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Published in: Journal of Bacteriology

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Document Version Publisher's PDF, also known as Version of record

Publication date: 1998

Link to publication in University of Groningen/UMCG research database

Citation for published version (APA): Buist, G., Venema, G., & Kok, J. (1998). Autolysis of Lactococcus lactis is influenced by proteolysis. *Journal of Bacteriology*, *180*(22), 5947 - 5953.

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Autolysis of Lactococcus lactis Is Influenced by Proteolysis

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Received 20 April 1998/Accepted 21 August 1998

The autolysin AcmA of Lactococcus lactis was shown to be degraded by the extracellular lactococcal proteinase PrtP. Autolysis, as evidenced by reduction in optical density of a stationary-phase culture and concomitant release of intracellular proteins, was greatly reduced when L. lactis MG1363 cells expressed the cell wallanchored lactococcal proteinase PrtP of the PI-type caseinolytic specificity (PI). On the other hand, lactococcal strains that did not produce the proteinase showed a high level of autolysis, which was also observed when the cells produced the secreted form of PI or a cell wall-anchored proteinase with PIII-type specificity. Autolysis was also increased when MG1363 expressed the cell wall-anchored hybrid PI/PIII-type proteinase PIac. Zymographic analysis of AcmA activity during stationary phase showed that AcmA was quickly degraded by PI and much more slowly by PrtP proteinases with PIII-type and intermediate specificities. Autolysis of L. lactis by AcmA was influenced by the specificity, amount, and location of the lactococcal proteinase. No autolysis was observed when the various proteinases were expressed in an L. lactis acmA deletion mutant, indicating that PrtP itself did not cause lysis of cells. The chain length of a strain was significantly shortened when the strain expressed a cell wall-anchored active proteinase.

Lactococcus lactis, like many other lactic acid bacteria, is a multiple-amino-acid auxotroph (6). For growth in milk, in which the concentrations of free amino acids and peptides are very low, these bacteria depend on an active proteolytic system which allows the degradation and release of amino acids from milk proteins (α_{S1} -, β -, and κ -caseins). The initial step in degradation is carried out by the extracellular cell wall-bound proteinase PrtP. The extreme C terminus of this protein contains the typical LPXTG motif of gram-positive cell surface proteins. After cleavage between the threonine and glycine residues, such proteins are covalently bound to the cross bridge in the peptidoglycan (23). The genes encoding PrtP have been cloned from several L. lactis strains and sequenced. Although the deduced amino acid sequences are over 98% identical, the proteolytic specificity of the proteinases toward milk caseins can be quite different. The lactococcal proteinases were initially divided into two major classes on the basis of the caseinolytic specificity: PIII-type proteinases (PIII) degrade α_{S1} -, β -, and κ -caseins, while the PI-type proteinases (PI) mainly degrade β -casein but with a specificity different from that of the PIII enzymes (30). Regions in the proteins determining cleavage specificity were identified by reciprocal exchange of fragments of the genes for PI of L. lactis Wg2 and PIII of strain SK11, and enzymes with new proteolytic properties were obtained (31). The proteinases of 16 different L. lactis strains have recently been classified into seven groups by examining their degradation of fragment f1-23 of α_{S1} -casein (9).

L. lactis expresses one major autolysin, AcmA, which is responsible for cell separation and autolysis during the stationary phase of growth (4, 5). In a previous study on AcmA, we observed that the C-terminal repeats of AcmA are removed by an unknown proteolytic activity (3), without affecting the enzyme activity.

Several observations indicate that processes in which auto-lysins take part are influenced by proteolysis. Competence, cell separation, and motility in *Bacillus subtilis* all involve cell wall hydrolase activities and were shown to be sensitive to extracel-lular proteinases (1, 35). A multiple proteinase-deficient strain of *B. subtilis* had a higher rate of turnover of pentidoglycan of *B. subtilis* had a higher rate of turnover of peptidoglycan a than proteinase-proficient strains (12). Growth of the protein-ase-deficient strain in the presence of subtilisin resulted in reduced peptidoglycan turnover and filament formation. Pro-teinase-hyperproducing strains showed an even more diminteinase-hyperproducing strains showed an even more diminished turnover and formed filaments. Addition of a proteinase inhibitor to cultures of these strains resulted in an increase in peptidoglycan turnover. Coxon et al. (7) showed that protease deficiency of *B. subtilis* was associated with an increased ten-dency of cells to lyse as they approached stationary phase. Salt-induced lysis of exponential phase cells of *Staphylococcus* aureus was inhibited by a brief pretreatment with subtilisin. Analysis of cell wall-lytic enzymes by renaturing sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) revealed that all lytic bands had disappeared after incubation with the proteinase (33). Loss of cell wall-lytic activity was also observed upon addition of protease to exponentially growing cells of Staphylococcus haemolyticus, which resulted in the formation of multicells (34). Lysis of Streptococcus pneumoniae induced by the addition of D-cycloserine or benzylpenicillin is inhibited by addition of trypsin to the growth medium (20). On the other hand, proteolytic activities have also been shown to be enhancers of cell wall-lytic activities. A muramidase of Enterococcus hirae is extracellular proteolytically activated from a 130-kDa latent form to a 87-kDa active form (13). The autolysin ATL of S. aureus is expressed as a bifunctional protein with amidase and glucosaminidase domains which is processed by an extracellular protease to generate two peptidoglycan hydrolases (17).

Haandrikman et al. (11) reported that the major secreted protein Usp45 of L. lactis MG1363 was absent in a culture of cells expressing the PI proteinase of strain Wg2, suggesting that this enzyme not only hydrolyzes caseins but is also able to degrade secreted proteins. In this paper, we show that PrtP degrades AcmA and that the rate and degree of autolysis of L.

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TABLE 1. Bacterial strains and plasmids used in this study

Strain or plasmid	Relevant phenotype(s) or genotype(s) ^a	Reference
Strains		
L. lactis subsp. cremoris		
MG1363	Plasmid-free strain	10
MG1363 <i>acmA</i> ∆1	Derivative of MG1363 carrying a 701-bp SacI/SpeI deletion in acmA	5
Plasmids		
pGK13	Em ^r Cm ^r , pWV01-based lacto- coccal cloning vector	14
pGKV500	Em ^r , pWV01 derivative specify- ing sPI of <i>L. lactis</i> Wg2	16
pGKV552	Em ^r , specifying aPI of <i>L. lactis</i> Wg2	11
pGKV1552	Em ^r , pGKV552 specifying aPI*	11
pGKV552abc	Em ^r , specifying aPlabc	31
pGKV552ac	Em ^r , specifying aPIac	31

^a Cm^r, chloramphenicol resistance; Em^r, erythromycin resistance.

lactis are dependent on the specificity, location, and amount of PrtP produced.

MATERIALS AND METHODS

Bacterial strains, plasmids, and growth conditions. The strains and plasmids used in this study are listed in Table 1. L. lactis was grown in M17 broth (Difco Laboratories, Detroit, Mich.) or whey-based medium (8) at 30°C as standing cultures. Twofold-diluted M17 (0.5× M17) agar plates contained 1.5% agar and 0.95% β-glycerolphosphate (Sigma Chemical Co., St. Louis, Mo.). In all cases, 0.5% glucose and 5 µg of erythromycin (Boehringer GmbH, Mannheim, Germany) per ml were added.

General DNA techniques and transformation. Plasmid DNA was isolated by using a QIAGEN plasmid DNA isolation kit and protocol (QIAGEN GmbH, Hilden, Germany) or by the alkaline lysis method as described by Sambrook et al. (25), with modifications suggested by Seegers et al. (27). Electrotransformation of L. lactis was performed with a Gene Pulser (Bio-Rad Laboratories, Richmond, Calif.) as described by Leenhouts and Venema (19). All chemicals were of analytical grade and were from Merck (Darmstadt, Germany) or BDH (Poole, United Kingdom).

SDS-PAGE and detection of AcmA. Five milliliters of whey cultures and 2 ml of M17 cultures were subjected to centrifugation; 5 ml of the whey supernatants and 0.5 ml of the M17 supernatants were dialyzed against several changes of demineralized water, lyophilized, and dissolved in 1 and 0.25 ml, respectively, of denaturation buffer (2). The cell pellets were washed with 1 ml of corresponding fresh medium and resuspended in 1 ml of denaturation buffer. Cell extracts were prepared as described by van de Guchte et al. (29).

AcmA activity was detected by a zymogram staining technique using SDS-12.5% polyacrylamide gels containing 0.2% autoclaved, lyophilized Micrococcus lysodeikticus ATCC 4698 cells (Sigma) as described before (5).

SDS-PAGE (12.5% polyacrylamide gel) was carried out according to Laemmli (18) with the Protean II Minigel system (Bio-Rad). The Bio-Rad prestained high-range molecular weight marker and the Amersham high-range SDS-PAGE Rainbow colored protein molecular weight marker (Amersham Life Science Inc., Buckinghamshire, United Kingdom) were used as protein size references. SDSpolyacrylamide gels were stained with Coomassie brilliant blue (Bio-Rad)

Optical density measurements and enzyme assays. Overnight cultures of L. lactis in M17 were diluted 100-fold in prewarmed M17, and the optical densities at 600 nm were followed over time in a Novaspec II spectrophotometer (Pharmacia Biotech AB, Uppsala, Sweden).

The presence of intracellular X-prolyl dipeptidyl aminopeptidase (PepX) in culture supernatants was measured by using the chromogenic substrate Ala-Prop-nitroanilide (Bachem Feinchemikalien AG, Bubendorf, Switzerland) as described before (3).

Proteinase activity was determined by using the chromogenic peptide MeO-Suc-Arg-Pro-Tyr-pNA (Chromogenix AB, Mölndal, Sweden) as described by Mierau et al. (22).

RESULTS

AcmA is degraded by PrtP. To examine the possible degradation of AcmA by the lactococcal proteinase PrtP, the PrtPnegative strain L. lactis MG1363(pGK13) [MG1363 (P⁻)] and two proteinase-producing strains were grown overnight in whey-based medium. *L. lactis* MG1363(pGKV552) expresses the PI-type PrtP (16). This full-length protein (cell wall-anchored PI [aPI]) is attached to the cell by a C-terminal cell wall anchor sequence (11). The proteinase produced by MG1363 (pGKV500) lacks this anchor and PI; consequently, it is secreted into the growth medium (and hence referred to as sPI [secreted PI]). As expected, proteinase activity was absent in the cell extract and supernatant of MG1363 (P⁻), while all sPI activity was present in the supernatant. Most of the proteinase activity of MG1363 expressing aPI [MG1363 (aPI)] fractionated with the cells (results not shown). These activity measurements were corroborated by SDS-PAGE (12% polyacrylamide gel). Analysis of cell extract and supernatant samples of the three strains showed that PrtP is present in high amounts in the supernatant of MG1363 (sPI), while a lower amount is seen in that of MG1363 (aPI). In the latter strain, most of the proteinase is detected in the cell extract (Fig. 1A). The major B secreted lactococcal protein Usp45 (28) is clearly present in the supernatant of MG1363 (P^-) but absent in these fractions



FIG. 1. (A) Protein profiles of cell (c) and supernatant (s) fractions of whey-grown L. lactis MG1363 containing pGK13 (P⁻), pGKV500 (sPI), or pGKV552 (aPI) in an SDS-12.5% polyacrylamide gel stained with Coomassie brilliant blue. PrtP, lactococcal proteinase; Usp45, secreted protein of unknown function (28). (B) Analysis of AcmA activity in the samples used for panel A by renaturing SDS-PAGE (12.5% polyacrylamide gel) in the presence of 0.15% M. lysodeikticus autoclaved cells. AcmA, N-acetylmuramidase of L. lactis; arrowhead, major degradation product of AcmA (5). Molecular masses (in kilodaltons) of standard proteins for both gels are shown on the left. Five microliters of each sample was loaded onto the gels.

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FIG. 2. (A) Growth and lysis by optical density (OD) measurement at 600 nm of cultures of *L. lactis* MG1363 containing pGK13 (P^- ; \times), pGKV500 (sPI; *), pGKV552 (aPI; \diamond), pGKV552 (aPI; \diamond), pGKV552abc (aPlac; Δ), or pGKV552ac (aPlac; \Box). *L. lactis* MG1363*acm*AΔ1 containing either of these plasmids exhibited very similar growth curves of which only one, MG1363*acm*A\Delta1(pGK13), is shown (\blacksquare). The strains were examined during 3 days of incubation at 30°C. Lettered arrowheads indicate the time points at which samples were taken from the cultures for analysis of AcmA activity (Fig. 3) and protein release (Fig. 2B and data not shown). (B) Analysis of PepX activity (in arbitrary units [A.U.]) in the culture supernatants of the strains presented in panel A. Symbols are as defined for panel A.

of the proteinase producers. When samples were run on an SDS-polyacrylamide gel containing *M. lysodeikticus* autoclaved cells, AcmA activity was detected in the cell extracts of all cultures (Fig. 1B). A band of cell wall hydrolytic activity corresponding to mature AcmA (40 kDa) is present in the supernatant of MG1363 (P⁻) but not in the supernatants of the sPI-and aPI-producing cultures. These results show that AcmA is subject to proteolytic degradation by PrtP.

Autolysis of various proteinase-producing strains of *L. lactis.* To investigate the influence of AcmA degradation on autolysis of *L. lactis*, the optical densities at 600 nm and release of the intracellular proteins were followed over time. The strains used in this experiment were MG1363 (AcmA⁺) and MG1363*acmA* $\Delta 1$ (AcmA⁻) harboring either pGK13 (negative control), pGKV500 (producing sPI), pGKV552 (specifying aPI), and pGKV1552 (encoding an inactive [Asp30 \rightarrow Asn30] form of PI, aPI^{*}). Plasmids pGKV552ac and pGKV552abc specify hybrid proteinases (aPIac and aPIabc, respectively) in which regions a and c and regions a, b, and c in PI have been replaced by the corresponding domains of the PIII-type proteinase of *L. lactis* SK11 (31). The hybrid proteinase specified by pGKV552abc has PIII-type specificity, whereas the hybrid



FIG. 3. Zymographic analysis of AcmA activity. Each lane contains 10 µl of the supernatant of an MG1363 culture in M17 producing the indicated proteinase sampled at the time points A, B, and C shown in Fig. 2A. Molecular masses (in kilodaltons) of standard proteins are shown on the left.

proteinase specified by pGKV552ac has a specificity intermediate between that of PI and PIII (31). As MG1363 (P⁻) grows to a lower optical density in whey than a Prt⁺ strain, M17 medium was used to obtain identical culture densities for all strains.

Overnight cultures of the strains were diluted 100-fold in fresh M17 medium, and the optical densities at 600 nm and release of the intracellular PepX of various strains were followed over time (Fig. 2). All strains grew equally well except for L. lactis MG1363 (aPI), which reached approximately the same maximum optical density but grew somewhat more slowly. Nearly no reduction in optical density during the stationary phase of growth was observed for any of the MG1363acmA $\Delta 1$ mutants, irrespective of which proteinase was produced, and for MG1363 (aPI). Reduction in optical density was highest for MG1363 expressing the hybrid proteinase aPIac or the inactive proteinase aPI* (Fig. 2A). An intermediate reduction in optical density was observed with the cultures of MG1363 producing no proteinase, sPI, or aPIabc.

Release of PepX activity was in full agreement with the above results (Fig. 2B): the higher the reduction in optical density measured at 600 nm, the more PepX activity was detectable in the culture supernatant.

Release of intracellular proteins was also examined by SDS-PAGE (12.5% polyacrylamide gel) analysis of supernatant samples of the various MG1363 cultures taken at the end of the exponential growth phase and after 1 and 3 days of incubation at 30°C (Fig. 2A). Coomassie brilliant blue staining (results not shown) of supernatant fractions of all strains except MG1363 (aPI) revealed a protein banding pattern typical for that of intracellular proteins of L. lactis (4). Hardly any protein was visible in the supernatants of MG1363 (aPI). The largest amount of protein was detected in the supernatant of the 3-day samples of MG1363 (aPI*) and MG1363 (aPIac). No band was detected at the position of lactococcal PrtP, whereas in all samples a band corresponding to Usp45 was present, indicating that, as expected, only little PrtP is formed in M17 medium (21). This amount of activity was still detectable by an enzymatic assay using the chromogenic peptide substrate MeO-Suc-Arg-Pro-Tyr-*p*NA (not shown).

Apparently, when the proteinase remains attached to the cell wall, autolysis is strongly influenced by the specificity of the proteinase: a PI-type proteinase results in strongly reduced autolysis, whereas autolysis is maximal when cells produce a proteinase with a specificity intermediate between those of PI and PIII.

Proteinase specificity is an important parameter in AcmA degradation. To examine whether proteinase specificity is important for AcmA degradation, we analyzed the pattern of active degradation products of AcmA. For this purpose, samples taken from all MG1363 cultures at the end of the exponential growth phase and after 1 and 3 days of incubation at 30°C (Fig. 2A) were examined by zymographic analysis. Figure 3 shows that the banding pattern of extracellular AcmA activity of MG1363 (P⁻) and of cells producing aPI* are very similar. Analysis of the supernatants of the other cultures shows that AcmA is degraded to various degrees during the 3-day incubation. In the supernatant of MG1363 (aPI), degradation of AcmA is already clearly observed at the end of the exponential

AcmA is already clearly observed at the end of the exponential growth phase. Degradation proceeds much more slowly in su-pernatant fractions of MG1363 producing aPIabc or aPIac. Similar amounts of both the unprocessed (46.1-kDa) and mature (40.3-kDa) forms of AcmA are present in the cell extracts of the three samples of all strains except MG1363 (aPI). In the latter strain, the amounts of both forms of AcmA were significantly lower at days 1 and 3. A band of activity of one of the major degradation products of AcmA was clearly present in the cell extracts of all three samples of MG1363 present in the cell extracts of all three samples of MG1363 (aPI) and in the cell extract of the strain secreting sPI sampled at day 3 (not shown). These results indicate that degradation of AcmA is influenced by the specificity of the proteinase PrtP.

The proteinase PrtP is involved in cell separation. In the course of the experiments described above, we observed a difference in sedimentation among MG1363acmA $\Delta 1$ strains \aleph producing the various proteinases. In contrast to MG1363 N (AcmA⁺), the mutant MG1363acmA $\Delta 1$ grows in extremely § long chains and sediments during overnight growth (5). Sedimentation of cells was also observed in overnight cultures of MG1363acmA $\Delta 1$ producing aPI* or sPI. This phenomenon was not seen with MG1363acmA $\Delta 1$ expressing an active cell wall-anchored proteinase (aPI, aPIabc, or aPIac). No sedimentation occurred in cultures of L. lactis MG1363 expressing either of these proteinases. Light microscopic analysis showed that all MG1363 strains had the same average chain length [Fig. 4; only shown for MG1363 (P⁻)]. As shown in Fig. 4, the length of the chains of MG1363acmA $\Delta 1$ is hardly influenced by the presence of an inactive cell-anchored proteinase [compare strains MG1363acmA $\Delta 1$ (P⁻) and MG1363acmA $\Delta 1$ (aPI*)]. In the presence of the unanchored, secreted form (sPI), the average chain length became only slightly shorter. However, the chains were found to be significantly shorter when cells produced an active proteinase that remained bound to the cell wall. The shortest chains (containing mainly two cells) were detected when MG1363acmA $\Delta 1$ expressed a hybrid anchored proteinase (aPIabc [Fig. 4] or aPIac [not shown]). In fact, the average chain length in these cultures was very similar to that of MG1363 (Prt⁻ AcmA⁺). These results were highly repro-



FIG. 4. Light microscopic analysis of the chain lengths in overnight M17 cultures of the indicated strains. Magnification, ×1,000.

ducible, and the same effects were observed when the various strains were grown in whey-based medium.

DISCUSSION

The results presented in this work show that the autolysin AcmA of *L. lactis* is subject to degradation by the lactococcal proteinase (caseinase) PrtP. AcmA activity, which is normally present in the supernatant of a culture of *L. lactis* MG1363 grown on whey-based medium, was completely absent in the supernatant of a strain secreting sPI. One specific degraded form of AcmA, barely visible in Fig. 1B but clearly present after prolonged renaturation, was present in the supernatants of MG1363 (P^-) and MG1363 producing aPI. This truncated form of AcmA is formed through cleavage of the C-terminal

cell wall binding domain by an as yet unidentified proteolytic activity in *L. lactis*. The protein is still active but, depending on the exact position of the cleavage site, may have reduced cell binding capacity (3). Reduced binding may explain the presence of this protein in the supernatant of MG1363 (aPI), as there it would escape degradation by aPI. The extent of degradation of AcmA by sPI or aPI is much less when the strains were grown in M17 (compare Fig. 1 and 3). This is due to lower expression of PrtP in this medium compared to whey permeate: the PrtP protein was detected in the supernatant fraction of the whey cultures by SDS-PAGE but was not visible in this fraction of M17 cultures (reference 31 and data not shown). Also, Usp45 was still present in the M17 culture media but had been degraded in the proteinase-producing whey cultures.

The inactive mutant aPI* does not degrade AcmA, either when AcmA is present in the cell walls (not shown) or when it is secreted in the culture medium (Fig. 3). Zymographic analvsis of AcmA activity, followed over time, in the supernatants of M17 cultures of MG1363 expressing different types of proteinases showed that AcmA degradation was highest when the cells expressed a PI-type proteinase. AcmA hydrolysis was significantly lowered when a hybrid proteinase was expressed (aPIabc or aPIac) with a caseinolytic specificity different from that of PI. The mature form of AcmA, which is normally located in the cell wall of L. lactis (5), was present in undiminished amounts in all samples in the cell walls of strains producing no PrtP, sPI, aPI*, or the two hybrid proteinases (results not shown). As AcmA is a good substrate for sPI and is fully degraded by the proteinase once the protein is released in the medium (Fig. 3), we assume that sPI has no access to the cell wall-located autolysin. When MG1363 expressed aPI, the amount of AcmA activity on the cells gradually decreased in time (not shown). While mature AcmA was still present on the cells at day 3, it was not detectable in the supernatant of these cells. It may be that AcmA, due to conformational constraints, is inaccessible for PrtP when the autolysin is bound to its substrate. After hydrolysis of peptidoglycan, AcmA would release and be subject to degradation by PrtP. Such a phenomenon has been described for partially purified glucosaminidase of B. subtilis. This enzyme preparation was unstable due to the action of contaminating proteinases. Addition of purified cell walls of *B. subtilis* resulted in partial resistance to proteolytic attack (24). Trypsin and α -chymotrypsin degradation of the pneumococcal LytA amidase and CPL1 lysozyme could be slowed down by the addition of choline, which induces a conformational change in these choline-dependent enzymes. Accordingly, the choline-independent CPL7 lysozyme was not protected by addition of this agent (26).

Due to AcmA degradation, autolysis was nearly absent in cells expressing a PI-type anchored proteinase. Although the reduction of the optical density at 600 nm did not reveal lysis, PepX and protein release in time from MG1363 (aPI) were evident. Detachment of the proteinase from the cell wall by removal of the cell wall anchor sequence (as in sPI) resulted in an extent of autolysis similar to that for the PrtP-negative strain MG1363. As AcmA is actively degraded in the supernatant of cells producing sPI, we presume that it is the cell wall-located autolysin that is involved in cell lysis. As detailed above, this fraction of AcmA is not affected by sPI. We expected autolysis of the aPI*-producing strain to be the same as in a PrtP-negative strain. The amounts of AcmA in the supernatant and on cells of MG1363 (aPI*) were the same as those for MG1363 (P⁻). Nevertheless, a higher level of cell lysis was observed with MG1363 (aPI*). This observation suggests that anchoring to the cell walls of PrtP destabilizes the cells in an as yet unknown way. The anchored hybrid proteinase aPIac also caused increased lysis of producing cells, while aPIabc had no effect on autolysis compared with MG1363 (P⁻). At this moment it is not clear what causes the difference in autolytic behavior of the two strains. The AcmA degradation patterns of cell extracts (not shown) and supernatants are very similar. The enzymes aPIabc and aPIac differ in only 9 of 1,902 amino acid residues. These residues are located in a domain in PrtP, which has previously been found to be of minor importance with respect to specificity of casein degradation (31) but may influence cell wall stability once the proteinase is anchored to the cells. When aPIac was expressed in an acmA deletion mutant, this strain did not autolyse; thus, the proteinase is not itself involved in degradation of the cell wall to an extent that cell lysis would occur.

The average chain length of MG1363*acm* $A\Delta 1$ expressing either of the hybrid proteinases aPIac and aPIabc was shorter than that of the same strain expressing aPI and nearly identical to that of MG1363. It is possible that cell wall proteins are degraded by the proteinases, thus possibly destabilizing the cell wall. This supposition seems to be strengthened by the observation that expression of sPI by MG1363*acm* $A\Delta 1$ does not result in such a drastic shortening of the average chain length.

The results presented in this report show that PrtP degrades AcmA and that the rate and degree of autolysis of *L. lactis* are dependent on the specificity, location, and amount of PrtP produced.

ACKNOWLEDGMENTS

We thank Henk Mulder for preparing the photographs and Anne de Jong for advice on and support of the computer work.

This study was supported by Unilever Research Laboratorium, Vlaardingen, The Netherlands. Jan Kok was the recipient of a fellowship from the Royal Netherlands Academy of Arts and Sciences.

REFERENCES

- Ayusawa, D., Y. Yoneda, K. Yamane, and B. Maruo. 1975. Pleiotropic phenomena in autolytic enzyme(s) content, flagellation, and simultaneous hyperproduction of extracellular α-amylase and protease in a *Bacillus subtilis* mutant. J. Bacteriol. 124:459–469.
- Béliveau, C., C. Potvin, J. Trudel, A. Asselin, and G. Bellemare. 1991. Cloning, sequencing, and expression in *Escherichia coli* of a *Streptococcus faecalis* autolysin. J. Bacteriol. 173:5619–5623.
- 3. **Buist, G.** 1997. AcmA of *Lactococcus lactis*, a cell-binding muramidase. Ph.D. thesis. University of Groningen, Groningen, The Netherlands.
- Buist, G., H. Karsens, A. Nauta, D. van Sinderen, G. Venema, and J. Kok. 1997. Autolysis of *Lactococcus lactis* caused by induced overproduction of its major autolysin, AcmA. Appl. Environ. Microbiol. 63:2722–2728.
- Buist, G., J. Kok, K. J. Leenhouts, M. Dabrowska, G. Venema, and A. J. Haandrikman. 1995. Molecular cloning and nucleotide sequence of the gene encoding the major peptidoglycan hydrolase of *Lactococcus lactis*, a muramidase needed for cell separation. J. Bacteriol. 177:1554–1563.
- Chopin, A. 1993. Organization and regulation of genes for amino acid biosynthesis in lactic acid bacteria. FEMS Microbiol. Rev. 12:21–37.
- Coxon, R. D., C. R. Harwood, and A. R. Archibald. 1991. Protein export during growth of *Bacillus subtilis*: the effect of extracellular protease deficiency. Lett. Appl. Microbiol. 12:91–94.
- 8. de Vos, W. M., P. Vos, H. de Haard, and I. Boerrigter. 1989. Cloning and expression of the *Lactococcus lactis* subsp. *cremoris* SK11 gene encoding an extracellular serine proteinase. Gene **85:**169–176.
- 9. Exterkate, F. A., A. C. Alting, and P. G. Bruinenberg. 1993. Diversity of cell envelope proteinase specificity among strains of *Lactococcus lactis* and its relationship to charge characteristics of the substrate-binding region. Appl. Environ. Microbiol. **59**:3640–3647.
- Gasson, M. J. 1983. Plasmid complements of *Streptococcus lactis* NCDO 712 and other lactic streptococci after protoplast-induced curing. J. Bacteriol. 154:1–9.
- Haandrikman, A. J., R. Meesters, H. Laan, W. N. Konings, J. Kok, and G. Venema. 1991. Processing of the lactococcal extracellular serine proteinase. Appl. Environ. Microbiol. 57:1899–1904.
- Jolliffe, L. K., R. J. Doyle, and U. N. Streips. 1980. Extracellular proteases modify cell wall turnover in *Bacillus subtilis*. J. Bacteriol. 141:1199–1208.
- Kariyama, R., and G. D. Shockman. 1992. Extracellular and cellular distribution of muramidase-2 and muramidase-1 of *Enterococcus hirae* ATCC 9790. J. Bacteriol. 174:3236–3241.
- Kok, J. 1992. Special-purpose vectors for lactococci, p. 97–102. *In* G. Dunny, P. P. Cleary, and L. L. McKay (ed.), Genetics and molecular biology of streptococci, lactococci, and enterococci. American Society for Microbiology, Washington, D.C.
- Kok, J., and W. M. de Vos. 1994. The proteolytic system of lactic acid bacteria, p. 169–210. *In* M. J. Gasson and W. M. de Vos (ed.), Genetics and biotechnology of lactic acid bacteria. Blackie and Professional, London, England.
- Kok, J., J. M. van Dijl, J. M. B. M. van der Vossen, and G. Venema. 1985. Cloning and expression of a *Streptococcus cremoris* proteinase in *Bacillus subtilis* and *Streptococcus lactis*. Appl. Environ. Microbiol. 50:94–101.
- Komatsuzawa, H., M. Sugai, S. Nakashima, S. Yamada, A. Matsumoto, T. Oshida, and H. Suginaka. 1997. Subcellular localization of the major autolysin, ATL and its processed proteins in *Staphylococcus aureus*. Microbiol. Immunol. 41:469–479.

- Laemmli, U. K. 1970. Cleavage of structural proteins during the assembly of the head of bacteriophage T4. Nature (London) 227:680–685.
 Leenhouts, K. J., and G. Venema. 1993. Lactococcal plasmid vectors, p.
- Leenhouts, K. J., and G. Venema. 1993. Lactococcal plasmid vectors, p. 65–94. *In* K. G. Hardy (ed.), Plasmids: a practical approach. Oxford University Press, Oxford, England.
- López, R., C. Ronda-Lain, A. Tapia, S. B. Wacks, and A. Tomasz. 1976. Suppression of the lytic and bactericidal effects of cell wall inhibitory antibiotics. Antimicrob. Agents Chemother. 10:697–706.
- Meijer, W., J. D. Marugg, and J. Hugenholtz. 1996. Regulation of proteolytic enzyme activity in *Lactococcus lactis*. Appl. Environ. Microbiol. 62:156–161.
- Mierau, I., E. R. S. Kunji, K. J. Leenhouts, M. A. Hellendoorn, A. J. Haandrikman, B. Poolman, W. N. Konings, G. Venema, and J. Kok. 1996. Multiple-peptidase mutants of *Lactococcus lactis* are severely impaired in their ability to grow in milk. J. Bacteriol. **178**:2794–2803.
- Navarre, W. W., and O. Schneewind. 1994. Proteolytic cleavage and cell wall anchoring at the LPXTG motif of surface proteins in Gram-positive bacteria. Mol. Microbiol. 14:115–121.
- Rashid, M. H., M. Mori, and J. Sekiguchi. 1995. Glucosaminidase of *Bacillus subtilis*: cloning, regulation, primary structure and biochemical characterization. Microbiology 141:2391–2404.
- Sambrook, J., E. F. Fritsch, and T. Maniatis. 1989. Molecular cloning: a laboratory manual, 2nd ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.
- Sanz, J. M., E. Díaz, and J. L. García. 1992. Studies on the structure and function of the N-terminal domain of the pneumococcal murein hydrolases. Mol. Microbiol. 6:921–931.
- 27. Seegers, J. F. M. L., S. Bron, C. M. Franke, G. Venema, and R. Kiewiet. 1994.

The majority of lactococcal plasmids carry a highly related replicon. Microbiology **140**:1291–1300.

- van Asseldonk, M., G. Rutten, M. Oteman, R. J. Seizen, W. M. de Vos, and G. Simons. 1990. Cloning of *usp45*, a gene encoding a secreted protein from *Lactococcus lactis* ssp. *lactis* MG1363. Gene 95:155–160.
- 29. van de Guchte, M., J. Kodde, J. M. B. M. van der Vossen, J. Kok, and G. Venema. 1990. Heterologous gene expression in *Lactococcus lactis* subsp. *lactis*: synthesis, secretion, and processing of the *Bacillus subtilis* neutral protease. Appl. Environ. Microbiol. 56:2606–2611.
- 30. Visser, S., F. A. Exterkate, C. J. Slangen, and G. J. C. M. de Veer. 1986. Comparative study of action of cell wall proteinases from various strains of *Streptococcus cremoris* on bovine α_{S1}-, β-, and κ-casein. Appl. Environ. Microbiol. 52:1162–1166.
- 31. Vos, P., I. J. Boerrigter, G. Buist, A. J. Haandrikman, M. Nijhuis, M. B. de Reuver, R. J. Siezen, G. Venema, W. M. de Vos, and J. Kok. 1991. Engineering of the *Lactococcus lactis* serine proteinase by construction of hybrid enzymes. Protein Eng. 4:479–484.
- Wilkinson, M. G., T. P. Guinee, D. M. O'Callaghan, and P. F. Fox. 1994. Autolysis and proteolysis in different strains of starter bacteria during Cheddar cheese ripening. J. Dairy. Res. 61:249–262.
- Yabu, K., and S. Kaneda. 1995. Salt-induced cell lysis of *Staphylococcus aureus*. Curr. Microbiol. 30:299–303.
- Yabu, K., Y. Nishiyama, and T. Ochiai. 1997. Protease-induced multicell formation in *Staphylococcus haemolyticus*. Microbiol. Immunol. 41:799–803.
- 35. Yoneda, Y., and B. Maruo. 1975. Mutation of *Bacillus subtilis* causing hyperproduction of α -amylase and protease and its synergistic effect. J. Bacteriol. 124:48–54.