Constitutive expression of Camelus bactrianus prochymosin B in Pichia pastoris

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Abstract

Camel chymosin can be efficiently employed to produce cheese. Traditionally the rennet enzyme produced by the glands of the fourth stomach of ruminant animals (abomasum) is used in cheese making. Full-length Camelus bactrianus (Bactrian camel) prochymosin gene was synthesized and constitutively expressed in Pichia pastoris cells under glyceraldehydes-3-phosphate dehydrogenase (GAP) promoter. It was purified by sequential anion and cation exchange chromatography. SDS-PAGE analysis resulted in two bands, approximately 42 and 35 kDa. The 42 kDa band vanished when the sample was treated with endoglycosidase H, indicating that the recombinant protein is partially glycosylated. Optimal pH for the activity of the highest-purity recombinant chymosin was pH 4.5 for cow's milk and pH 4.8 for mare's milk. The range 45-50 °C and 70 °C for cow's and mare's milk types, respectively, was found to be the most appropriate for maximal relative milk-clotting activity. Concentration of CaCl₂ that ensured the stability of the chymosin milk-clotting activity was between 20 and 50 mM with an optimum at 30 mM. Milk-clotting activity of camel recombinant chymosin and ability to make curd was successfully tested on fresh mare's milk. Pichia pastoris strain with integrated camel chymosin gene showed high productivity of submerged fermentation in bioreactor with milk-clotting activity 1412 U/mL and 80 mg/L enzyme yield. These results suggest that the constitutive expression of the camel chymosin Camelus bactrianus in the yeast Pichia pastoris has good prospects for practical applications.

1. Introduction

One of the earliest biotechnological applications of enzymes is cheese production [1]. The proteolytic enzymes pepsin and chymosin [2], previously known as "rennet," [3] have been identified as active ingredients in this process. Chymosin (EC 3.4.23.4) is an aspartic peptidase that belongs to the pepsin-like family [4]. Chymosin particularly targets the κ-casein peptide link between Phe105 and Met106 (Figure 1A), triggering the formation of destabilized casein micelles, milk coagulation, and separation of the milk into curds and whey [5]. Although plants [6, 7, 8, 9, 10, 11] and microorganisms [12, 13, 14, 15, 16, 17] can be used as different sources of milk-coagulating enzymes, plant and microbial rennet are characterized by nonspecific proteolytic activity against casein, and animal rennet is more frequently used in the dairy sector for cheese making. Because of the high specific milk-clotting activity, chymosin is considered the most efficient protease for the cheese-making industry. Today, fermentation-produced chymosin accounts for more than 90% of rennet utilized, and it has the added benefit of being both kosher and halal acceptable [18]. There are chymosins, obtained from different animals, that arouse interest, for example the yak (Bos grunniens) [19] and Altai maral (Cervus elaphus) [20].

Chymosin refers to a specific action aspartic proteases [21]. Analysis of the amino acid sequences of the milk for camel, horse, human, pig, cow, goat, and sheep showed that the cleavage site of κ-casein by chymosin is the sequence SFXAIPPKK, where X is methionine in the bovids and isoleucine in other mammals (Figure 1B). Highly conserved region 97–116 of κ-casein between species [22], that κ-casein proteolysis and subsequent milk coagulation are important biological processes. Since chymosin possesses high substrate specificity, this difference may explain why camel's milk cannot be coagulated with bovine chymosin [23, 24]. At the same time, it is interesting to test the activity of camel's chymosin on mare's milk.

Mare's κ-casein contains 165 amino acids residues. Isoelectric point of equine and human κ-casein, 8.03 and 8.68 respectively, are higher than
bovine κ-casein at 5.93, and they have a net positive charge at physiological pH, whereas bovine κ-casein has a net negative charge [25]. In terms of hydrophobic dispersion along the polypeptide chain, human κ-casein appears to be more comparable to horse than bovine κ-casein [26].

Camel chymosin can be employed to produce cheese from cow’s milk [27, 28, 29, 30] and camel milk [24, 31]. In comparison, camel chymosin has 70% more milk-clotting activity than bovine chymosin and is more thermostable than bovine chymosin [23, 29], making it useful and appealing for commercial cheese manufacture. Chymosin from Camelus dromedarius is characterized well [23, 29, 32], and its recombinant version is available as commercial milk-clotting product Chy-MAX M (Ch. Hansen). The recombinant chymosin of C. dromedarius has been successfully expressed in the filamentous fungus Aspergillus niger [23] and in the yeast P. pastoris [33]. Chymosin from the Central Asian camel Camelus bactrianus is not characterized, but its full nucleotide sequence is known [34]. Analysis of the nucleotide sequence of the two camel chymosins shows that the C. bactrianus chymosin gene differs from the dromedary chymosin gene at three positions, C685G, A880T, A881C, which has 70% more milk-clotting activity than bovine chymosin and is more thermostable than bovine chymosin [23, 29, 32].

The aim of this work was secretory expression of C. bactrianus chymosin B in the yeast P. pastoris and biochemical characterization of the resultant enzyme. P. pastoris (also known as Komagataella phaffii), an eukaryotic expression system, is a methylotrophic yeast suitable as a host organism for heterologous expression of proteins owing to simplicity of its genetic manipulation, rapid growth on an inexpensive medium to high cell densities, and capability for complex post-translational modifications [37, 38]. The P. pastoris yeast expression system is a good choice for obtaining a heterologous recombinant protein with prospects for large-scale microbial production [39], and C. bactrianus chymosin holds promise for the cheese-making industry.

At the best of our knowledge, a comprehensive research on dairy products from minor species (equine, camel, yak) is not yet available in the scientific literature, due to a lack of economic relevance. The most well-known dairy product made from mare’s milk is koumiss. It is most well-known and commonly consumed in Central Asia. In the winter, temperatures in their native habitat, for example in Trans-Altaï region of Gobi desert, can drop as low as –35°C [36]. This arises interest to obtain and study C. bactrianus chymosin.

2. Materials and methods

2.1. Vectors, strains, cloning enzymes and chemicals

Vector pGAPZαA (Invitrogen, USA) was used to construct the expression cassette. Restriction enzymes EcoRI and NotI, phosphatase FastAP, T4 DNA ligase, and Phusion High-Fidelity DNA Polymerase (Thermo Fisher, USA) were employed for the amplification and cloning of the target gene. Escherichia coli strain DH5α was acquired from Thermo Fisher Scientific. P. pastoris GS115 cells were purchased from Invitrogen (USA). The chemical reagents used in this study were of molecular biology or pure analytical grade and purchased from Sigma-Aldrich (St. Louis, USA) and AppliChem (Darmstadt, Germany). The vectors and enzyme were stored at -20°C, the strains were stored at -80°C and chemicals were stored accordingly manufactured recommendations.

2.2. Synthesis of the gene and vector construction

The amino acid sequence of the C. bactrianus prochymosin gene was retrieved from GenBank (accession No. JARL00000000.1) [34]. The nucleotide sequence was codon optimized for expression in yeast. The gene was synthesized by the Macrogen Company (Korea) and was shipped in the pTOP_Blunt_V2 vector. The prochymosin gene was amplified from pTOP_Blunt_V2/ProchymCB with PCR primers ProchymCB-EcoRI (5′-CCGGAATTCCTTGGAATTACTAGAATCCTGATTG-3′) and ProchymCB-NotI (5′-ATAGTTTGGGCCCGCTTAGATGGCTTTAGCCAATCCAAATCGTGTTTAGC-3′) and was cloned into the pGAPZαA vector at the EcoRI/NotI sites, resulting in shuttle plasmid pGAPZαA/ProchymCB. The encoded prochymosin protein carries an N-terminal α-factor signal peptide for secretion in yeast culture.

2.3. Transformation of P. pastoris cells and producer strain preparation

The pGAPZαA/ProchymCB vector was linearized with endonuclease PagI (Thermo Fisher, USA) in Buffer O (Thermo Fisher, USA). The linearized vector was purified by phenol/chloroform extraction followed by ethanol precipitation. P. pastoris GS115 cells were electroporated with the linearized vector as follows. Fresh competent P. pastoris cells were prepared and transformed according to the instructions included with the EasyselectTM Pichia Expression Kit manual (Invitrogen). To 80 μL of the cells, 3 μg of purified linearized plasmid DNA was added, and then the suspension was pulsed in a 0.2 cm electroporation cuvette at 2 kV for 4.8 ms with electroporator (MicroPulser™, Bio-Rad, USA). Clones of P. pastoris GS115/pGAPZαA/ProchymCB were selected on an agar plate with zeocin (200 μg/mL). After that, the clones were screened for the presence of the insert by PCR with primers GAPFw (5′-GTCCTTATTCAATCAATTGACA-3′) and AO1rv (5′-GCAATGGCCATTCGTGACATCC-3′). The clones positive for pGAPZαA/ProchymCB insert were analyzed for chymosin expression by a milk-clotting assay. Clone with the highest

Figure 1. A - Domain structure of kappa-casein of cow’s milk; B - Comparative sequence of chymosin sensitive region of kappa-casein molecules of some mammals.
chymosin expression was preserved with glycerol at −80 °C and served as a producer strain.

2.4. Determination of stability of the gene integration in the genome of P. pastoris

Determination of the stability of the gene integration in the yeast genome was performed accordingly [42], the strain was cultivated on the YPD medium without any antibiotic for 20 generations. Colonies were then plated to YPD agar containing zeocin and grown for 3 days. The viable colonies on each plate were then counted. The genomic DNA of the transformants was analyzed by PCR to confirm integration of the cassette in the genome of P. pastoris.

2.5. Determination of an optimal medium for prochymosin expression

Cultivation of the Pichia pastoris GS115/pGAPZαA/ProchymCB strain was evaluated in eight media (Table 1). The cultivation was carried out according to the EasySelect™ Pichia pastoris Kit (Invitrogen, USA) recommendations in 250 mL flasks with 50 mL of a medium at 28 °C and 250 rpm for 72 h. Then, 1 mL of the culture was collected every 24 h and centrifuged at 13200 × g, for 20 min, at 4 °C (Eppendorf Centrifuge 5415R, Hamburg, Germany). The supernatants were stored at 4 °C and were subjected to the milk-clotting assay.

2.6. Enzyme production in shake flasks

Pichia pastoris GS115/pGAPZαA/ProchymCB strain cells were inoculated into a 50 mL flask containing 5 mL of YPD with zeocin (100 μg/mL) and were cultured overnight at 30 °C and 250 rpm in a shaking incubator (KS 4000 i control, IKA, Germany). The overnight culture was inoculated into 50 mL of YPD in a 500 mL shake flask and was incubated at 30 °C and 250 rpm for overnight. The overnight culture was inoculated into 500 mL of YPD + CAS + BMz (10g/L) in a 5 L shake flask and was incubated in a shaking incubator (Climo-Shaker ISF1-X, Kuhner, Switzerland) at 28 °C and 250 rpm for 120 h with daily addition of 40% glucose (0.8% final) and citrate-phosphate buffer pH 4.0 (100 mM final). The cells were collected by centrifugation at 3500 × g, for 20 min, at 4 °C (Avanti J-26XP with rotor JA-10, Beckman Coulter, USA) and discarded. The supernatant was used for enzyme purification.

2.7. Deglycosylation

Deglycosylation was performed accordingly [43]. The Pichia pastoris GS115/pGAPZαA/ProchymCB culture of was grown in 200 mL of the YPD + CAS + BMz (10g/L) medium in a flask for 144 h at 28 °C and 250 rpm. The cells were collected by centrifugation (3500 × g, 4 °C, 15 min) (Avanti J-26XP with rotor JA-10, Beckman Coulter, USA) and discarded. The supernatant was clarified by centrifugation at 4000 × g, for 1 h at 4 °C (Avanti J-26XP with rotor JA-20, Beckman Coulter, USA) and filtered through a 0.22 μm filter, and 20 mL of the filtrate was loaded onto 10 kDa MWCO (molecular-weight cutoff) protein concentrators (Thermo Fisher, USA). After concentration the chymosin (1 mg/mL) was denatured at 95 °C for 5 min and deglycosylated with endoglycosidase H (New England Biolabs, USA) at 37 °C for 16 h. The reaction was stopped by heating at 65 °C. The result was analyzed by western blotting.

2.8. Western blotting with an anti-prochymosin antibody

For western blotting, we applied polyclonal antibodies raised in rabbits against calf prochymosin. Western blotting was performed according to the standard protocol [44]. Briefly, protein samples were separated by SDS-PAGE in a 12% (w/v) gel and transferred to a polyvinylidene fluoride (PVDF) membrane. The membrane was stained with Ponceau S dye (0.1% (w/v) of Ponceau S in 5% (v/v) acetic acid) to confirm protein transfer. Membrane was blocked by 5% (w/v) skim milk powder in Tween 20/Tris-buffered saline (TBST: 50 mM Tris-HCl pH 7.6, 150 mM NaCl, 0.1% (w/v) Tween 20). The C. bactriana prochymosin protein was detected with the rabbit polyclonal antibody against calf prochymosin (with 1:5000 dilution) as a primary antibody and a goat anti-rabbit IgG antibody conjugated with horseradish peroxidase (Sigma-Aldrich Chimie S.a.r.l., Lyon, France) (with 1:10,000 dilution) as a secondary antibody. The bands were detected by the Enhanced Chemi-Luminescence Detection Kit (AppliedChem GmbH, Darmstadt, Germany), and an X-ray film was then exposed to the membrane (AgfaPhoto GmbH, Germany).

2.9. Purification of recombinant camel chymosin

Ion exchange chromatography was used to purify recombinant camel chymosin from yeast culture [45]. The supernatant of GS115/pGAPZαA/ProchymCB culture was passed through a 0.22 μm filter and pH was changed to 4.5 with 25 mM sodium acetate, followed by incubation at room temperature for 24 h with stirring to activate all produced chymosin. Activated culture supernatant was diluted 3 times with 25 mM sodium acetate, and pH was lowered to 3.0 with 1M HCl to make chymosin net charge positive. The mixture was loaded onto anion-exchange column, using peristaltic pump P1 (Pharmacia, Sweden), with DEAE-Sepharose FF (GE Healthcare, USA) equilibrated with 25 mM NaCl in 50 mM sodium citrate buffer (pH 3.0); this step was defined as clarification. The flow through the column was collected and loaded onto cation-exchange column, using peristaltic pump P1, with SP-Sepharose FF (Sigma-Aldrich, USA) equilibrated with 25 mM NaCl in 50 mM sodium citrate buffer (pH 3.0). The column was next washed with 5 mL of 25 mM NaCl in 50 mM sodium citrate buffer (pH 3.0) and 15 mL of 50 mM NaCl in 25 mM sodium acetate buffer (pH 5.5) to change the net charge of chymosin to negative. Then, the protein was eluted with 750 mM NaCl in 25 mM sodium acetate buffer (pH 5.5), and was collected by 1 mL fractions. The fractions with milk-clotting activity were combined, and the concentration of NaCl was lowered to 25 mM by dilution with 25 mM sodium acetate buffer (pH 5.5) and loaded, using peristaltic pump P1, onto the strong cation-exchange column with Q-Sepharose FF.
(Sigma-Aldrich, USA) equilibrated with 25 mM NaCl in 25 mM sodium acetate buffer (pH 5.5). The column was washed with 25 mM NaCl in 25 mM sodium acetate buffer (pH 5.5), and chymosin was eluted by means of a 50–2000 mM gradient of NaCl in 25 mM sodium acetate buffer (pH 5.5). The fractions were analyzed by the milk-clotting assay and the most active fractions were applied to SDS-PAGE. The SDS-PAGE was conducted according to the Laemmli method [46] in a Mini-PROTEAN Tetra cell (Bio-Rad Laboratories Inc., USA).

2.10. Determination of protein concentration

Protein concentration was determined by Bradford method [47] using Bio-Rad assay reagent (Bio-Rad Protein Assay Day, Munich, Germany) and bovine serum albumin as the standard. The measurements were carried out in three repetitions, with the average of the three repetitions being reported as the specified result.

2.11. The milk-clotting assay

The milk-clotting assay was carried out in accordance with ref. [48]. This assay was performed with powdered cow's skim milk reconstituted at 12% (w/v) in 0.025 M sodium acetate buffer, pH 6.0, as a substrate. The enzymatic reactions for clone selection were carried out at least in triplicate at 37 °C in test tubes with 20 μL of an enzyme solution and 1 mL of the substrate. The milk clots were visualized by turning the tubes upside down. One unit of milk-clotting activity was defined as the quantity of the enzyme required for clotting 1 mL of cow's skim milk in 40 min at 35 °C. Eq. (1) below shows how chymosin activity units (U) were calculated:

\[
A = \frac{V_{\text{milk}}}{V_{\text{chymosin}}} \times \frac{2400}{T_{\text{mc}}}
\]  

(Equation 1)

there \(V_{\text{milk}}\) is milk volume (mL), \(V_{\text{chymosin}}\) is volume of added chymosin (mL), and \(T_{\text{mc}}\) is milk-clotting time (sec).

The milk-clotting assay on mare's milk was carried out by using freeze-dried mare's milk (SaumalBioTech LLP, Kazakhstan). Powdered milk was reconstituted at 20% in deionized water, preheated to 40 °C, defatted by centrifugation at 3000 × g for 15 min (Eppendorf Centrifuge 5415R, Hamburg, Germany). Final composition of the substrate solution was brought to next: 10% mare milk, 0.2M buffer solution with an appropriate pH, 30 mM CaCl₂. Purified recombinant camel chymosin was diluted to 50 ng/μL for cow's milk and to 2 g/μL for mare's milk with 10 mM sodium acetate buffer (pH 5.5) and 10 μL of enzyme was incubated with 1 mL of the milk substrate at 37 °C. For cow's milk clotting time in each tube was measured by time first flakes observed by inversion of tube. For mare's milk reaction at each time point (15 s interval) was stopped by addition of pepstatin A in a final concentration in tube 50 μg. Stopped reaction was then centrifuged at 3500 × g for 3 min to divide precipitated casein. And time of the most transparent supernatant was recorded as clotting time.

2.12. Measurement of optimal substrate pH for the milk-clotting activity

To this end, skim milk powder was dissolved in either 50 mM sodium acetate buffer (pH 4.5–5.5), 50 mM imidazole buffer (pH 6.0–6.5), or 50 mM Tris-HCl buffer (pH 7.0–8.0). The milk solutions were placed as a substrate in 1.5 mL tubes. Maximum enzyme activity was defined as 100% activity, and the other samples were evaluated based on that. The tests were carried out in three repetitions, and the outcome was determined as the average of the three repetitions.

2.13. Determination of optimal temperature for the milk-clotting activity

The milk-clotting activity was measured over the temperature range of 0–70 °C (with 5 °C intervals) at 25 mM sodium acetate buffer (pH 6.0) for cow's milk and at 200 mM sodium acetate buffer (pH 5.0) for mare's milk. Substrate and enzyme solutions were preheated to the reaction temperature for 5 min, and the solutions were mixed. Maximum enzyme activity was defined as 100% activity, and the other samples were evaluated based on that. The tests were carried out in three repetitions, and the outcome was determined as the average of the three repetitions.

2.14. Effect of CaCl₂ on the milk-clotting activity

The effect of calcium on the milk-clotting activity was performed as described by [23]. Cow's milk-clotting activity were examined in an assay at 37 °C in the presence of CaCl₂ at a concentration from 0 to 160 mM, dissolved in 25 mM sodium acetate buffer (pH 6.0). Enzyme activity in the absence of metal ions was assumed to be 100%, and the other samples with varied metal ions were tested based on that. The tests were carried out in three repetitions, and the outcome was determined as the average of the three repetitions.

2.15. Effects of metal ions on the milk-clotting activity

The effects of various metal ions on the milk-clotting activity were examined in an assay at 37 °C in the presence of each of these 11 chlorides: CaCl₂, CoCl₂, NiCl₂, FeCl₂, BaCl₂, ZnCl₂, MgCl₂, MnCl₂, LiCl, KCl, and NaCl at a concentration of 10 mM, dissolved in 25 mM sodium acetate buffer (pH 6.0). Enzyme activity in the absence of metal ions was assumed to be 100%, and the other samples with varied metal ions were tested based on that. The tests were carried out in three repetitions, and the outcome was determined as the average of the three repetitions.

2.16. Enzyme production in a bioreactor

A 10 L fermenter (Biotstat, Sartorius, Germany) was utilized to determine the ability of the yeast strain to produce C. bactri anus recombinant chymosin on a large scale. A single colony was inoculated into 5 mL of YPD broth and grown at 28 °C and 250 rpm. After 24 h, the culture was transferred to 30 mL of YPD broth and grown in the same condition for 24 h. The culture was inoculated into 300 mL of the YPD + CAS + BMz (10g/L) medium and grown at 28 °C and 250 rpm for 24 h. Then, the culture was inoculated into 3 L of the YPD + CAS + BMz (10g/L) medium in the 10 L bioreactor. Standard procedures were utilized to operate the fermenter under the following cultivation conditions: 28 °C, 400 rpm, aeration 2–4 L/min, pH 4.0, fed carbon sources: 3% glucose and 5% 100 mM citrate-phosphate buffer (pH 4.0) (added each 24 h), and cultivation duration: 144 h.

Using Eqs. (2) and (3), the biomass and enzyme production rates (ΔBiomass max and ΔEnzyme max) were computed as the ratio of the increase in wet cells or activity to the cultivation period during which the rise in biomass/activity occurred.

\[
\text{Biomass}_{\text{max}} = \frac{\Delta \text{WCCW}}{T}
\]  

(Equation 2)

\[
\text{ΔEnzyme}_{\text{max}} = \frac{\Delta \text{EC}}{T}
\]  

(Equation 3)

where ΔWCCW is weight of produced wet cells (g) in period of time, ΔEC as enzyme activity of produced cells (U) in period of time, T is cultivation period (h). ΔWCCW was calculated as a difference of wet cells weight before and after cultivation period. ΔEC was calculated as an activity unit yield difference before and after cultivation period.

2.17. Software and statistical analysis

All tests were carried out in triplicate. In the quantitative study, the mean values and standard deviations were obtained using the GraphPad.
3. Results and discussion

The prochymosin gene of *C. bactrianus* was inserted into a yeast expression vector pGAPZuA under the control of the constitutive GAP promoter to construct plasmid pGAPZuA/ProchymCB. Twenty transformants were checked by PCR for the presence of the chymosin gene in genomic DNA. Ten transformants with the most intense glow of the PCR product under UV light were chosen and grown in YPD broth. The supernatants were analyzed by the milk-clotting assay, the results are indicated in Figure 2, milk-clotting activity units (10g/L) broths. The supernatants were analyzed by the milk-clotting assay, the results are indicated in Figure 2, milk-clotting activity units were calculated using Eq. (1). Six clones expressed chymosin with a good milk-clotting activity (Figure 2, Supplementary materials: Figure S1). Clone N8 showed maximum milk-clotting activity (1379 U/mL) and was chosen as a producer strain of camel chymosin.

A basal salt medium (BSM) proposed through Invitrogen is usually hired to acquire the excessive-density cultures, however it isn’t always the proper complement because of its excessive ionic strength, medium additives precipitation, and imbalanced nutrient concentration [49]. The media is troublesome to organize for big scale uses and needs the addition of vitamin H that is pricey vitamin, and of mixture of micronutrients, media is troublesome to organize for big scale uses and needs the addition of vitamin H that is pricey vitamin, and of mixture of micronutrients, several of which can inhibit recombinant protein production [50]. Well-known medium for the cultivation of *Pichia pastoris* yeast is YPD, containing peptone and yeast extract as a nitrogen source and glucose is added as a carbon source. The addition of beet molasses to YPD at a concentration of 0.5% and 1% increases the level of chymosin expression by 2.6 and 11.3%, respectively (Table 2). A further increase in the molasses content up to 2% strongly increases the expression level of the recombinant chymosin. The absence of glucose in the medium also had a negative effect, which is explained by the very low content of glucose in molasses - about 0.28 % of dry matter [51], which is needed to activate the *P. pastoris* GAP promoter [52]. Biotin is the sole vitamin in the commonly used rich growth medium for *P. pastoris* [53] to increase growth rate and biomass yield [54], but other complex vitamin sources may N-glycosylate proteins [60]. The most popular and significant form of post-translational modification is N-glycosylation [61]. Glycosylation of various chymosins expressed in yeast has been observed for: buffalo (*Bubalus arnee bubalis*) [62], camel (*Camelus dromedaries*) [33], cow (*Bos Taurus*) [50], goat (*Capra hircus*) [63], yak (*Bos grunniens*) [19] and this is not unusual for yeast-produced proteins [62].

The calculated molecular mass of *C. bactrianus* unglycosylated chymosin is 35.6 kDa. It follows from Figure 3 that the N-glycosylation increases the molecular mass by approximately 7 kDa. Using a mannose oligosaccharide linked to asparagine by two N-acetylglucosamines, *P. pastoris* may N-glycosylate proteins [60]. The most popular and significant form of post-translational modification is N-glycosylation [61]. Glycosylation of various chymosins expressed in yeast has been observed for: buffalo (*Bubalus arnee bubalis*) [62], camel (*Camelus dromedaries*) [33], cow (*Bos Taurus*) [50], goat (*Capra hircus*) [63], yak (*Bos grunniens*) [19] and this is not unusual for yeast-produced proteins [62].

The combination of anion exchange and cation exchange chromatography made it possible to purify chymosin from the medium and to concentrate the recombinant protein. By calculation, was established that bacterial camels’ mature chymosin has isoelectric point = 4.87 and was found to not bind to DEAE-Sepharose and binds to SP-Sepharose at pH 3.0 and to Q-Sepharose at pH 5.5 (Figure 4, Supplementary materials: Figure S3). As shown in Figure 4A recombinant chymosin purified on SP-Sepharose column migrated in SDS-PAGE as 2 protein bands (with molecular weights of 42 and 35 kDa). This reflects the glycosylation pattern. As we see in Figure 4B Q-Sepharose allowed to concentrate the chymosin and to get rid of other proteins.

Optimal pH for the activity of highest-purity *C. bactrianus* recombinant chymosin was pH 4.5 and 4.0 for cow’s and mare’s milk, respectively (Figure 5). With rising pH, the relative clotting activity decreased. At pH
The milk-clotting activity of the camel recombinant chymosin depending on the growth medium.

| Medium                  | Milk-clotting activity, U/mL |
|-------------------------|------------------------------|
| YPD                     | 1043 ± 18                    |
| YPD + BMz(5g/L)         | 1071 ± 33                    |
| YPD + BMz(10g/L)        | 1176 ± 37                    |
| YPD + BMz(20g/L)        | -                            |
| YP + BMz(10g/L)         | -                            |
| YPD + CAS + BMz(10g/L)  | 1411 ± 45                    |
| YPD + CAS + CrMz(10g/L) | 1212 ± 27                    |
| D + CAS                 | -                            |

Table 2.

Figure 3. Western blotting analysis of hourly accumulation of the camel recombinant chymosin in culture media, and analysis of the culture supernatant of strain GS115/pGAPZuA/ProchymCB before (lane 8) and after deglycosylation (lane 9). 1: bovine chymosin; 2: negative control; 3: 24 h; 4: 48 h; 5: 72 h; 6: 96 h; 7: 120 h; 8: 144 h; 9: 144 h supernatant treated with Endo H.

Figure 4. Purification of the camel recombinant chymosin on SP-Sepharose FF (A) and Q-Sepharose FF (B).
Protein production was started from the first 24 h of cultivation and reached 375 U/ml. Total yield of the recombinant enzyme was 80 mg per liter of yeast culture. The resulting supernatant at 144 h cultivation contained 1412 U/mL activity.

According to our calculations, using Eqs. (2) and (3), the maximum of biomass production rate of 4.29 g/h is observed on the third day. Maximum of enzyme production rate of 89,075 U/h was at the fourth day. After 96 h of growth, no further growth in chymosin production was observed, which coincided with the beginning of the stationary phase of growth. In compare, *C. dromedarius* and Yak chymosins with methanol-induced expression in *P. pastoris* under AOX1 promoter, maximum enzyme yield was observed at 144 h and 184 h respectively [33, 67].

Table 3. Effects of metal ions on the milk-clotting activity of the camel recombinant chymosin.

| Metal ion Concentration, mM | Relative activity, % (±s.t.d.) |
|-----------------------------|-------------------------------|
| Control                     | 100 ± 3.95                    |
| Ca²⁺ (CaCl₂) 10 mM          | 298.8 ± 16.04                 |
| Cr²⁺ (CrCl₂) 10 mM          | 182.7 ± 8.21                  |
| Ni²⁺ (NiCl₂) 10 mM          | 19.9 ± 0.39                   |
| Fe²⁺ (FeCl₂) 10 mM          | 338.7 ± 12.99                 |
| Ba²⁺ (BaCl₂) 10 mM          | 373.5 ± 20.06                 |
| Zn²⁺ (ZnCl₂) 10 mM          | 98.5 ± 4.54                   |
| Mg²⁺ (MgCl₂) 10 mM          | 188.2 ± 5.31                  |
| Mn²⁺ (MnCl₂) 10 mM          | 518.4 ± 19.02                 |
| Li⁺ (LiCl) 10 mM            | 162.8 ± 2.73                  |
| K⁺ (KCl) 10 mM              | 97 ± 4.61                     |
| Na⁺ (NaCl) 10 mM            | 105 ± 4.06                    |

Figure 5. Optimal substrate pH for the milk-clotting activity of the camel recombinant chymosin.

Figure 6. Optimal temperature for the milk-clotting activity of the camel recombinant chymosin.

Figure 7. The impact of CaCl₂ concentration on the cow’s milk-clotting activity of the camel recombinant chymosin.

Figure 8. Pilot scale production of *C. bactrianus* recombinant chymosin by yeast *P. pastoris*.
Table 4. Biomass and enzyme production rate.

| Cultivation period, h | Volume, mL | Wet cells weight, g | Activity, U | ΔBiomassmax, g/h | ΔEnzymemax, U/h |
|----------------------|------------|---------------------|-------------|------------------|-----------------|
| 0–24                 | 3000       | 69                  | 1 125 000   | 2,88             | 46 875          |
| 24–48                | 3400       | 170                 | 2 567 000   | 4,21             | 60 083          |
| 48–72                | 3900       | 273                 | 4 067 700   | 4,29             | 62 529          |
| 72–96                | 4500       | 345                 | 6 205 500   | 3,00             | 89 075          |
| 96–120               | 5200       | 377                 | 7 342 400   | 1,33             | 47 371          |
| 120–144              | 6000       | 382                 | 8 472 000   | 0,21             | 47 067          |

These results indicate that C. bactrianus chymosin expression in P. pastoris under the control of the constitutive (GAP) promoter is effective. C. bactrianus chymosin was tested in mare's milk coagulation. It is known that mare's milk differs significantly from that of other dairy mammals [68]. The whey protein fraction represents almost 40% in mare's milk and less than 20% in cow's milk. Mare's milk could be defined typically as albuminous milk. Due to this, there is a big problem in processing and clot formation of mare's milk. Previous investigation on rennet-induced coagulation of mare's milk reported that it does not form a gel during renneting, by bovine chymosin and is not suitable for cheese production [26].

Future studies will focus on the kinetic and thermodynamic parameters of the C. bactrianus chymosin, as it was shown for other enzymes [16, 69, 70], optimal parameters for large-scale and industrial fermentation of GS115/pGAPZaA/ProchymCB strain will be found out and cheese making conditions from mare's and camel's milk using C. bactrianus chymosin will be assayed.

4. Conclusions

In this study, a recombinant chymosin from the two-humped camel C. bactrianus was characterized for the first time. The C. bactrianus chymosin gene was cloned and expressed in P. pastoris under the control of the GAP promoter and purified from culture via a combination of cation and anion exchange chromatography. C. bactrianus recombinant chymosin manifested high milk-clotting activity (9605 U/mg) with optimum conditions for cow's milk-clotting activity at 45 °C, pH 4.5, and 30 mM CaCl2 and for clotting mare's milk at 70 °C, pH 4.0. The dependence of this activity on metal ions was investigated too. The activity of cow's milk clotting is increased by Mn2⁺, Fe3⁺, Ba²⁺, Co²⁺, and Mg2⁺ and strongly suppressed by Ni2⁺ and no effect on mare's milk clotting was observed. Recombinant-chymosin production in a 10 L fermenter indicates that the camel chymosin constitutive expression P. pastoris yeasts could be used for large scale production for milk-clotting enzyme manufacturing with 80 mg/L of enzyme yield. The proposed expression system has high efficiency of the production of C. bactrianus recombinant chymosin. The inclusion of the constitutive GAP promoter to control the heterologous expression may be a good solution for the production of a recombinant enzyme for the food industry [71]. Our research on the optimal medium composition allowed us to find out conditions for the recombinant-chymosin manufacture under the control of this constitutive promoter. C. bactrianus recombinant chymosin was used to prepare cheese from bovine milk; this confirms the applicability of the new enzyme to a cheese-making process. C. bactrianus recombinant chymosin appears to be a good alternative to the milk-clotting enzymes currently used in the cheese industry. Milk-clotting activity on a mare's milk has the prospect for processing mare's and donkey's milk.

Declarations

Author contribution statement

Zhibger Akinchev: Conceived and designed the experiments; performed the experiments; analyzed and interpreted the data; wrote the paper.
