Expression of *Bacillus subtilis* neutral protease gene (*nprE*) in *Saccharomyces cerevisiae*

LIN-Fa Wang† and Rodney J. Devenish*

Department of Biochemistry, and the Centre for Molecular Biology and Medicine, Monash University, Clayton, Victoria 3168, Australia

(Received 10 August 1992; accepted 19 October 1992)

Expression in the yeast *Saccharomyces cerevisiae* of the intact *nprE* gene of *Bacillus subtilis*, which encodes the pre-pro-NprE neutral protease precursor, resulted in intracellular accumulation of unprocessed precursor without detectable secretion or processing of the expressed gene product. When sequences specifying the signal peptide of yeast invertase were fused upstream of sequences encoding the mature NprE enzyme, *nprE* gene products were secreted into the culture medium. The secreted protein products were, however, highly glycosylated and biologically inactive.

Introduction

The α-amylase genes from various sources (Filho et al., 1986; Sato et al., 1986; Rothstein et al., 1987), including *Bacillus* (Ruoheinen et al., 1987; Nonato & Shishido, 1988; Kovaleva et al., 1989), have been successfully expressed in *Saccharomyces cerevisiae*, but no work has yet been reported on the expression and secretion of heterologous, especially bacterial, protease genes in yeast. *B. subtilis* secretes a large number of extracellular proteases during the late phase of growth (Priest, 1977). All share a common structural feature, a pre-peptide located between the coding regions for signal peptide (or pre-peptide) and the mature enzyme (Stahl & Ferrari, 1984; Yang et al., 1984; Bruckner et al., 1990; Rufo et al., 1990). The neutral protease (NprE) of *B. subtilis* accounts for more than 70% of total activity of all extracellular proteases, and has the largest pre-peptide of 197 amino acid residues (Yang et al., 1984).

We have recently demonstrated that the *B. subtilis* neutral protease gene (*nprE*) can be used as a reporter gene with applications in molecular biology and biotechnology (Wu et al., 1991). By removing its original ribosome binding site (RBS), we have also achieved expression of the gene in *Escherichia coli* such that protease is secreted from cells (Wang et al., 1990). In this communication, we report our studies on the expression and secretion of the *B. subtilis* *nprE* gene in yeast.

Methods

Yeast strain 334 MATa pep4-3 prb1-1122 ura3-52 leu2-3, 112 reg1-501 gal1 (Hovland et al., 1989) and plasmid pLF1 (Nagley et al., 1988) were used for this study. The PCR primers used to engineer the recombinant genes are listed in Table 1. A wild-type *nprE* gene cassette lacking the native RBS was obtained by PCR amplification using primers P1 and P2 as described previously (Wang et al., 1990), and subcloned into the unique *BglII* site of pLF1 to form expression vector pLF-wtNPR (Fig. 1). Similarly, the Sphi-BamHI fragment encoding the mature NprE protease was obtained by PCR amplification with primers P2 and P3, and then joined to a BamHI-Sphi fragment encoding the yeast invertase signal peptide (Carlson & Botstein, 1982) obtained by direct PCR amplification of yeast chromosomal DNA with primers P4 and P5. The hybrid gene cassette was cloned into pLF1 to form pLF-ImNPR (Fig. 1).

Yeast transformation of plasmid DNA was carried out essentially as described by Klebe et al. (1983). For expression studies, yeast cells were grown at 30°C in liquid YEPD medium for 48 h. The culture was separated into cell pellet and culture medium by centrifugation using a Sorvall SS-34 rotor at 10000 r.p.m. for 10 min at 4°C. The cell pellet was resuspended in 7 M-guanidine hydrochloride and then vortexed in the presence of glass beads to produce a total cell lysate. Portions of lysate equivalent to the cell pellet from 300 μl of the original culture were mixed with an equal volume of 2× Laemmli sample buffer and then subjected to SDS-PAGE (10%, w/v) analysis. Culture medium aliquots (300 μl) were concentrated by precipitation with acetone before analysis by SDS-PAGE (10%, w/v).

Protease assays and immunoblots were carried out as described previously (Wang et al., 1990; Wu et al., 1991). Endoglycosidase D and endo-α-N-acetylgalactosaminidase were purchased from Boehringer Mannheim Biochemica, and were used according to the supplier's instructions; 300 μl culture medium fractions were incubated with 2 μl...
Table 1. PCR oligonucleotide primers

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5’ → 3’)</th>
<th>Position*</th>
<th>Target gene†</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1</td>
<td>CCATGGATCCAAATATAA TGGTTAGGTAAG</td>
<td>2-16</td>
<td>NPR</td>
</tr>
<tr>
<td>P2</td>
<td>GCCTGGATCCCTTTTATGCAA</td>
<td>1676-1700</td>
<td>NPR (c)</td>
</tr>
<tr>
<td>P3</td>
<td>CAAAATAAGTAGCATGGCGCCGC CGCCAC</td>
<td>645-673</td>
<td>NPR</td>
</tr>
<tr>
<td>P4</td>
<td>TCACCAAGATGATGCTTTTGCAAGC</td>
<td>1-17</td>
<td>IVT</td>
</tr>
<tr>
<td>P5</td>
<td>TTGTATGCAGATTTTTGGCTGC</td>
<td>39-67</td>
<td>IVT (c)</td>
</tr>
</tbody>
</table>

* Nucleotide A of the ATG translational initiation codon is designated residue number 1.
† NPR, gene for B. subtilis neutral protease; IVT, gene for yeast invertase; c, complementary strand.

Results and Discussion

Extracellular proteases, like extracellular amylases, are a class of enzymes with great potential for use as reporter enzymes in studies of gene regulation and protein secretion. By comparison with methods used to assay amylases, those for protease detection are much easier, and can be carried out in situ without the necessity for staining procedures (Wang & Doi, 1987; Wu et al., 1991).

Encouraged by the direct expression and secretion of B. subtilis α-amylases in S. cerevisiae (Ruohonen et al., 1987; Nonato & Shisido, 1988; Kovaleva et al., 1989) and by our recent demonstration of direct expression and secretion of B. subtilis neutral protease in E. coli (Wang et al., 1990) we began to test the feasibility of using the B. subtilis nprE gene in yeast as a reporter system. The structures of the two expression plasmids constructed in the course of this study are illustrated in Fig. 1. The sequence integrity and functionality of the PCR-derived nprE gene cassette was tested by secretion of functional protease from E. coli as described in our previous paper (Wang et al., 1990) before it was inserted into the vector pLF1 for expression studies in yeast. The resulting recombinant vector, pLF-wtNPR, was introduced into yeast strain 334 by PEG transformation (Klebe et al., 1987) and its presence in the resulting transformants was confirmed by PCR analysis using primers P1 and P2 with total yeast DNA as template (data not shown).

Liquid YEPD cultures were set up as described in Methods to study the expression and secretion of the nprE gene in yeast. When the culture medium from recombinant yeast strain 334[pLF-wtNPR] was analysed for protease production using either plate or liquid assay methods (Wang & Doi, 1987; Wang et al., 1990) no significant activity was detected over the control recombinant strain 334[pLF1]. This result could have been due to any one of the following reasons: (i) failure to

\[
\text{enzyme (1 U µl}^{-1}\text{) at 37 °C for 30 min in the presence of 0.1% SDS, before being precipitated with 4 vols cold acetone and analysis by immunoblotting.}
\]
Fig. 2. Test for intracellular expression of the nprE gene in yeast made by Western blot analysis on 10% (w/v) SDS-PAGE. Lanes 1 to 3 display samples of total cell lysate prepared from yeast strains 334, 334[pLF1], and 334[pLF-wtNPR], respectively. Positions of molecular weight mass markers (Rainbow, Amersham) run on the same gel are shown on the left.

express the nprE gene; (ii) the nprE gene was expressed, but the product not secreted; (iii) the gene product was expressed and secreted but was inactive owing to post-translational modification or incorrect processing of the pre-pro-enzyme. To distinguish between these possible causes, immunoblot analysis was carried out using affinity-purified rabbit anti-NprE sera (Wang et al., 1990; Wu et al., 1991). As shown in Fig. 2, when the intracellular fraction (i.e. total cell lysate) was analysed, a specific band of about 65 kDa was detected only for the strain carrying the functional nprE gene (lane 3), while the lower faint band (around 42 kDa) appeared in all lanes, indicating its probable identity as a cross-reactive protein originating from the yeast host cells. The size of the NprE-related polypeptide expressed from strain 334[pLF-wtNPR] corresponds to that predicted for the unprocessed pre-pro-NprE polypeptide (Yang et al., 1984). A similar immunoblot was also performed for extracellular fractions (i.e. culture medium), but no positive signal was detected for cells carrying pLF-wtNPR (data not shown). These results would suggest that the nprE gene was expressed in yeast under the control of the PGK1 promoter, but that the native signal peptide failed to function as a targeting signal to direct the gene product into the yeast secretory pathway. Supporting evidence includes, first, the absence of any NprE-related polypeptide in the culture medium, and secondly, the presence of a relatively sharp band of the unprocessed pre-pro-NprE inside the cell. If the signal peptide of the B. subtilis neutral protease was able to

Fig. 3. Western blot analysis of nprE gene products expressed from strain 334[pLF-ImNPR]. Lanes 1 and 2: total cell lysate from strains 334[pLF-ImNPR] and 334[pLF1], respectively. Lane 3: positive control of mature NprE protease expressed from B. subtilis. Lanes 4 and 5: culture medium from strains 334[pLF-ImNPR] and 334[pLF1], respectively.

function in the initial phases of the secretion process, its gene product may still have remained inside the cell, but the polypeptide would most likely have been glycosylated (see Ruohonen et al., 1987 and later discussion in this paper), which would have resulted in the diffusion or smearing of the signals on Western blots.

The failure to achieve secretion may also have resulted from sequence features of the polypeptide rather than its signal peptide. To examine further the extent to which the mature neutral protease enzyme could be secreted from yeast, plasmid pLF-ImNPR was constructed by direct fusion of the coding sequence of the yeast invertase signal peptide immediately upstream of sequences encoding the mature NprE protease via PCR-mediated gene construction (see Methods for details). The sequence integrity of this PCR-derived hybrid gene was confirmed by direct DNA sequencing (data not shown) before the gene cassette was used in yeast expression studies.

When this hybrid gene was expressed in yeast, no protease activity was detected. However, when the intracellular and extracellular fractions were analysed by immunoblotting, it was found that the majority of the NprE-related polypeptides were present in the culture medium, i.e. they had been secreted (Fig. 3). In addition, instead of forming a sharp band on Western blots as observed for the intracellular pre-pro-NprE expressed from pLF-wtNPR, the signals obtained for both intracellular and extracellular NprE-polypeptides showed, in this case, extensive diffusion and smearing, indicating a high level of glycosylation of the nprE gene products. The major diffused band present in the intracellular
fraction (lane 1) had a molecular mass of about 45 kDa, which is bigger than the unmodified mature neutral protease (about 38 kDa, lane 3) or the unmodified pre-NprE with the invertase signal peptide attached (about 40 kDa). These results indicated that the secretion process was very efficient, with all of the gene products either being secreted or having at least entered the secretion pathway as indicated by the absence of unmodified nascent pre-NprE polypeptides of the same mobility as seen in the pLF-wtNPR expression experiment (Fig. 2). These results support the contention that the failure of secretion of the native pre-pro-NprE in yeast most likely results from the failure of the native signal peptide to function in yeast rather than any incompatibility of the NprE protease with the yeast secretory pathway.

There are three potential N-linked glycosylation sites in the mature NprE enzyme, Asn–Ile–Ser (residues 18–20), Asn–Glu–Ser (residues 166–168), and Asn–Asp–Thr (residues 177–179). The Ser and Thr contents of the NprE protease are 10.0 ± 0.5 and 9.5 ± 0.5, respectively. These features suggest that it is potentially a good substrate for N- and O-linked glycosylation. To confirm that the nprE gene products expressed from pLF-ImNPR were actually glycosylated, supernatant samples were treated with commercially available endoglycosidases. Sharpening of the Western blot signals would have indicated degradation of the carbohydrate moieties attached to the NprE proteins. The results obtained after the treatment of supernatant fractions with endoglycosidase D are shown in Fig. 4. As indicated by arrows, several bands appeared in the digested sample (lane 2) which were absent in the control sample (lane 1). No sharp band was observed in the region around 38 kDa (the size predicted for the unmodified NprE protease), suggesting either that the endoglycosidase D digestion was incomplete or that more than one type of glycosylation was present on the NprE protein or both. Interestingly, digestion of samples with endo-α-N-acetylgalactosaminidase had no effect on the mobility of NprE proteins as determined by Western blot (data not shown).

In conclusion, the results presented in this communication illustrated two important points: (i) different signal peptides, derived from the same organism (Bacillus), may behave very differently in a heterologous system (yeast); and (ii) in contrast to the B. amyloliquefaciens amylase, whose glycosylated form was still active (Ruohonen et al., 1987; Kovaleva et al., 1989), the glycosylated B. subtilis neutral protease secreted from yeast is biologically inactive.

We thank Dr R. H. Doi for providing anti-neutral protease sera, Dr R. A. Sclafani for providing yeast strain 334 and Professors P. Nagley and A. W. Linnane for providing us with a stimulating research environment in which to carry out this work. This study was supported in part by a Monash University Special Research Fund Grant (19.122.032) to L.E.W. and ARC Grant (20.122.029) to R.J.D. and L.F.W.

References


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