | 1E-5    |
| Phenylacetaldehyde           | 0·003   |
| Octanol                      | 0·0001  |
| 2-heptanone                  | 1E-13   |
| Tolualdehyde                 | 1E-13   |
| Isomaltoe                    | 1E-13   |

(a) Alcohol (control)  

(b) Almond

(c) AR2000

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B. anomalus β-glucosidase
Optimal pH and temperature and thermostability of \( \beta \)-glucosidase enzymes

After large scale production of the YV396 \( \beta \)-glucosidase, we determined optimal pH and temperature, thermostability and apparent \( K_M \) and \( k_{cat} \) values of the \textit{B. anomalus} \( \beta \)-glucosidase, and compared it to the commercially available almond GH1 and \textit{A. niger} (AR2000) GH3 \( \beta \)-glucosidases (Pozzo \textit{et al.} 2010; Zhao \textit{et al.} 2013).

### Table 1: Compound Comparison

| Compound                  | P-value      |
|---------------------------|--------------|
| Methyl salicylate         | < 1E-13      |
| Benzyl alcohol            | < 1E-13      |
| Dodecanal                 | 0.001        |
| Linalool                  | 5E-05        |
| 2-nonanone                | 0.04         |

### Table 2: Compound Comparison

| Compound                  | P-value      |
|---------------------------|--------------|
| 2-heptanone               | 0.002        |
| Nonanoic acid             | 0.01         |
| 2-undecanone              | 2E-05        |
| Delta-decalactone         | 1E-05        |
| Gamma-decalactone         | 0.0004       |
| Dodecanal                 | 5E-10        |
| 2-nonanone                | 0.0007       |
| Benzyl alcohol            | 2E-13        |
| p-tolualdehyde            | 0.0005       |

### Table 3: Compound Comparison

| Compound                  | P-value      |
|---------------------------|--------------|
| Methyl salicylate         | < 1E-13      |
| Benzyl alcohol            | 6E-07        |
| Dodecanal                 | < 1E-13      |
| Gamma-dodecalactone       | 6E-10        |
| Dodecanoic acid           | 0.003        |
| Linalool                  | 0.0003       |
| Gamma-decalactone         | 1E-05        |
| Delta-decalactone         | 0.005        |
| p-tolualdehyde            | 2E-05        |

### Figure 5: Volatile compounds in fruit milk beverages with different enzyme treatments (n = 4).

The aroma of milk beverages treated with the \textit{Brettanomyces anomalus} \( \beta \)-glucosidase differed significantly from fruit milk beverage without enzyme (a), with Almond \( \beta \)-glucosidase (b) or AR2000 (c). Colour codes indicate relative concentrations, which were calculated as described in Fig. 4.
The β-glucosidases showed striking differences in their optimal pH and temperatures (Fig. 2). Optimal activity for A. niger (AR2000, used in wine industry), almond and B. anomalus β-glucosidase was reached at pH 4.5, 5.0 and 5.75 and at 58°C, 50°C and 37°C respectively. Also their thermostability differed: the B. anomalus β-glucosidase showed very low relative thermostability compared to both AR2000 and Almond glucosidase, although its absolute enzymatic activity was higher than Almond glucosidase and, when incubated at temperatures exceeding 50°C, higher than AR2000 (Fig. 3 and Fig. S7). Further enzymatic engineering can be useful to increase the thermostability and thus enzymatic activity of the B. anomalus glucosidase.

**Apparent K_M and k_cat values of the β-glucosidases**

K_M and k_cat parameters describe the enzyme’s affinity for a substrate and the amount of substrate each enzyme site can convert per unit of time respectively. We determined these parameters for the different enzymes (B. anomalus, AR2000 and almond) for the substrates cellobiose, amygdalin and salicin at pH 4.5 at 17°C (simulating the end of beer fermentations) and pH 5.75 at 37°C (optimal conditions for the B. anomalus β-glucosidase). We report k_cat and K_M as apparent values since the purity of enzyme solutions used is below 100%. Enzymatic activity plots and a summary of the apparent K_M and k_cat values are shown in Fig. S8 and Table 1. All enzymes followed Michaelis–Menten kinetics. AR2000 showed low K_M values, indicating its high affinity for β-glucosides. Measured k_cat values confirmed that the B. anomalus enzyme works optimal at pH 5.75 and 37°C. The almond glucosidase had particularly high k_cat values for amygdalin, a glycoside commonly found in Rosaceae, proving that this enzyme is very suitable to hydrolyse these glycosides. The almond β-glucosidase showed high K_M values for amygdalin and salicin as well.

**Triangle tests of beers and milk beverages treated with β-glucosidase enzymes**

In a next step, the β-glucosidases were added to beer fermentations (with hop or cherries) and tropical or forest fruit milk beverages in order to assess their effect on the aroma of these beverages. These beverages were chosen as their pH (4.5–5.2) was close to the pH optimum of the B. anomalus glucosidase. After production, a test panel evaluated the aroma of the beverages using triangle tests. We found significant differences between cherry beers with the B. anomalus enzyme and cherry beers with AR2000 or without enzyme. Also, differences were found between hopped beers with B. anomalus β-glucosidase and without enzyme (Table 2). However, the hop extract...
was made by CO₂ extraction of hop pellets which may lead to lower concentrations of glycosides in the hop extract, since glycosides are water-soluble (Sarry and Günata 2004). Beers with hop pellets and different enzymes did not show any difference. Among the milk beverages, only forest fruit milks with different enzyme treatments were significantly different (Table 2). Given these results, cherry beers and forest fruit milk beverages were analysed with HS-SPME GC-MS and preference tests.

Volatile compounds released by β-glucosidase enzymes in cherry beers and forest fruit milk beverages

HS-SPME-GC-MS results of cherry beers and forest fruit milks are shown in Figs 4 and 5. Although many volatile compounds were found, we focused our analyses on relative changes in the amount of aglycones.

Beers with B. anomalus β-glucosidase contained more eugenol (clove, honey aroma) and benzyl alcohol (sweet, flower) than untreated beers. Also, the B. anomalus glucosidase released more eugenol and less linalool (citral, flower; Mosciano 1994, accessed 13th January 2015) than AR2000. When looking at absolute concentrations of different aglycones (Table 3), it is clear that benzaldehyde (almond, cherry), linalool, eugenol, beta-damascenone (honey, apple, peach) and geraniol (rose, lemon, flower) for all enzymes exceeded their odour threshold concentration (Meilgaard 1975; Acree and Heinrich 2014, accessed 12th October 2014).

Forest fruit milk beverages with B. anomalus β-glucosidase contained more methyl salicylate (peppermint, wintergreen; Meilgaard 1975), benzyl alcohol and linalool than untreated beverages or beverages with AR2000. Compared to almond β-glucosidase, the B. anomalus enzyme released more benzyl alcohol (Fig. 5). Considering that across β-glucosidase enzyme treatments different aglycones exceeded their threshold concentration (Table 3), the resulting aroma of the beverages might be different. For this reason, beverages with different β-glucosidases were subjected to double-blind consumer preference tests to see if the aroma released by a specific enzyme was preferred.

Preference tests of beers and milk beverages treated with β-glucosidase enzymes

In preference tests, the sensory panel indicated their preference for cherry beers with B. anomalus glucosidase over cherry beers with AR2000 (Fig. 6a). They described the beer with B. anomalus β-glucosidase as fruitier and more cherry- and honey like, which corresponds to the increased release of eugenol (honey). Consumers also preferred forest fruit milk with the B. anomalus enzyme over milk with almond β-glucosidase (Fig. 6b) which they attributed to a more spicy character of these beverages, as methyl salicylate exceeded its threshold concentration, while the aroma of almond treated milk was more neutral. Since spicy aromas usually are not associated with fruited milks, this also explains why 70% of the test panel preferred the untreated sample over the sample with the B. anomalus enzyme (although this preference was not significant).

Discussion

Screening of 428 yeast strains for β-glucosidase activity revealed strong activity in Brettanomyces strains. Although these yeasts are generally considered safe for food production, using them is often not possible as they are notorious for the formation of off-flavours (Steensels et al. 2015). This study is the first to identify the β-glucosidase-encoding genes in B. anomalus and B. bruxellensis, and thus represents a first step towards using (enzymes of) these yeasts without their off-flavours. After genome sequencing and identification of the β-glucosidase-encoding genes, we expressed the codon-optimized genes in E. coli and isolated the resulting enzymes. Since the B. anomalus β-glucosidase showed the highest enzymatic
activity, we thoroughly characterized this protein by comparing it to commercially available β-glucosidases (AR2000 and almond β-glucosidase).

AR2000, B. anomalus and almond β-glucosidase worked optimally at pH 4.5 and 58°C, pH 5-75 and 37°C and pH 5 and 50°C respectively. For the A. niger (AR2000) β-glucosidase, optimal conditions between pH 4–4.5 and 55°C and 70°C, are described (Placzek 2014, accessed 12th November 2015). Although we found strongly reduced enzymatic activity at pH 3 and significant activity above pH 5, the packaging of AR2000 claims that it works optimally between pH 2-8 and 5, which might be a matrix effect. The results for almond β-glucosidase do not agree with values described by Woodward and Wiseman (1982) (optimal pH of 5-6 on cellobiose). As the B. anomalus enzyme has a lower optimal temperature than AR2000 and almond glucosidase, it can be validated in ‘green’ food and biofuel applications as it does not require heating. However, although kcat values indicated considerable activity at lower pH, the relative high pH optimum of the B. anomalus β-glucosidase might be an issue for bioflavouring as most food products are slightly acidic. Hence, we tested the enzymes in applications that are characterized by a higher pH value like beer fermentations (pH 5-4 at the start of fermentation) and fruit milks (pH 5-5).

Cherry beers with B. anomalus β-glucosidase contained significantly more benzyl alcohol and eugenol, which contribute to the typical aroma of cherry beers, which was preferred over samples with AR2000 (Schmid and Wiseman (1982) (optimal pH of 5-6 on cellobiose)). As the B. anomalus enzyme has a lower optimal temperature than AR2000 and almond glucosidase, it can be validated in ‘green’ food and biofuel applications as it does not require heating. However, although kcat values indicated considerable activity at lower pH, the relative high pH optimum of the B. anomalus β-glucosidase might be an issue for bioflavouring as most food products are slightly acidic. Hence, we tested the enzymes in applications that are characterized by a higher pH value like beer fermentations (pH 5-4 at the start of fermentation) and fruit milks (pH 5-5).

The aroma of forest fruit milks also differed depending on the β-glucosidase used. Since the forest fruit was a mixture of different fruits, it was difficult to predict which aglycones were expected. However, all molecules listed in Fig. 5 (except p-toluualdehyde) contribute to a characterizing strawberry aroma (Zabetakis and Holden 1997). The amount of glycosides hydrolysed was clearly influenced by the different pH and temperature optimum of the enzymes, as the almond enzyme released more aglycones in cherry beers, while the B. anomalus β-glucosidase was much more active in fruited milk.

In conclusion, we identified, purified and characterized a novel B. anomalus β-glucosidase that releases naturally present but hidden food flavours. Industrial application of this enzyme can be met by adding it during food production or by using the immobilization tag of the enzyme. Additionally, this study provides the first high-quality genome sequence of B. anomalus and B. bruxellensis, opening the door for further exploration of their genomic properties.

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Conflict of Interest

No conflict of interest declared.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

**Figure S1** Synthetic DNA construct used to introduce the different glucosidase genes in the pET28 plasmid.

**Figure S2** Codon-optimized \(\beta\)-glucosidase genes of *Brettanomyces anomalus* (YV396) and *Brettanomyces bruxellensis* (YV397)

**Figure S3** Western blot (anti-His) of cytoplasmic and inclusion body fraction before and after induction of *B. anomalus* \(\beta\)-glucosidase synthesis.

**Figure S4** Enzymatic product formation by AR2000 (\(\sim 4.32 \times 10^{-3}\) g l\(^{-1}\) and \(\sim 2.16 \times 10^{-3}\) g l\(^{-1}\)), Almond \(\beta\)-glucosidase (\(\sim 3.78 \times 10^{-2}\) \(\mu\)mol l\(^{-1}\) and \(\sim 1.89 \times 10^{-2}\) \(\mu\)mol l\(^{-1}\)) and *B. anomalus* \(\beta\)-glucosidase (\(\sim 3.78 \times 10^{-2}\) \(\mu\)mol l\(^{-1}\) and \(\sim 1.89 \times 10^{-2}\) \(\mu\)mol l\(^{-1}\)) for cellobiose, amygdalin and salicin at pH 4.5 and 17°C, pH 5 and 17 or 37°C and pH 5.75 and 37°C at different incubation times.

**Figure S5** Temperature program used during GC-MS analysis.

**Figure S6** Alignment of the amino acid sequences of the GH3 \(\beta\)-glucosidase enzymes of *Brettanomyces bruxellensis*, *Brettanomyces anomalus* and *Kluyveromyces marxianus*.

**Figure S7** Absolute enzymatic activity of the heat-treat ed glucosidases.

**Figure S8** Michaelis–Menten plots for AR2000 (\(\sim 4.32 \times 10^{-3}\) g l\(^{-1}\) and \(\sim 2.16 \times 10^{-3}\) g l\(^{-1}\)), Almond \(\beta\)-glucosidase (\(\sim 3.78 \times 10^{-2}\) \(\mu\)mol l\(^{-1}\) and \(\sim 1.89 \times 10^{-2}\) \(\mu\)mol l\(^{-1}\)) and *B. anomalus* \(\beta\)-glucosidase (\(\sim 3.78 \times 10^{-2}\) \(\mu\)mol l\(^{-1}\) and \(\sim 1.89 \times 10^{-2}\) \(\mu\)mol l\(^{-1}\)), almond and *B. anomalus* \(\beta\)-glucosidase for cellobiose, amygdalin and salicin at pH 4.5 17°C and pH 5.75 37°C.

**Table S1** \(\beta\)-glucosidase activity of screened yeast strains.

**Table S2** Composition of agar media used for yeast screening.

**Table S3** Main results from the qualitative screening of 428 yeast strains for \(\beta\)-glucosidase activity on various agar media.