Kinetics of the secretion of *Bacillus subtilis* levanase overproduced during the exponential phase of growth

Laurence Leloup, Jérôme Le Saux, Marie-Françoise Petit-Glatron and Régis Chambert

Author for correspondence: Régis Chambert. Tel: +33 1 44 27 49 17. Fax: +33 1 44 27 59 94. e-mail: chambert@sccr.jussieu.fr

Institut Jacques Monod – CNRS, Universités Paris 6 et Paris 7, Laboratoire Génétique et Membranes, Tour 43-2, place Jussieu, F-75251 Paris Cedex 05, France

The *Bacillus subtilis* levanase structural gene sacC was expressed under the regulated control of sacR, the inducible levansucrase leader region, in a degU32(Hy) strain. In this genetic context, exocellular levanase is overproduced (0.5% of total protein) during the exponential phase of growth upon induction by sucrose at 37 °C and pH 7. No precursor form that comprised a signal peptide was detected in pulse-chase experiments. The subsequent release of the cell-associated processed protein is a slow event ($t_{1/2} = 80 \pm 10 \text{s}$). The unfolding-folding transition of pure levanase monitored in vitro by the resistance to proteolysis was achieved within the same time range ($t_{1/2} = 50 \text{s}$) under the same conditions of pH and temperature. Calcium ions, which modulate the rate and the yield of refolding, have a low affinity for the protein. Comparison of these results with those obtained previously with levansucrase and $\alpha$-amylase overproduced in the same genetic and physiological context suggests that the precursor processing is more efficient in levanase and $\alpha$-amylase than in levansucrase. This discrepancy could lie in information borne by the signal peptide sequence of these exoproteins. However, the rate of the ultimate stage of release of these three proteins, which includes the passage through the cell wall, is correlated with the rate of folding and appears to be independent of their molecular size.

**Keywords:** levanase, protein secretion, late stage of secretion, folding, *Bacillus subtilis*

**INTRODUCTION**

We have recently shown (Leloup et al., 1997) that the expression of the structural gene *amyE* under the control of the inducible levansucrase leader region *sacR* in a degU32(Hy) strain leads to an overproduction of $\alpha$-amylase. Such a context was suitable for the characterization in vivo of the sequential events of the secretion process of $\alpha$-amylase.

Comparison of the levansucrase and $\alpha$-amylase secretion mechanisms drew our attention to the fact that cleavage of the $\alpha$-amylase signal peptide was more rapid than that of levansucrase (Petit-Glatron et al., 1987; Leloup et al., 1997). Nevertheless, for both proteins, the final stage of mature protein release is slow, $t_{1/2} = 60 \text{s}$ and 120 s for levansucrase and $\alpha$-amylase, respectively, and is seemingly coupled with the folding.

The efficiency of the membrane translocation process could therefore be specific to each exoprotein family, such as proteins produced and secreted during the exponential phase (levansucrase) or the stationary phase (levanase or $\alpha$-amylase), whereas the mechanism underlying the final release could be common. In the present work, we examined the levanase SacC secretion pathway to test the validity of this general outline. We chose this protein for the following reasons. (i) Its signal sequence region (Schörgendorfer et al., 1987) displays, like $\alpha$-amylase, a hydrophobicity twice that of levansucrase (Scotti et al., 1996). Is this feature therefore correlated with the efficiency of the translocation mechanism? (ii) This protein has a higher molecular mass, 73 kDa, than levansucrase (50 kDa). Does such a difference affect the rate of release, which includes the passage through the cell wall, or does this rate only correlate with the unfolding kinetics of the protein? The answers to these questions could provide new information about each stage of the general mechanism of secretion we propose (Petit-Glatron et al., 1987; Leloup et al., 1997). *Bacillus subtilis* levanase is secreted at a low level during
the stationary phase of growth in the wild-type strain 168 Marburg (Fuchs, 1959), and \textit{degU32(Hy)} mutations inducing overproduction of several exocellular enzymes do not cause overproduction of levanase (Kunst et al., 1977).

Characterization \textit{in vivo} of the sequential events that lead up to secretion of levanase is not possible by conventional biochemical methods in this context of low production. We therefore placed \textit{sacC} under the regulated control of \textit{sacR}, the levanosucrase leader region, in a \textit{degU32(Hy)} strain. Under these conditions levanase is overproduced and it is then possible to compare its secretion with that of \(\alpha\)-amylase and levanosucrase in the same genetic context and under the same growth conditions.

**METHODS**

**Strains and media.** The \textit{Bacillus subtilis} strain GM96201 contained a \textit{sacR-sacC} fusion which was introduced into the chromosome of strain GM96100, as previously described (Leloup et al., 1997). Bacteria were grown at 37°C in minimal medium (Chambert & Petit-Glatron, 1984) supplemented with 1% (w/v) glucose and 0.5 mM CaCl2. Strains and plasmids are listed in Table 1.

**Plasmid and DNA manipulation.** The DNA fragments carrying \textit{sacC}, the levanosucrase structural gene including its signal sequence, or \textit{sacR}, the \textit{sacB} (levanosucrase structural gene) leader region, were amplified by PCR (Petit-Glatron & Chambert, 1992) with the chromosomal DNA of strain QB112 or pLS50 (Steinmetz et al., 1985) as templates and A and B or C and D oligonucleotides for \textit{sacC} and \textit{sacR}, respectively: A, 5’ AAGGAGACGTCAACGATG 3’ (forward primer), and B, 5’ AAAGGCCTGCAGAACACCTGATGATTTGG 3’ (reverse primer), with \textit{SacI} and \textit{SacII} sites (shown in bold) in A and B, respectively; C, 5’ CGCGGATCCCTTTTTTACCCCATCAC- ATATAACCTG 3’ (forward primer), and D, 5’ CATCCTT-GACGTCTCCTT 3’ (reverse primer), with \textit{BamHI} and \textit{AatII} sites (in bold) at the 5’ end of C and D, respectively.

The restriction sites were included in the primers to facilitate subsequent in-frame fusion of the regulatory region and the coding sequence, and insertion into the appropriate vector. The amplified fragments were purified by electroelution after electrophoresis on an agarose gel. The fragments were then blunt-ended with \textit{Pfu} DNA polymerase treatment and inserted into the \textit{SvaI} site of the pCR (+) vector according to the supplier’s recommendations (Stratagene), resulting in plasmids pGMC9 and pGMC10. The \textit{BamHI}–\textit{AatII} fragment (\textit{sacR}) of pGMC9 was purified and ligated into pGMC10 (containing the \textit{sacC} fragment) digested with the same endonucleases. The resulting plasmid was used to transform \textit{Escherichia coli} TG1. In these plasmids, which were purified from \textit{E. coli} transformants and exhibited fragments of the expected size after digestion by various endonucleases, we verified the in-frame fusion between the two DNA fragments by sequencing the amplified fragments, using the Sequenase kit (USB) and double-stranded plasmid DNA. The presence of active levanase was assayed in the cell extracts of \textit{E. coli} transformants. One appropriate plasmid, pGMC11, was selected and used for construction of strain GM96201.

**Enzyme assay.** Activity of levanase was assayed in the presence of uniformly labelled \([^{14}C]\)levan in 0.1 M sodium phosphate, pH 6, at 37°C. \([^{14}C]\)Fructose released from the levan hydrolysis was quantitatively analysed by TLC on silica gel TLC foils (Schleicher & Schuell) developed in butan-1-ol/propan-2-ol/water (3:12:4, by vol). Uniformly labelled \([^{14}C]\)levan was obtained from the action of immobilized levanosucrase on \([^{14}C]\)sucrose (Chambert & Petit-Glatron, 1993). One unit of enzyme activity was defined as the amount of enzyme that releases one \(\mu\)mol fructose min\(^{-1}\) in the presence of 10 mg labelled levan ml\(^{-1}\) (which corresponds to 0.06 M fructosyl unit). One enzyme unit corresponds to 8.5 \(\mu\)g pure levanase.

**Gel electrophoresis and immunoblotting.** Proteins were analysed by 10% (w/v) SDS-PAGE and the cell-associated

<table>
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<th>Table 1. Strains and plasmids</th>
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<td><strong>Strains</strong></td>
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<td>QB112</td>
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<td>GM96100</td>
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<td>GM96201</td>
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<tr>
<th><strong>Plasmids</strong></th>
<th><strong>Relevant genotype and phenotype</strong></th>
<th><strong>Source/reference</strong></th>
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<tr>
<td>pLS50</td>
<td>\textit{B. subtilis sacR–sacC} inserted into \textit{pJH101} (Cm(^R))</td>
<td>Steinmetz et al. (1985)</td>
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<tr>
<td>pGMC50</td>
<td>\textit{pLS50} (Km(^R))</td>
<td>Petit-Glatron &amp; Chambert (1992)</td>
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<tr>
<td>pGMC64</td>
<td>\textit{pGMC50} deleted of the \textit{BamHI}–\textit{EcoRV} fragment and with the \textit{sacR–sacC} fusion (\textit{BamHI}–\textit{Stul}) ligated at these sites</td>
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<td>pGMC9</td>
<td>\textit{pCR(+) [pBluescript II SK(+) sacR}</td>
<td>This work</td>
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<tr>
<td>pGMC10</td>
<td>\textit{pCR(+) [pBluescript II SK(+) sacC}</td>
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<tr>
<td>pGMC11</td>
<td>\textit{pCR(+) sacR–sacC}</td>
<td>This work</td>
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\(*\) This construct was created by Campbell-like integration.

\(^\dagger\) \textit{pJH101} derived from plasmid pBR322 by insertion of the 1 kb \textit{cat} gene from pC194 into the \textit{PvuII} site (Ferrari et al., 1983).
forms of levanase were analysed by immunoblotting as described previously (Petit-Glatron et al., 1987).

**Pulse-chase experiment.** Under the usual conditions of growth, *B. subtilis* cells were induced with sucrose (60 mM final concentration) and pulse-labelled at an OD$_{600}$ of 2 by adding 0.25 mCi (9 MBq) $^{[35]}$Smethionine (800 Ci mmol$^{-1}$) to 1 ml culture suspension maintained at 37 °C. After a pulse period of 30 s, non-radioactive methionine (4 mM final concentration) was added. Samples of 0.2 ml were withdrawn at intervals and all reactions were immediately stopped by diluting the samples threefold with ice-cold stopping buffer (0.1 M sodium phosphate, pH 7, containing 2.4 M KCl, 200 μg chloramphenicol ml$^{-1}$ and 0.2 mM PMSF). Cell suspensions were centrifuged and the supernatants were dialysed overnight at 4 °C against TNE buffer (50 mM Tris/HCl, pH 8, containing 150 mM NaCl and 5 mM EDTA) and diluted fivefold in TNET (TNE buffer containing 1%, v/v, Triton X-100). The bacterial pellets were washed with ice-cold stopping buffer without KCl, resuspended in 0.3 ml TNES (TNE buffer containing 2%, w/v, SDS) and diluted fivefold in TNET. Cells were disrupted by sonication. The suspensions were incubated for 5 min at 95 °C. Antibodies against *B. subtilis* levanase (20 μl) and 10% (w/v) Protein-A-Sepharose (80 μl) (Sigma) in TNET were then added to 0.3 ml dialysed supernatants and to the disrupted cells. After overnight incubation at 4 °C, the immunoprecipitates were recovered by centrifugation. The pellets were washed three times with 1 ml TNET and finally resuspended in electrophoresis sample buffer. The samples were boiled for 3 min and analysed by SDS-PAGE.

**Purification of levanase.** Exocellular levanase was purified from the supernatant of strain GM96201 harvested at the end of the exponential phase of growth. Supernatant was first concentrated 20-fold on a bioconcentrator miniplate (Millipore), then the enzyme was obtained in a pure state after the successive chromatographic steps, as follows.

**Step 1:** hydroxylapatite (HA) chromatography. The supernatant, concentrated from 31 culture, was loaded on a HA column (15 x 1 cm) pre-equilibrated with 0.1 M sodium phosphate, pH 6.5.

The column was washed with the same buffer, and proteins were eluted with a linear gradient of sodium phosphate from 0.1 to 0.8 M. Levanase activity was eluted by 0.5 M phosphate. Fractions containing activity were pooled, dialysed against 0.1 M sodium phosphate, pH 6.5, and re-concentrated up to 2 ml with vivaspin (Vivascience).

**Step 2:** gel-filtration chromatography. We ran gel-filtration chromatography using a 70 x 2.5 cm Biogel P150 column (200–400 mesh; Bio-Rad) with continuous elution at a constant rate of 45 ml h$^{-1}$ at 5 °C with 0.1 M sodium phosphate, pH 6. Fractions containing levanase activity were pooled and dialysed against 0.1 M sodium phosphate, pH 7.

**Step 3:** DEAE-Sepharose chromatography. The dialysed protein solution was loaded on a DEAE-Sepharose anion-exchange column (20 x 1 cm) pre-equilibrated with 0.1 M sodium phosphate, pH 7. The column was washed with the same buffer, and proteins were eluted with a linear gradient of 0–0.6 M NaCl. Levanase activity was eluted by 0.4 M NaCl.

Purified levanase migrated as a single protein band on 10% (w/v) SDS-PAGE and had an apparent molecular mass of 73 000 Da.

**Antibodies against levanase.** Antibodies raised in rabbits were prepared against pure levanase.

**RESULTS**

**Construction of strain GM96201 overproducing levanase during the exponential phase of growth**

The BamHI–Stul fragment of pGMC11 carrying the fusion of P$_{saeC}$ to the sacC structural gene was ligated between the BamHI/EcoRV sites in pGMK50 (Petit-Glatron & Chambert, 1992) to give pGMK64, an integrative plasmid in which the sacC gene is under the

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Fig. 1. Production of exocellular levanase in strain GM96201 during the exponential phase of growth. Bacteria from strain GM96201 induced (C) or not (●) with 60 mM sucrose were grown in minimal medium supplemented with 1% (w/v) glucose at 37 °C. Samples were withdrawn at intervals, their OD$_{600}$ was measured and then they were centrifuged. (a) Supernatants were dialysed against 0.1 M sodium phosphate, pH 6, and assayed for activity. (b) At an OD$_{600}$ of 2, a 1 ml aliquot of the induced culture suspension was pulse-labelled for 45 s with 0.9 MBq ($^{35}$S)methionine (800 Ci mmol$^{-1}$) and chased with a large excess of non-radioactive methionine (4 mM final concentration). Samples were removed at intervals and centrifuged. Supernatants were analysed by SDS-PAGE.
control of the sacB leader region. pGMK64 was integrated by a Campbell-like mechanism into the chromosome of GM96100 (Leloup et al., 1997). The transformants were selected from LB plates containing the appropriate antibiotic. We confirmed the presence of the sacR–sacC fusion in the transformants by PCR using appropriate oligonucleotides. One of the transformants containing the \( P_{\text{sacB}}-sacC \) fusion and exhibiting sucrose-inducible expression of levansucrase was chosen for further analysis and named GM96201.

When strain GM96201 was grown in the absence of sucrose, no levansucrase activity (Fig. 1a) or labelled levansucrase (pulse experiment, not shown) was detected in the supernatant. In the presence of sucrose, production of exo-levansucrase occurred at a constant differential rate of synthesis, 5 \( \mu \)g mg\(^{-1}\) cell dry weight h\(^{-1}\), during the exponential phase of growth. Pulse experiments (Fig. 1b) showed that levansucrase is one of the major proteins released into the culture supernatant. The molecular mass of the protein was 73 kDa as expected from the nucleotide sequence of the structural gene \( sacC \) (Schörgendorfer et al., 1987). We observed that the production of exo-levansucrase is around ten times lower than that of levansucrase (Chambert & Petit-Glatron, 1984). This point will be discussed later.

**Fig. 2.** Pulse labelling experiment. Bacteria of strain GM96201 were induced with 60 mM sucrose in the exponential phase of growth. At an OD\(_{600}\) of 2, a 1 ml aliquot of the culture was pulse-labelled for 45 s with 0.9 MBq \([^{35}S]\)methionine (800 Ci mmol\(^{-1}\)) and chased with a large excess of non-radioactive methionine (4 mM final concentration). Samples were removed at intervals, treated and analysed as described in Methods. (a) SDS-PAGE of labelled cell-associated and exocellular levansucrase. (b) At each time point, exocellular (O) or cell-associated (●) levansucrase is shown as a fraction of total levansucrase (exocellular + cell-associated). Quantification was performed with a Phosphor-Imager.

**Fig. 3.** Kinetics of the unfolding–refolding transition of \( B.\) subtilis levansucrase measured by resistance to proteolysis at 37 °C, 0.1 M sodium phosphate, at pH 7. Unfolding was promoted (arrow 1) by mixing 20 \( \mu \)l stock solution of purified levansucrase (1.5 mg ml\(^{-1}\) with 150 \( \mu \)l 10 M urea, pH 7. Samples (8 \( \mu \)l) were withdrawn at the times indicated and quickly mixed with 92 \( \mu \)l 0.1 M sodium phosphate, pH 7, containing 10 \( \mu \)g subtilisin ml\(^{-1}\). Refolding was initiated (arrow 2) after 30 min unfolding by mixing 40 \( \mu \)l unfolding mixture with 460 \( \mu \)l 0.1 M sodium phosphate, pH 7, containing 0.5 mM calcium. Samples (100 \( \mu \)l) were withdrawn at the times indicated and quickly mixed with 2 \( \mu \)l of a 0.5 mg subtilisin ml\(^{-1}\) solution, and incubated for 5 min. PMSF was then added to the samples. Aliquots (80 \( \mu \)l) of all the samples were subjected to SDS-PAGE. Aliquots (5 \( \mu \)l) were analysed for levansucrase activity. (a) SDS-PAGE analysis of the samples. (b) Quantification of the percentage of the folded protein was obtained from densitometric tracings of the Coomassie-blue-stained gels using the NIH Image program (O) and from measurements of the residual enzyme activity (●).

**Sequential stages of the levansucrase secretion process.** We characterized the discrete stages of the levansucrase secretion pathway from pulse–chase experiments (Fig. 2). We did not detect any unprocessed precursor of levansucrase whose expected molecular mass is 76 kDa (deduced from the nucleotide sequence). However, a cell-associated form with the same molecular mass as the exocellular protein (73 kDa) disappeared slowly from the cells and was concomitantly released into the culture supernatant. The \( t_{1/2} \) of this event was evaluated at 80 s ± 10 s and the yield of the release was almost 100%. Thus, this processed cell-associated levansucrase displays the features of a transient secretion intermediate of the protein. This precursor was tightly bound to the cells since it was not released after washing the cells at a high ionic strength (see Methods). We have previously characterized a very similar intermediate in the secretion process of levansucrase (Petit-Glatron et al., 1987) and
Kinetics of secretion of *Bacillus subtilis* levanase

73 kDa

![Graph](image)

**Fig. 4.** Unfolding-refolding transition of levanase mediated by Ca$^{2+}$ at 37 °C, pH 7. Stock solution (15 μl) of purified protein (1.5 mg ml$^{-1}$) was preincubated for 30 min in 105 μl 10 M urea in 0.1 M sodium acetate at pH 7. Refolding was initiated by mixing 60 μl of the unfolding mixture with 700 μl 0.1 M sodium acetate, pH 7, containing or not 20 mM calcium chloride. Samples (90 μl) were withdrawn at the time indicated and quickly mixed with 2 μl 1 mg subtilisin ml$^{-1}$ solution. Samples were submitted to SDS-PAGE analysis, and were assayed for levanase activity. (a, b) SDS-PAGE analysis of the samples of the refolding experiment (a) in the absence of calcium and (b) in the presence of calcium. (c) Curves obtained from scanning the coloured bands of the SDS gels. (d) Residual enzyme activities were measured in the presence of the same concentration of Ca$^{2+}$ (10 mM final concentration). ○, Refolding in the absence of calcium; ●, refolding in the presence of calcium.

α-amylase (Leloup *et al.*, 1997) when these proteins were overproduced. We pointed out that the time constant of the processed precursor release of these two proteins was of the same order of magnitude as the time constants of *in vitro* refolding of the proteins occurring under the same conditions of pH, temperature and ionic strength. We tested whether a similar correlation could be observed with levanase.

**Kinetics of unfolding–refolding of levanase *in vitro*: calcium is a modulator of the folding reaction**

The unfolding–folding transition was measured by monitoring the appearance and disappearance of the subtilisin sensitivity of the protein (Fig. 3). In the presence of 8.75 M urea, at pH 7 and 37 °C, unfolding occurred with a $t_{1/2}$ of 7 ± 1 min. After dilution of the denaturing agent in 0.1 M sodium phosphate, pH 7, at 37 °C, the unfolding–folding transition occurred with a high yield ($Y = 80\%$) and the $t_{1/2}$ of the reaction was 46 ± 5 s. The rate of levanase refolding is therefore approximately the same as that of *in vivo* release of the protein under the same conditions of pH, temperature and ionic strength. Similar observations have been made for levansucrase (Chambert *et al.*, 1995) and α-amylase (Leloup *et al.*, 1997). We proposed that the release and folding processes are coupled on the external side of the cytoplasmic membrane. Moreover, we identified the calcium ion as being a folding modulator that can assist such a coupling (Petit-Glatron *et al.*, 1993). In the case of levanase, we observed that the presence of calcium increases the rate and yield of refolding (Fig. 4). We examined the ability of levanase to bind calcium in comparison to other secreted proteins by the method of Maruyama *et al.* (1984) using a dot blot assay with pure proteins (Fig. 5). The radioactive spots were quantified by phosphor-imaging. The ability of levanase to bind calcium was approximately the same as that of levansucrase. Therefore, we can reasonably conclude that levanase, α-amylase and levansucrase have some common folding properties: calcium increased the rate of refolding of each protein; however, the affinities of the native proteins to calcium are very low. The folding catalyst effect of calcium could play an important role in the final stage of their secretion processes.

**DISCUSSION**

The expression of sacC under the control of the leader region of levansucrase led to an overproduction of exocellular levanase during the exponential phase of growth of *B. subtilis*. Pulse-chase experiments showed some differences in the timing of the cellular sequential events leading to secretion compared with those of levansucrase. In contrast to levansucrase, we observed no unprocessed intermediates in the sequential process of levanase secretion, suggesting that the precursor processing is very rapid. The same result was obtained...
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...for 15 min at room temperature in 0.1 M potassium acetate or 0.1 M potassium phosphate, pH 7, respectively. Then each piece of membrane was incubated for 1 h in 1 ml of the same buffer containing 10 μCi (370 kBq) 45CaCl₂. After 20 min washing in a large volume of buffer, the membranes were dried, exposed to a Kodak X-Omat film and quantified by phosphor-imaging (c).

For the expression of heterologous proteins this difference was generally explained by post-transcriptional events such as the inability of the secretory apparatus to recognize foreign proteins, the protease sensitivity of these proteins or competition for the secretion apparatus. We have demonstrated unambiguously that, for the expression of homologous α-amylase and levanase, none of these explanations applies because the proteins adopt in vitro a proteolytically insensitive structure at approximately the same rate as levansucrase and their levels of production remain the same in strains deleted or not for the levansucrase structural gene. Presumably the difference in the production level is mostly due to differences in transcription or translational efficiency. Preliminary experiments have indicated that the level of α-amylase and levanase mRNAs in fully induced strains correlates to some extent with the secretion process, i.e. the release of the processed protein. Levanase is the third model protein for which we have observed that the half-life of such an event is similar to the half-life of in vitro refolding of the protein occurring under the same conditions of pH, temperature and ionic strength. Furthermore, these two events occur within the same time range (t₁/₂ = 1–2 min) for the three proteins despite their large molecular mass difference. These results suggest that the rate of the release stage, which includes the crossing of the cell wall, is not dependent on protein size as it was previously postulated (Demchick & Koch, 1996), but rather dependent on their folding rate. The final point of interest concerns the difference in the levels of synthesis of levanase and levansucrase even though the two structural genes were under the same transcriptional control. Levanase synthesis is ten times lower than that of levansucrase. Similar results have been reported for α-amylase, whose level of synthesis is about five times lower (Leloup et al., 1997), and also for other heterologous model proteins expressed under the regulatable control of the levansucrase leader region (Simonen & Palva, 1993).

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