Efficient secretion of the model antigen M6–gp41E in Lactobacillus plantarum NCIMB 8826

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Four Lactobacillus strains (Lb. plantarum NCIMB 8826, Lb. paracasei LbTGS1.4, Lb. casei ATCC 393 and Lb. fermentum KLD) were tested for their ability to produce and secrete heterologous proteins. These strains were first screened with an a-amylase reporter under the control of a set of expression or expression/secretion signals from various lactic acid bacteria. With most of the constructions tested, the level of extracellular production was highest in Lb. plantarum NCIMB 8826, and lowest in Lb. paracasei LbTGS1.4. These two strains were next assayed using a model antigen consisting of the N-terminal part of the M6 protein from Streptococcus pyogenes fused to the linear epitope ELDKWAS from human immunodeficiency virus gp41 protein. Secretion of this heterologous protein was inefficient in Lb. paracasei LbTGS1.4, which accumulated a large intracellular pool of the unprocessed precursor, whereas Lb. plantarum NCIMB 8826 was able to secrete the antigen to a level as high as 10 mg l⁻¹.

Keywords: Lactobacillus, secretion, a-amylase, M6 protein, mucosal vaccines

INTRODUCTION

Lactobacilli are used in various processes such as dairy fermentation, baking, production of alcoholic beverages, sausage and meat preservation, pickling of vegetables and silage making. Some members of the genus are commensal colonizers of mammalian gastrointestinal systems and this colonization is thought to have beneficial effects on health and nutrition (Chassy, 1987; Fernandes et al., 1987; Marteau & Rambaud, 1993; Pouwels & Leer, 1993). Recently, lactobacilli have been considered as potential antigen delivery vehicles for the development of mucosal vaccines since they display a number of advantages. Lactobacilli are GRAS (generally recognized as safe) micro-organisms. Furthermore, lactobacilli are able to cross the digestive tract and some of them are maintained transiently or for a longer term as a stable population in the gut (Link-Amster et al., 1994; Marteau & Rambaud, 1993). Finally, they possess intrinsic adjuvancy and low immunogenicity (Pouwels et al., 1996; Perdigon et al., 1988). Lactobacilli are also an alternative to the other GRAS lactic acid bacterium Lactococcus lactis, which has also been evaluated as a live antigen carrier (Norton et al., 1994; Wells et al., 1993a), the main difference being the capacity of some lactobacilli to colonize the gut mucosa while no implantation can be detected for Lc. lactis (Klijn et al., 1995).

An important consideration in the development of a live mucosal vaccine concerns the location of the antigen (intracellular, extracellular or cell-wall anchored). With live enteric vectors such as Salmonella typhimurium or Escherichia coli, cell surface presentation has been considered advantageous for efficiently inducing an antibody response (Leclerc et al., 1991; Schorr et al., 1991). Cell surface exposure of antigens or epitopes in Gram-positive bacteria has only recently been investigated. An anchoring system for the mouth commensal bacterium Streptococcus gordonii has been developed on the basis of the fibrillar M6 protein (Oggioni et al., 1995; Fischetti et al., 1993), whereas anchoring in the non-pathogenic bacteria Staphylococcus xylosus and Staphylococcus carnosus is based on protein A from...
Staphylococcus aureus (Samuelson et al., 1995; Nguyen et al., 1993). Like nearly all surface molecules from Gram-positive bacteria, these two proteins follow the general secretion pathway for their external location, and anchoring per se takes place in a second step owing to the presence of an anchoring hydrophobic C-terminal domain associated with a conserved motif, LPXTGX (Scheuwein et al., 1995; Fischetti et al., 1993).

The general context of this work deals with the implementation of the M6 epitope carrier system in lactobacilli and more precisely concerns the evaluation of the secretion capacity of different Lactobacillus strains, since secretion is a prerequisite of the anchoring process. Secretion efficiency is clearly a limitation factor in the production of a model antigen comprising the N-terminal part of the M6 protein fused to an epitope derived from the human immunodeficiency virus (HIV) gp41 protein (M6-gp41E).

METHODS

Plasmids, bacterial strains and culture conditions. The plasmids and bacterial strains used in this study are listed in Table 1. Plasmids used in lactobacilli all replicated as rolling circles. E. coli MC1061 was used as host strain for the propagation of plasmids pTG2247, pTG2257, pTG2281 and pTG3237; the other plasmids were prepared from E. coli TG1. All Lactobacillus strains were routinely grown in MRS medium (Difco). Amylase activity was detected on plates by incorporating 0.2% (w/v) starch in MRS medium containing 0.2% (w/v) glucose instead of 20% (w/v) starch and subsequently visualizing starch degradation haloes by staining with iodine vapours (Merck). Antibiotics were used at the following concentrations: 100 µg chloramphenicol ml⁻¹ for LbTGSl.4, 7.5 µg chloramphenicol ml⁻¹ for Lactobacillus paracasei LbTGS1.4, 10 µg chloramphenicol ml⁻¹ for Lactobacillus plantarum NCIMB 8826; 7.5 µg chloramphenicol ml⁻¹ for Lactobacillus paracasei LbTGS1.4, Lactobacillus fermentum KLD and Lactobacillus casei ATCC 393.

DNA manipulations and transformation. All DNA manipulations were carried out as described previously (Hols et al., 1992, 1994). Plasmid DNA from Lb. plantarum was isolated as described by Posno et al. (1991). Plasmid DNA from Lb. paracasei LbTGS1.4 and Lb. casei ATCC 393 was isolated as described by O’Sullivan & Klaenhammer (1993). Preparation of plasmid DNA from Lb. fermentum was performed by the same procedure except that cells from a 2 ml overnight culture were harvested by centrifugation and resuspended in 400 µl 25% (w/v) sucrose containing 50 mg lysozyme ml⁻¹ and further incubated at 37 °C for 1 h. Electroporation of Lb. plantarum NCIMB 8826 was performed as described by Josson et al. (1989). Electroporation of Lb. paracasei LbTGS1.4 and Lb. casei ATCC 393 was conducted as follows. An overnight culture was diluted 100-fold in 300 ml MRS medium supplemented with 20 mM D/L-threonine. Cells were harvested by centrifugation and washed three times with ice-cold electroporation buffer (272 mM sucrose, 1 mM HEPES, pH 6.5) and finally resuspended in 3 ml of the same buffer. The cell suspension was mixed with 100 ng plasmid DNA and transferred into an electroporation cuvette (0.2 cm). A single pulse of 11 kV was applied with a capacitance of 25 µF and a resistance of 200 Ω. After electroporation, cells were directly diluted with 300 µl MRS and incubated for 2 h at 37 °C. Transformants were visible after 48 h incubation at 37 °C under aerobic conditions. Electroporation of Lb. fermentum KLD was performed according to the previous protocol with the following modifications: the overnight culture was diluted 100-fold in 50 ml MRS medium without threonine, cells were harvested at an OD₆₀₀ of 0.2 and the electroporation buffer was adjusted to pH 6.0 and the cells were resuspended in 1 ml of the same buffer after the washing steps. After electroporation, cells were directly diluted with 1 ml MRS and incubated for 2 h at 37 °C. Plates were incubated under anaerobic conditions (Gaspak System; BRL).

z-Amylase assay. Cell culture and fractionation and the z-amylase activity test (Phedebas kit; Pharmacia) were performed as described previously (Hols et al., 1994). Total protein in cell extracts was measured with Bio-Rad protein assay reagent on the basis of the Bradford (1976) staining procedure.

Western blotting and ELISA assay. Protein extracts and culture supernatants were prepared from MRS cultures grown for 9 h at 37 °C. Total protein extracts were prepared by sonication (Sonifier 450; Branson Ultrasonics); cell lysates were usually obtained after 12–16 × 30 s. Culture supernatants from Lb. plantarum NCIMB 8826 were directly mixed with Laemmli sample buffer, whereas those from Lb. paracasei LbTGS1.4 were filtered through 0.22 µm Millex HA (Millipore) membranes and precipitated with 10% (w/v) ice-cold TCA. After centrifugation, the pellets were washed once with ice-cold ethanol and resuspended in Laemmli sample buffer (Laemmli, 1970). Protein extracts and culture supernatants were separated by SDS-PAGE (Laemmli, 1970) and electrophoretically blotted onto nitrocellulose sheets as described by Towbin et al. (1979). The subsequent steps were carried out as specified by the suppliers of the Western blot AP system (Protoblot; Promega). Immunoblots were performed with anti-gp41E antibodies (human monoclonal 2F5; 1/7500 dilution) as primary antibodies (Durot, 1996). Rainbow markers (Amersham) were used as standards. The ELISA assay was based on the end-point method. Anti-gp41E antibodies (1/7500 dilution) and biotinylated anti-mouse antibodies (1/500 dilution; Amersham) were used as primary and secondary antibodies, respectively. After the addition of the streptavidin/peroxidase complex (1/1000 dilution; Amersham), staining was obtained by incubation with o-phenylenediamine (Sigma) plus hydrogen peroxide. Plate readings were made with a Vmax machine ( Molecular Devices) at 490 nm.

RESULTS

Screening of Lactobacillus strains for extracellular production using z-amylase as reporter

A first step towards the development of oral vaccines based upon recombinant live lactobacilli was to evaluate the ability of various commensal species to efficiently secrete a model foreign protein. We first selected Lb.
<table>
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<th>Table 1. Plasmids and bacterial strains</th>
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<tr>
<td>Plasmid or strain</td>
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<td><strong>Plasmids</strong></td>
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<td>pTG290</td>
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<tr>
<td>pGIT008</td>
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<tr>
<td>pGIP312.4</td>
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<tr>
<td>pGIP212.4</td>
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<td>pGIP212.7 and pGIP212.12</td>
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<tr>
<td>pNZ10x1</td>
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<td>pTG3237</td>
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<td>pGK13</td>
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<td><strong>Strains</strong></td>
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<td><em>E. coli</em> TG1</td>
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<tr>
<td><em>Lb. casei</em> ATCC 393</td>
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<tr>
<td><em>Lb. fermentum</em> KLD</td>
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<tr>
<td><em>Lb. paracasei</em></td>
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<tr>
<td>LbTGSl.4</td>
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<tr>
<td><em>Lb. plantarum</em> NCIMB 8826</td>
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**paracasei** LbTGSl.4, a murine isolate of vaginal origin able to colonize the gastro-intestinal and vaginal tracts of the mouse for about 1 and 3 weeks, respectively (Dutot, 1996); this strain was of interest since the mouse was chosen first as an experimental system. We also tested three additional strains: *Lb. fermentum* KLD from human faeces, *Lb. plantarum* NCIMB 8826 from human saliva and *Lb. casei* ATCC 393 from cheese.

A range of α-amylase constructs was tested in these different hosts. An α-amylase reporter (*amyL* from *Bacillus licheniformis* or *amyS* from *Bacillus stearothermophilus*) was brought under the control of different expression or expression/secretion signals originating from different LAB such as *Lb. plantarum* (pGIT008, pGIP212.7, pGIP212.12), *Lc. lactis* (pNZ10x1), *Streptococcus thermophilus* (pTG290) and *Enterococcus faecalis* (pGIP212.4, pGIP312.4) (Table 1). The genetic fusions were simple transcriptional fusions in pTG290 and pGIT008 or translational fusions between expression/secretion signals and the α-amylase reporter in all the other constructs tested.

Seven α-amylase constructs were transferred by electroporation into *Lb. plantarum* NCIMB 8826 and *Lb. paracasei* LbTGSl.4. The α-amylase phenotype was
tested on MRS plates containing 0.2% (w/v) glucose and 0.2% (w/v) starch (Fig. 1). All the recombinant strains displayed a positive phenotype in NCIMB 8826 and LbTGS1.4 with the exception of pGIP212.12 in LbTGS1.4. A visual comparison of the halo in NCIMB 8826 and LbTGS1.4 recombinant strains clearly showed that all the NCIMB 8826 recombinant strains displayed a larger halo size (Fig. 1). Fractionation experiments performed on these recombinant strains demonstrated that the α-amylase specific activity was generally higher in the supernatant of strain NCIMB 8826 (Table 2), confirming the observations made on the haloes. In most recombinant strains, α-amylase activity was mainly located in the supernatant, with the exception of pNZ10×1, where approximately 50% of the amylase activity was associated with the cell pellet (Table 2).

In addition to the transfer of a range of α-amylase constructs in Lb. plantarum NCIMB 8826 and Lb. paracasei LbTGS1.4, some constructions were also tested in Lb. fermentum KLD and Lb. casei ATCC 393. The transfer of plasmids pTG290, pGIP008 and pGIP312.4 was unsuccessful in Lb. fermentum, even after repeated trials. Plasmids pGIP212.4 and pNZ10×1 were successfully transferred in that strain but a majority of the primary transformants were α-amylase-negative, suggesting an early structural instability of these plasmids. This was confirmed for plasmid pNZ10×1, where a region of 1-2 kb was systematically deleted (data not shown). Four of these α-amylase constructs were successfully introduced and stably maintained in Lb. casei ATCC 393, and the results were similar to those obtained in Lb. paracasei LbTGS1.4 (Table 2).

**Construction of M6-gp41E expression/secretion plasmids**

To assess whether the observed differences in extracellular production would hold true for another protein, secretion of a model antigen (M6-gp41E) was evaluated in the two most divergent strains, Lb. plantarum NCIMB 8826 (high α-amylase production) and Lb.

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**Table 2. α-Amylase activity in culture supernatants and cell extracts of different recombinant Lactobacillus strains**

Activity is expressed as $10^{-2} \times \alpha$-amylase units (mg total protein)$^{-1}$, mean of two experiments. ND, Plasmid not transferred. –, Unsuccessful transfer or structural instability of the plasmid.

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Lb. plantarum NCIMB 8826</th>
<th>Lb. paracasei LbTGS1.4</th>
<th>Lb. casei ATCC 393</th>
<th>Lb. fermentum KLD</th>
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<tr>
<td></td>
<td>Supernatant</td>
<td>Cell extract</td>
<td>Supernatant</td>
<td>Cell extract</td>
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<tr>
<td>pTG290</td>
<td>52</td>
<td>&lt;1</td>
<td>12</td>
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<tr>
<td>pGIT008</td>
<td>148</td>
<td>12</td>
<td>7</td>
<td>&lt;1</td>
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<tr>
<td>pGIP312.4</td>
<td>87</td>
<td>11.5</td>
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<td>pGIP212.7</td>
<td>18</td>
<td>&lt;1</td>
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<tr>
<td>pGIP212.12</td>
<td>38</td>
<td>22</td>
<td>&lt;1</td>
<td>&lt;1</td>
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<tr>
<td>pNZ10×1</td>
<td>102</td>
<td>135</td>
<td>142</td>
<td>119</td>
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corresponds to promoter P25. The transcriptional start mapped
from the Lb. pentosus

The precursor protein (pre-M6-gp41E) was PCR-amplified from plasmid pTG2257 using the following oligonucleotides: OTG5508 (5'-AATGTCTAGAAT- AACAAGATAGACCTAT-3'), which included an XbaI site (underlined), and OTG5592 (3'-CGAATT- CTTCTTCCGTTGTGCCTGCAGCCGAG-5'), which included a SacI site (underlined). The 572 bp DNA fragment of the PCR reaction was cleaved by XbaI and SacI and inserted in the corresponding site of the expression vector pTG2247 (Table 1). A restriction map of the resulting pTG2281 plasmid and the organization of the fusion are presented in Fig. 2(a) and Fig. 2(b), respectively. We used the P25 promoter from S. thermophilus (Slos et al., 1991) and the Shine–Dalgarno sequence from the Lactobacillus pentosus ldhD gene (Taguchi & Ohta, 1991) to drive expression of the pre-M6-gp41E protein. To direct secretion through the general secretion pathway, we took the genuine 42 amino acid M6 signal sequence (Hollingshead et al., 1986) except that amino acids 2 and 3 (Ala and Lys) were changed into Ser and Arg as a result of using the XbaI site in the ATG fusion. This modification was expected to have a negligible effect on the function of the N-terminal part of the signal sequence since the number of positively charged amino acids remained unchanged in the ‘n’ region of the signal sequence (Simonen & Palva, 1993).

A number of Lb. plantarum translation initiation regions (TIRs) have been reported to contain additional motifs (5'-boxes) besides the Shine–Dalgarno sequences that could interact with the 5'-end of the 16S rRNA (Hols et al., 1994). Therefore, in an attempt to improve translation, a modified version of plasmid pTG2281 was constructed by inserting a 5'-box in front of the ldhD Shine–Dalgarno sequence. A synthetic linker composed of the complementary oligonucleotides OTG5930 (5'-TGAATGTCTAGAAGTTCTTGCGAAAGC-3') and OTG5931 (3'-CTAGACATATG- AACACGAATAGACCTAT-5') (compatible XbaI and SacI termini underlined) was inserted between the same sites in pTG2281. The resulting plasmid, pTG3237, contains in front of the ldhD Shine–Dalgarno sequence 14 nucleotides comple-

underlined. The start (ATG) and stop (TAA) codons of the M6-gp41E ORF are underlined. The lower box corresponds to the fusion protein M6-gp41E; the enclosed box delineates the signal sequence. The C-terminal epitope gp41E (bold) is underlined. The 572 bp DNA fragment of the PCR reaction was cleaved by XbaI and SacI and inserted in the corresponding site of the expression vector pTG2247 (Table 1). A restriction map of the resulting pTG2281 plasmid and the organization of the fusion are presented in Fig. 2(a) and Fig. 2(b), respectively. We used the P25 promoter from S. thermophilus (Slos et al., 1991) and the Shine–Dalgarno sequence from the Lactobacillus pentosus ldhD gene (Taguchi & Ohta, 1991) to drive expression of the pre-M6-gp41E protein. To direct secretion through the general secretion pathway, we took the genuine 42 amino acid M6 signal sequence (Hollingshead et al., 1986) except that amino acids 2 and 3 (Ala and Lys) were changed into Ser and Arg as a result of using the XbaI site in the ATG fusion. This modification was expected to have a negligible effect on the function of the N-terminal part of the signal sequence since the number of positively charged amino acids remained unchanged in the ‘n’ region of the signal sequence (Simonen & Palva, 1993).

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mentary to the 5'-end of the 16S rRNA of \textit{Lb. casei} [\(\Delta G = -24\ \text{kcal mol}^{-1} (-100.8\ \text{kJ mol}^{-1})\); Fig. 2c] and 11 nucleotides complementary to the 5'-end of the 16S rRNA of \textit{Lb. plantarum} [\(\Delta G = -13.0\ \text{kcal mol}^{-1} (-54.6\ \text{kJ mol}^{-1})\); Fig. 2c].

**Secretion of M6–gp41E in \textit{Lb. plantarum NCIMB 8826} and \textit{Lb. paracasei LbTGS1.4} transformed with plasmid pTG2281**

Plasmid pTG2281 was transferred by electroporation into \textit{Lb. plantarum} NCIMB 8826 and \textit{Lb. paracasei} LbTGS1.4. A Western blot analysis was performed with cell extracts and supernatant fractions of the two recombinant strains (Fig. 3). The precursor M6–gp41E protein (apparent \(M_r 29000\)) was detected as the main band in the cell extract (Fig. 3, lane 1) and as a minor band in the supernatant (Fig. 3, lane 3) of \textit{Lb. paracasei} LbTGS1.4 (pTG2281). The main band (apparent \(M_r 23000\)) detected in 1 ml supernatant from LbTGSl.4 (pTG2281) corresponds to the mature protein. This band moved slightly behind the M6–gp41E reference protein (Fig. 3, lane 5), which had been purified from \textit{E. coli} and had a lower \(M_r\) than the mature M6–gp41E produced in this work due to modifications of its N-terminus [RVFP to (M)VFP] and C-terminus (LWNYKA to LWN) (P. Slos, unpublished results). The situation was completely different with the recombinant \textit{Lb. plantarum} NCIMB 8826 strain. The same amount of total cell protein (30 \(\mu\)g) as in the case of the LbTGS1.4 recombinant strain was loaded (Fig. 3, lane 10) and no precursor was detected. Furthermore, in only 24 \(\mu\)l of unconcentrated supernatant from NCIMB 8826 (pTG2281), we detected a strong band with the same mobility as the major band observed in the TCA precipitate from 1 ml supernatant of LbTGS1.4 (Fig. 3, lane 11). These observations clearly showed that the pre-M6–gp41E precursor accumulated intracellularly in \textit{Lb. paracasei} LbTGS1.4 and only a small fraction of the unprocessed protein was converted into the mature form which could be detected exclusively in the supernatant fraction after strong concentration. Conversely, the precursor protein did not accumulate in \textit{Lb. plantarum} NCIMB 8826 cells and large amounts of the mature form were detected in the culture supernatant. The amounts of extracellular M6–gp41E proteins were quantified by ELISA essay; 0.2 mg M6–gp41E \(1^{-1}\) was produced by the recombinant LbTGSl.4 strain whereas extracellular production amounted to 1.2 mg \(1^{-1}\) in strain NCIMB 8826.

**Secretion of M6–gp41E in \textit{Lb. plantarum NCIMB 8826} and \textit{Lb. paracasei LbTGS1.4} transformed with plasmid pTG3237**

Plasmid pTG3237 was also transferred in \textit{Lb. paracasei} LbTGSl.4 and in \textit{Lb. plantarum} NCIMB 8826. Quantification by ELISA of the extracellular amount of M6–gp41E was performed directly on the supernatants of the two recombinant strains. The amount of the protein produced was low and highly variable among the five different LbTGSl.4 clones studied. However, a 10-fold increase of extracellular production (10.4 mg \(1^{-1}\)) was observed for NCIMB 8826 in comparison to the same strain harbouring plasmid pTG2281. Two bands corresponding to the precursor and the mature forms were detected by Western blotting in the cell fraction of NCIMB 8826 (pTG3237) (Fig. 3, lane 6), whereas no band was detected in the same amount of cell protein in NCIMB 8826 (pTG2281) (Fig. 3, lane 10). In the extracellular fraction, a single band corresponding to the mature form was detected with a loading of only 2 \(\mu\)l...
unconcentrated supernatant (Fig. 3, lane 8). Thus, modifying the region in front of the Shine–Dalgarno sequence dramatically boosted the expression level in NCIMB 8826, with the result that a large increase in extracellular production of M6–gp41E was achieved, together with the retention of small amounts of the precursor and the mature forms inside the cell.

**DISCUSSION**

The main goal of this study was to investigate the secretion capacity of a few strains of lactobacilli for their future use as a live antigen carrier in the development of mucosal vaccines. These strains were initially selected on the basis of different criteria such as genetic amenability (e.g. transformation), human or dairy origin and their capacity to colonize the gastro-intestinal tract of the mouse used as a model system.

The first part of this work was an evaluation of the capacity of four selected strains to deliver an α-amylase reporter extracellularly under the control of a set of expression or secretion signals from different LAB. This reporter was chosen since it has already been used with success for secretion studies in a number of LAB (Hols et al., 1992, 1994; Perez-Martinez et al., 1992; Van Asseldonk et al., 1993). This preliminary screening made clear that *Lb. plantarum* NCIMB 8826 displayed the highest level of amylase activity in the extracellular medium for the majority of the amylase constructs tested. *Lb. paracasei* LbTGS1.4 and *Lb. casei* ATCC 393 were much less efficient in this respect and *Lb. fermentum* KLD was excluded from further investigations as a majority of the plasmids tested in this strain were structurally unstable. Variation observed in the level of extracellular amylase production could of course result from differences in the level of transcription and/or translation, in mRNA stability or in the copy number of the plasmids. However, a range of expression/secreation signals from various LAB were assayed in this screening which all led to better amylase production in *Lb. plantarum* NCIMB 8826. Previously, *Lb. plantarum* NCIMB 8826 had already been shown to be an efficient secretor in comparison to other *Lb. plantarum* strains (Fitzsimons et al., 1994; Hols et al., 1994). Therefore, we believe that the low level of extracellular amylase production in strains LbTGS1.4 and ATCC 393 results from low efficiency of the secretion process. However, fractionation experiments did not confirm the expectation that a high amylase activity would be found in the intracellular fraction of the two strains except when plasmid pNZ10x1 was used, but in this case high intracellular amylase activity was observed in all the different hosts tested. Formally, we cannot exclude the accumulation of a inactive precursor, but preliminary Western blot experiments conducted with the pGTO08 amylase construct (*amyL* under the control of an *Lb. plantarum* promoter) in strains NCIMB 8826, LbTGS1.4 and ATCC 393 did not show any accumulation of the precursor (data not shown). Similar observations have been made using the same *amyL* reporter in *E. coli*, where it was shown that a defect in the secretion machinery (limitation in signal peptidase I) did not result in the accumulation of the precursor, whereas another reporter protein (*β-lactamase*) accumulated to a large extent (van Dijl et al., 1988). One of the explanations suggested by the authors is an extremely tight coupling between the processing and the synthesis of α-amylase (van Dijl et al., 1988). Alternatively, a proteolytic degradation of improperly folded precursors cannot be excluded.

We have retained the two most opposite strains (NCIMB 8826 and LbTGS1.4) for testing their capacity to secrete the model antigen M6–gp41E cloned in the expression vector pTG2281. Strain NCIMB 8826 was confirmed to be an efficient secretor; only the mature form of the antigen was present in the culture supernatant, and no precursor was detected intracellularly. The hypothesis that the low extracellular amylase activity in strain LbTGS1.4 would result from the poor secretion capacity of this strain was also clearly supported. A high intracellular retention (more that 90%) of the M6–gp41E precursor was observed in this case in LbTGS1.4 and the mature form of the protein could hardly be detected in the supernatant, where the precursor was also observed. The presence of the precursor in the culture supernatant is probably the result of cell lysis and we cannot exclude the possibility that the mature product observed in this case would result from non-specific proteolytic cleavage. At this stage, we can only speculate on these observations. The strong secretion defect observed for LbTGS1.4 could be due to poor efficiency of the secretion machinery itself or to a limitation or a complete lack of specific chaperones, as already postulated for a number of heterologous proteins refractory to secretion in Gram-positive bacteria (Simonen & Palva, 1993).

We constructed a second expression/secreation vector (pTG3237) with the objective of enhancing the level of expression of M6–gp41E by improving the TIR of the previous construction. Sequence analysis of TIR regions from a number of expression signals from *Lb. plantarum* had previously revealed the occurrence of motifs complementary to the 5'-end of 16s rRNA and the suggestion was made that these could strengthen translation initiation through mRNA–16s rRNA interactions (Hols et al., 1994). We therefore introduced a ‘5'-box' upstream of the Shine–Dalgarno sequence and this indeed improved antigen production by a factor of 10 in *Lb. plantarum*. However, this modification could have indirect effects such as modifying the secondary structure of the TIR region or improving mRNA stability, which are known to contribute to translation efficiency (de Smit & van Duin, 1990; Schauder & McCarthy, 1989). A search for specific modifications of secondary structures in the TIR region revealed the presence of a hairpin with a ΔG = −4.6 kcal mol⁻¹ (−19.32 kJ mol⁻¹) (Tinoco et al., 1973) encompassing the Shine–Dalgarno sequence in pTG2281 while the same region in pTG3237 was devoid of any structure. A more systematic approach would of course be needed to show a
direct correlation between the interaction with the 5'-end of the 16S rRNA and/or the presence of secondary structures and the level of expression.

The transfer of this improved M6–gp41E expression vector in LbTGS1.4 did not result in stable expression of the antigen. We can speculate that increasing the expression level could result in toxic effects through the antigen. We can speculate that increasing the vector in LbTGS1.4 did not result in stable expression of structures and the level of expression. The transfer of this improved M6-gp41E expression seems to be a limiting factor as the data with the precursor is probably the consequence of an overloading of the cell pellet has been observed in a number of cases dealing with heterologous secretion in Gram-positive bacteria (Simonen & Palva, 1993). In LAB, the most illustrative example is the secretion of the tetanus toxin fragment C (TTFC) in Lc. lactis, where a large amount of the mature protein remains associated with the cell fraction (Wells et al., 1993b). Most authors suggest that the cell wall may act as a barrier to the diffusion of some proteins (Simonen & Palva, 1993; Wells et al., 1993b; Saunders et al., 1987).

Production of 10 mg M6–gp41E (1 culture medium)⁻¹ is among the highest extracellular production levels for heterologous protein reported in LAB. To our knowledge, secretion in the milligram range was only published previously with a controlled expression system (based on the T7 promoter) in the case of TTFC (3 mg l⁻¹) and murine interleukin 2 (3 mg l⁻¹) in Lc. lactis (Steidler et al., 1995; Wells et al., 1993b). The major difference observed in the capacity of Lb. plantarum NCIMB 8826 and Lb. paracasei LbTGS1.4 to secrete the M6–gp41E protein was recently confirmed with another antigen, the cholera toxin B subunit (P. Slos, unpublished results). These achievements open the way for further developments in the construction of LAB as live vectors for mucosal immunization.

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