Proline iminopeptidase Pepl overexpressing Lactobacillus casei as an adjunct starter in Edam cheese

Sahar Navidghasemizad, Timo M Takala, Tapani Alatossava, and Per EJ Saris*

Department of Food and Environmental Sciences, University of Helsinki, Helsinki, Finland

Keywords: Edam cheese, bitterness, Pepl, proline

Introduction

Bitterness is a common taste defect in semi-hard cheeses, affecting their acceptance and marketing. One reason for bitter taste in cheese is an incomplete hydrolysis of hydrophobic peptides produced during cheese ripening. The major milk protein, casein, and in particular its subunits, α- and β-casein, are an abundant source of proline. Primary degradation of casein by rennet, plasmin and starter proteinases results in the production of hydrophobic oligopeptides rich in proline. Proline as a free amino acid may have a sweet taste, but the accumulation of proline-rich oligopeptides rich in proline. For instance, cheeses made with adjunct cultures of L. casei, L. paracasei, Lb. plantarum, or Lb. helveticus were less bitter than those made without adjuncts. The efficiency of lactobacilli to reduce bitterness in cheese may be further increased by the overexpression of heterologous peptidase. So far, various peptidases have been overexpressed in L. lactis, and studied in cheese models. Courtin et al. cloned genes encoding six different peptidases from Lb. helveticus into L. lactis. The authors found increased levels of PepX, PepQ, and PepW, which led to a 3-fold increase in free amino acids in the model cheese. Similarly, Guldfeldt et al. overexpressed PepC, PepN, PepO, and PepV from different strains of Lactococcus in L. lactis subsp. cremoris. In other studies, proline-specific peptidases have been overexpressed in Lc. lactis. However, there are no studies regarding the overexpression of peptidases in Lactobacillus. Lactobacilli show higher potential as hosts for the expression of peptidases since they have a better protease producing potential and they survive and increase in numbers in ripened cheese.

In this study the growth of genetically modified Lactobacillus casei LAB6, overexpressing proline iminopeptidase Pepl and its capacity to increase free proline was investigated during ripening of Edam cheese. The strain successfully survived 12 weeks of ripening period in cheese. The food-grade plasmid pLBB604, carrying the pepI gene, was stable and Pepl enzyme was active in LAB6 cells isolated at different stages of the ripening process. However, HPLC analyses indicated that Lb. casei LAB6 could not increase the amount of free proline in ripened cheese.

*Correspondence to: Per EJ Saris; Email: per.saris@helsinki.fi
Submitted: 05/17/2013; Revised: 06/20/2013; Accepted: 06/25/2013
http://dx.doi.org/10.4161/bio.25543

Bioengineered 4:6, 408–412; November/December 2013; © 2013 Landes Bioscience

Volume 4 Issue 6

408
overexpressed constructed PepI. In this study, the LAB6 strain was used as an adjunct starter for Edam cheese production. The objectives were to examine: (1) the growth trend of Lb. casei strain LAB6, (2) the stability of the plasmid pLEB604 in carrying the pep gene in the strain LAB6 during ripening period, and finally (3) free proline content in the final ripened Edam cheese.

## Results and Discussion

### Identification of Lb. casei LAB5 and LAB6

The growth of Lb. casei LAB5 and Lb. casei LAB6 in cheese and their plasmid stability were analyzed during the ripening process. Strains were isolated from cheese at different steps of ripening and identified by comparing the RAPD profiles of LAB5 and LAB6 to other casei-type lactobacilli in cheeses using primer 6789 (Fig. 1). The RAPD profiles of LAB5, LAB6, and their host Lb. casei E were identical to each other, but were different from other Lb. rhamnosus and Lb. casei strains. Therefore, it was possible to identify and track the plasmid stability of Lb. casei LAB 5 and LAB 6 strains during the ripening process.

### Survival of starters and Lb. casei LAB5 and LAB6 adjuncts in cheese

During the whole ripening period, according to RAPD profile of isolated colonies, only Lb. casei LAB5 or LAB6 were found on LAB5 plates. All the isolated LAB5 and LAB6 colonies showed lactose positive phenotype on lactose indicator plates, verifying the presence of the lactose-selectable food-grade plasmids in the strains. If the strains had lost the plasmids, they would have also lost the ability to metabolize lactose. All colonies from LAB6, but not from LAB5, showed PepI activity (data not shown). These tests confirmed that during the whole ripening period, both pLEB604 and pLEB604 plasmids were stable and functional in LAB5 and LAB6, respectively, during the ripening period.

The growth of starters LAB5 and LAB6 during ripening period is shown in Figure 2. Both strains LAB5 and LAB6 survived the whole 12 weeks ripening period, and their growth trend was similar in both batches. Strains started to grow from approximately 10^5 cfu g^-1 and by three weeks the bacterial counts stabilized to approximately 10^7–10^8 cfu g^-1. The starters Lactococcus and Leuconostoc number remained constant throughout the ripening period in the first batch, whereas in the second batch, their number started to decrease after 28 d and they were not detectable after 10 and 12 weeks in cheeses made using LAB6 or LAB5 strains, respectively. In the research by Courtin et al.,^23^ it was shown that the number of lactococci in cheeses was not detectable after 28 d of ripening. This is in parallel with the fact that long ripening time results in lactose depletion in the cheese. The lack of energy source for lactococci and starter cultures cause cells to lyse. Therefore, in this study, for the first batch, the starter counts, for an unknown reason, did not decrease even after 84 d. Cheese batches were made on two different days. The variation in quality of milk used for the cheese making process such as milk composition, and its lactose content as well as the original microflora can affect the starter number in final product.

The iminopeptidase activity was higher in cheeses made with PepI-producing strain LAB6 than those of cheeses containing the vector strain LAB5 (Fig. 3). This indicates that LAB6 increased PepI activity in cheese, thus the strain may promote proline release from peptides.

### Amino acid profile of Edam cheeses made with Lb. casei LAB5 and LAB6 adjuncts

The amino acid content of cheeses was determined qualitatively and quantitatively using HPLC. The amounts of free amino acid in both cheeses containing LAB5 and LAB6 were approximately at the same level (Fig 4), meaning LAB6 could not increase the free proline content in the cheese. The PepI is an intracellular enzyme, and its release to cheese matrix requires the lysis of LAB6 cells. If the cells are not lysed extensively, then PepI cannot be effectively released to hydrolyze the substrates. A longer ripening period of the cheese may promote the lysis of LAB6 cells and increase the peptide hydrolysis. Low temperature may also decrease enzyme activity. Edam cheeses in this study were ripened at temperatures of about 12 °C. Considering the fact that proline iminopeptidase has an optimum activity at 37 °C, lower temperature during ripening can significantly slow down the enzyme activity.26,27 Furthermore, the low amounts of substrate could contribute to the lack of change in the proline profile of the cheese. Proline specific enzymes i.e., PepQ, PepP, PepX, and PepP enzymes isolated and identified from different lactic acid bacteria act very specifically and cleave proline only from a certain position in a polypeptide chain which is specific for that individual enzyme (Table 1). Therefore, in order to release proline from cheese substrate in an efficient way, a combination of different proline-specific enzymes would be required. In future experiments, the
LAB6 strain, which showed good potential to grow and produce peptidase activity in cheese, could be engineered to produce several proline-specific enzymes in combination with a mutation weakening the cell wall for increased lysis during ripening of the cheese.

Materials and Methods

Bacterial strains and growth conditions. LAB5 and LAB6—variants of *Lactobacillus casei* E, a NSLAB strain isolated from a high quality Edam cheese—were used as adjunct starter cultures. LAB5 carries the plasmid pLEB600, a lactose-selectable food-grade vector. LAB6 is a PepI expressing strain carrying the plasmid pLEB604, in which a gene encoding the PepI enzyme is cloned under the control of a constitutive promoter in pLEB600. For cheese making, starter cultures *Lactococcus lactis* (Chr. Hansen) and *Lactobacillus casei* were added as freeze-dried cultures. For colony counting, the starters were grown in M17 agar plates (Oxoid) containing 0.5% glucose. Strains *Lactobacillus casei* 163 (Danisco), *Lactobacillus casei* defensis (Danisco), *Lactobacillus rhamnosus* LC-705 (Valio Ltd), *Lactobacillus rhamnosus* LCAB1, and *Lactobacillus rhamnosus* 1/6 (Valio Ltd) were used for RAPD (random amplification of polymorphic DNA) profile comparisons. Lactobacilli other than LAB5 and LAB6 strains were grown in MRS (Oxoid). LAB5 and LAB6 were grown in modified MRS supplemented with 1% lactose at 37 °C. Modified MRS lacks glucose and beef extract. In order to follow the plasmid stability during the cheese ripening, lactose utilization was determined from the *Lactobacillus casei* strains. Modified MRS plates containing 1% lactose and 0.05 mg ml⁻¹ bromocresol purple were used to determine the lactose utilization of LAB5 and LAB6 colonies isolated from cheese. The lactose indicator plates were incubated for 48 h at 37 °C. A color change from purple to yellow around lactose-positive colonies was an indication for acid production by the strains.

Cheese manufacture and ripening. Edam cheese was prepared in two batches. For each batch, cheeses were made using two different strains as adjunct starter: (1) control cheese containing *Lactobacillus casei* LAB5 with no PepI expressing enzyme, (2) the cheese prepared with *Lactobacillus casei* LAB6, containing PepI expressing enzyme. About 120 L of milk was used to produce 9.6 kg of cheese. The cheeses were processed essentially according to the method used to produce Edam cheese by Valio Ltd. In brief, milk was pasteurized for 15 sec at 75 °C, then it was cooled down to 32 °C and starters *Lactococcus lactis* and *Leuconostoc mesenteroides* (Chr. Hansen Ltd), as well as adjunct *Lactobacillus casei* LAB5 or LAB6 were added to milk. Adjuncts were added approximately 3 x 10⁸ mL⁻¹ milk, representing about 10% of the amount of starters. After 30 min, rennet was added and coagulation took place for 35 min. Then, the curd was cut, and stirred for 11 min. After that, 30 L of whey was discarded and stirring continued for 16 min at 31 °C. Next, 10 L water was added to curd and temperature was raised up to 36 °C and stirring continued for 60 min. Thereafter, curd was ready for pressing. Molding process was performed for 3 h at room temperature. After that, curds were salted in 16% brine at 12 °C for 8–10 h and then vacuum packaged and ripened at 12 °C for 84 d. Final packages contained about 800 g cheese which were stored at 12 °C. To monitor the ripening, samples were taken every second week.

Identification of *Lactobacillus casei* LAB5 and LAB6 in cheese samples. For isolation and identification of *Lactobacillus casei* strains LAB5 and LAB6, 10 g of cheese was homogenized in a Stomacher (Lab Blender 400, Seward Medical) with 90 ml of 0.9% NaCl. Serial dilutions of homogenates were plated on LBS agar (BBL, Becton Dickinson), and the plates were incubated anaerobically (AnaeroChamber® A, Merck) in jars at 37 °C for 48 h, after which the colonies were counted. After isolation, LAB5 and LAB6 were identified by RAPD technique. Three different...
RAPD primers (kindly provided by Valio Ltd) were tested for identification of LAB5 and LAB6 from other closely related species and strains. The most suitable primer, named 6789, was chosen for RAPD-PCR identifications. The chosen RAPD-PCR primer sequence was: 5′-GCT CGT ATG TTG TGT GG-3′. To identify the lactobacilli, 100 random LBS colonies per cheese sample were chosen for colony RAPD-PCR. PCR amplification was directly performed on fresh isolated cells grown on LBS agar plates. RAPD analyses were performed in a 50 μL reaction volume consisting of 5 μL 10× PCR reaction buffer containing 15 mM MgCl₂ (Finnzymes), 1 μL dNTP (10 mM; Finnzymes), 1 μL of the primer (20 μM), 0.5 μL DNA polymerase (DyNAzyme II DNA polymerase 2 U μL⁻¹, Finnzymes) and 42.5 μL of deionized sterile water. RAPD-PCR amplification was performed in an Eppendorf Master Cycler gradient apparatus (Eppendorf AG). The temperature profile in thermocycler was as follows: initial denaturation at 94 °C for 2 min, then 35 cycles of 95 °C for 20 sec, 37 °C for 30 sec, 72 °C for 2 min, and final extension in 72 °C for 10 min. Amplification products were analyzed electro- phoretically in 1% (w/v) agarose gels containing ethidium bromide (0.5 μg mL⁻¹) and visualized under UV light.

PepI enzyme activity in Lb. casei LAB5 and LAB6 isolated from cheese.
Fifty colonies of LAB5 and LAB6 isolated from cheese samples were tested by a rapid method for the detection of PepI activity. Colonies from LBS plates were randomly picked and incubated overnight in MRS broth. Next day, the cultures were washed with 50 mM potassium phosphate buffer (pH 7). The pellets were suspended in 100 μL of the same buffer. Then, 25 μL of 20 mM l-proline-ρ-nitroanilide substrate (Sigma-Aldrich) was added, and the tubes were incubated at 37 °C for 30 min. PepI producing strains degrade the substrate and release yellow-colored nitroaniline, which was measured spectrophotometrically at 410 nm.

Detection of PepI activity from cheese.
Fifty milligrams of grated cheeses were mixed with 1.5 mL 50 mM potassium phosphate buffer (pH 7). The mixtures were centrifuged for 10 min at 16,000 g. The supernatants were filtered and measured spectrophotometrically at 410 nm.

Qualitative and quantitative determination of free amino acids in the ripened cheeses.
Amino acid analyses were performed from full ripened cheeses (84 d). The free amino acid composition was determined as phenylthiocarbamate derivatives using the Waters Pico Tag method and the preparation of the derivatives and the HPLC analysis were performed according to the instructions of Millipore Corporation by external service (MTT Agrifood Research Center).

Disclosure of Potential Conflicts of Interest
No potential conflict of interest was disclosed.

Acknowledgments
We want to thank Mr Jyri Rekonen at Viikki dairy pilot plant for the manufacture of the test cheeses.
