The Response of Lactococcus lactis to Membrane Protein Production

Ravi K. R. Marreddy, Joao P. C. Pinto, Justina C. Wolters, Eric R. Geertsma, Fabrizia Fusetti, Hjalmar P. Permentier, Oscar P. Kuipers, Jan Kok, Bert Poolman

1 Department of Biochemistry, Groningen Biomolecular Sciences and Biotechnology Institute, Netherlands Proteomics Centre & Zernike Institute for Advanced Materials University of Groningen, Groningen, The Netherlands, 2 Department of Molecular Genetics, Groningen Biomolecular Sciences and Biotechnology Institute, University of Groningen, Groningen, The Netherlands

Abstract

Background: The biogenesis of membrane proteins is more complex than that of water-soluble proteins, and recombinant expression of membrane proteins in functional form and in amounts high enough for structural and functional studies is often problematic. To better engineer cells towards efficient protein production, we set out to understand and compare the cellular consequences of the overproduction of both classes of proteins in Lactococcus lactis, employing a combined proteomics and transcriptomics approach.

Methodology and Findings: Highly overproduced and poorly expressed membrane proteins both resulted in severe growth defects, whereas amplified levels of a soluble substrate receptor had no effect. In addition, membrane protein overproduction evoked a general stress response (upregulation of various chaperones and proteases), which is probably due to accumulation of misfolded protein. Notably, upon the expression of membrane proteins a cell envelope stress response, controlled by the two-component regulatory CesSR system, was observed.

Conclusions: The physiological response of L. lactis to the overproduction of several membrane proteins was determined and compared to that of a soluble protein, thus offering better understanding of the bottlenecks related to membrane protein production and valuable knowledge for subsequent strain engineering.

Introduction

Membrane proteins are involved in many essential cellular processes such as transport of nutrients, sensing of environmental changes, energy transduction and scaffolding of cell structure. Due to their important roles in various diseases these proteins are clinically important as potential drug targets. To date, 60% of all available pharmaceutical drugs target membrane proteins [1]. Even though 20 to 30% of all genes encode integral membrane proteins (IMPs) [2], the structures of relatively few of these proteins have been elucidated at high resolution. Expression hosts such as Escherichia coli, yeast (Pichia pastoris and Saccharomyces cerevisiae) or higher eukaryotic cells (mammalian and insect cells) are often used for membrane protein production [3,4]. However, the production of proteins in a functional state and in sufficient yields for structural analysis is often a problem. Emerging systems like cell-free protein expression offer interesting possibilities but producing the protein in the native state is still a bottleneck [5]. Over the past decade, the Gram-positive bacterium Lactococcus lactis has emerged as an alternative host for membrane protein production [6–12]. L. lactis is amenable to genetic manipulation and well-tunable promoter systems are available [9,13,14]. The organism shows a limited proteolytic activity and, as a Gram-positive bacterium, contains a single membrane with a high fraction of glycolipids. The easier targeting of proteins to the single (cytoplasmic) membrane, compared to Gram-negative bacteria, facilitates activator/inhibitor studies of expressed proteins [12,15]. In addition, the limited number of endogenous transporters simplifies complementation studies.

The choice of host cells for production of recombinant membrane protein depends on various factors, such as gene source (codon bias, tRNA levels), protein complexity and the requirements for a particular folding environment, post-translational modifications, and others. Production of proteins can often be improved by trial-and-error approaches to screen for the best promoter, inducer levels and growth media [16,17] or by screening a wide variety of homologues [18,19]. Alternatively, production levels can be increased by selecting strains with limited proteolytic activity and, as a Gram-positive bacterium, is amenable to genetic manipulation and well-tunable promoter systems are available [9,13,14]. The organism shows a limited proteolytic activity and, as a Gram-positive bacterium, contains a single membrane with a high fraction of glycolipids. The easier targeting of proteins to the single (cytoplasmic) membrane, compared to Gram-negative bacteria, facilitates activator/inhibitor studies of expressed proteins [12,15]. In addition, the limited number of endogenous transporters simplifies complementation studies.

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improved protein production potential [20–22] or by screening for stable variants of a given protein [23]. To understand and ultimately alleviate the bottlenecks in membrane protein production, one needs to comprehend the response of the host cells, as has been done for yeast [24] and E. coli [25]. This knowledge leads to a better understanding of the cellular bottlenecks affecting the production of membrane proteins and hints towards strategies to engineer strains [26,27].

Here, we used transcriptomics and proteomics approaches to determine the response of *L. lactis* to the production of membrane proteins. We evaluated the effects of (over)expression of diverse proteins, including the osmoregulatory ABC transporter OpuA, the plant sucrose transporter StSUT1 and the human γ-secretase component PS1Δ9; these proteins display different levels of expression and compromise growth to different extents. We evaluated the expression of these membrane proteins against the water-soluble substrate receptor OpuAC. (Over)expression of the diverse membrane proteins elicited a similar response in *L. lactis*, which was distinct from the stress evoked by the production of OpuAC.

**Results**

**Recombinant protein production and cell growth**

The genes of the following proteins were cloned in the lactococcal nisin-inducible gene expression plasmid pNZ8048 [28] and introduced in *Lactococcus lactis* NZ9000: the ABC transporter OpuA from *L. lactis*, the sucrose transporter StSUT1 from *Solanum tuberosum*, the human PS1Δ9, a presenilin variant missing exon 9, and the soluble glycine betaine-binding protein OpuAC from *L. lactis*. Each of the proteins was engineered to contain a C-terminal hexa-histidine tag, facilitating their detection by immunoblotting. *L. lactis* NZ9000 containing the empty vector pNZ8048 was used as a control. All the physiological and omics analyses were carried out on biological triplicates, and the cells were grown in pH-controlled bioreactors. The cells were induced in the mid-exponential phase of growth (OD_{600}=0.5) with 10 ng/mL of nisin A. The growth rate (μ_{max}) of the cells prior to induction was 0.75±0.01. The addition of the inducer had an effect on growth, as determined in control cells carrying the empty vector and for OpuAC producing cells (the μ_{max} values were 0.60±0.06 and 0.59±0.02, respectively). A stronger effect on cell growth was observed in cells producing OpuA (μ_{max}=0.44±0.04 hr⁻¹) and StSUT1 (μ_{max}=0.35±0.01 hr⁻¹), while growth of cells producing PS1Δ9 (μ_{max}=0.19±0.03 hr⁻¹) was affected most severely (Figure 1A).

To determine protein production levels, membrane fractions extracted from these cells, harvested 2 h after induction, were analyzed by SDS-PAGE. Analysis of the Coomassie-stained gel (Figure 1B) and the immunoblot (Figure 1C) showed very high levels of OpuA. Similar to previous studies [29], the levels of the individual subunits of OpuA (OpuAA and OpuABC) were estimated to be >10% of total membrane protein (i.e. ~15 mg/L of cell culture). In contrast, the production levels of the erythrocystic membrane proteins PS1Δ9 and StSUT1 were at least an order of magnitude lower. Both these proteins were only detectable on the immunoblot (Figure 1C). OpuAC, the soluble substrate-binding protein of OpuA, was also overexpressed and detectable on the immunoblot (Figure 1C). OpuAC, the soluble water-soluble substrate receptor OpuAC. (Over)expression of the expression and compromise growth to different extents. We evaluated the expression of these membrane proteins against the water-soluble substrate receptor OpuAC. (Over)expression of the diverse membrane proteins elicited a similar response in *L. lactis*, which was distinct from the stress evoked by the production of OpuAC.

**Transcription and Translation Kinetics/Dynamics**

To optimize the sampling for the proteomics and transcriptomics studies, we determined the time-dependent profiles of transcription of *opuA* and a number of genes involved in the stress response (Figure 2D), in addition to the profile of synthesis of the OpuA protein (Figure 2E). Figure 2D shows that transcripts for the individual subunits of OpuA (OpuAA and OpuABC) can be readily detected after induction, rising quickly and reaching a plateau about 15 min after induction. The transcripts levels of cesR and dnuK, indicators of a specific and a general stress response in *L. lactis*, respectively [see below and accompanying paper [32]], were determined in parallel. The kinetics of formation of these

**Strains and constructs for “Omics” analyses**

To probe the basis for the difference in the production levels of the four proteins under study, we determined the physiological response of *L. lactis* to the protein synthesis burden by performing a combined proteomic and transcriptomic analysis. Contrary to PS1Δ9 and StSUT1, which have no known activity in *L. lactis* and/or of which the substrates are not present in our experimental setup, a high level of OpuA results in the accumulation of glycine betaine, thus producing an effect in itself. At high concentrations, glycine betaine can have a favorable effect on the stability/folding of proteins. To avoid such an effect, we expressed a point-mutated version of OpuA, OpuA(H223A), in which the histidine residue at position 223, essential for the ATPase activity [30], was replaced by alanine.

OpuA is easily produced by *L. lactis* NZ9000 and is therefore expected to request a considerable amount of resources for transcription as well as for translation and membrane targeting/insertion. To account for the putatively significant transcription burden of the genes encoding OpuA and OpuA(H223A), we also made pNZ4401 (NZ9000 derivative lacking the chromosomal *opuA* gene [31]) carrying the proper expression plasmids, confirmed that wildtype OpuA was functionally expressed and that OpuA(H223A) was inactive (Figure 2B). Induced expression of wildtype OpuA and OpuA(H223A) had similar negative effects on the growth of *L. lactis* (Figure 2C). Growth of the mRNA control strain *L. lactis* NZ9000 (pNZ4401mRNA) was only slightly inhibited upon induction with nisin A, and similar to that of the empty vector control strain *L. lactis* NZ9000 (pNZ4408). These data suggest that the nisin A-induced increase of transcription does not, per se, have a major impact on the physiology of *L. lactis*. Growth of the producer organism is inhibited only when either the functional or the inactive form of OpuA is produced at high levels.

Thus, to characterize the response of *L. lactis* to membrane protein production burden, proteomics and transcriptomics studies were performed with the *L. lactis* NZ9000 strains carrying pNZopuA(H223A), pNZPS1Δ9, pNZStSUT1 or pNZOpuAC. *L. lactis* NZ9000 (pNZopuA(H223A)) was used as a dedicated control for *L. lactis* NZ9000 (pNZopuA(H223A)), whereas *L. lactis* NZ9000 (pNZ4408) was used for the same purpose in the three other situations.

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transcripts was similar to that of opuA. Expression of the branched chain amino acid permease gene bcaP, which was used as a control, remained indeed constant throughout the growth curve [Brouwer et al., manuscript in preparation]. Although increased transcript levels could be detected after one min of induction, it took longer to observe the overproduced proteins by Western blotting (Figure 2E). OpuABC was detected after 4 to 8 min of induction and increased gradually over a period of 120 min. Therefore, sampling for transcriptomics was dense (0, 2, 8, 16, 32 and 64 min after induction), but less frequent for proteomics, (0 min, 16 min (onset of detectable OpuA) and 64 min (high accumulation of OpuA)). Time points later than 64 min were not examined as many indirect physiological phenomena were expected to obscure the (initial) response to the overproduction of the membrane proteins.

**Time-resolved stress response and experimental approach**

*L. lactis* was grown in pH-regulated bioreactors. Cell samples were collected in methanol at −40 °C to immediately quench further synthesis or breakdown of mRNA. For the proteomic studies, translation in harvested cells was stopped immediately by adding 100 μg/mL of chloramphenicol. Transcriptome data from biological triplicates was acquired by hybridizing the dual dye-labelled cDNA, obtained by reverse transcription of purified mRNA, onto SuperAmine glass slides spotted with duplicates of around 2500 ORF amplicons from *L. lactis* subsp. cremoris MG1363 [33]. Principal component analysis (PCA) of the transcriptome data showed that the cells reacted immediately to the induction of the production of OpuA(H223A), using NZ9000 (pNZopuAmRNA) as the reference, and significant differential expression of genes was observed already after 8 min. The PCA analysis also showed that the changes gradually increased up to 64 min after induction (Figure 3A). After preliminary analysis of the transcriptome data, the 64 min time point seemed to be most relevant and, for pragmatic reasons, was therefore selected for the transcriptomics analysis in the experiments comparing *L. lactis* strains NZ9000 (pNZPS1Δ9), NZ9000 (pNZStSUT1) and NZ9000 (pNZOpuAC).

The soluble proteome of control cells [strains NZ9000 (pNZ8048) or NZ9000 (pNZopuAmRNA)] and of cells overproducing membrane protein was analyzed by two-dimensional differential in gel electrophoresis (2D-DIGE), in combination with nanoLC-MS/MS (see Materials and Methods). Upon OpuA(H223A) overproduction, the PCA analysis on the spot volumes of proteins, that were matched across all the 2D gel images, revealed no significant differences for the time points 0 and 16 min. Differences between the induced *L. lactis* NZ9000 (pNZopuAmRNA) and NZ9000 (pNZOpuA(H223A)) cultures became significant only after 64 min. The soluble proteomes were then distinct from each other and from the earlier time points (Figure 3B) and were further analyzed.

Membrane proteomes extracted from control cells (strains NZ9000 (pNZ8048) or NZ9000 (pNZopuAmRNA)) and from cells overproducing membrane protein, collected after 64 min of induction, were analyzed by 2D-liquid chromatography, separating iTRAQ-labelled peptides by strong cation exchange (SCX)
and reverse phase nano-liquid chromatography (RP-nLC). Proteins were identified by MALDI-MS/MSMS of the peptides. To improve the identification and quantification of low abundant membrane proteins, the membrane fractions were extracted with urea/K-EDTA and subsequently with cholate to remove the majority of specifically-bound cytosolic proteins (for more detail, see Text S1 and Figure S1). Nevertheless, highly abundant soluble cytosolic proteins (glycolytic enzymes, ribosomal proteins) could still be detected in the membrane proteome fraction and these were included in the proteome dataset. The significant differentially expressed proteins and all identified proteins (raw data) are listed in Table S1.

The physiological response of L. lactis to membrane protein overproduction

The physiological response of L. lactis to recombinant membrane protein overproduction was determined by analyzing the changes in the transcriptome and the proteome in biological triplicates and by comparing these datasets (Table S1). The level at which the target genes/proteins were overexpressed/overproduced is depicted in Figure 4A. Only opuA/OpuA were endogenous and can, de facto, be said to be up- or downregulated. As the other target genes are not present in the control strain, the obtained ratios serve only as qualitative controls for the production of the target proteins in each experiment. Only native L. lactis genes were represented on the DNA microarray slides and therefore only data on the overexpression of opuA was obtained in this way. That the expression fold change was negative only signifies that the mRNA control strain NZ9000 [pNZopuAmRNA] overexpressed the opuA transcript to a higher extent than NZ9000 [pNZOpuA(H223A)] due to the lack of additional burden of overproducing the OpuA protein. A global quantification of the statistically significant observations is presented in Table 1. More changes were observed at the level of the transcriptome than at the proteome level, which is commonly observed in omics studies [34,35]. The correlation between the transcriptome and proteome datasets are represented as Venn diagrams and the Pearson coefficients are given in Figure 3C, showing that for about ~25% to 40% of the identified differentially expressed proteins the corresponding genes in the transcriptome were identified to be differentially regulated. The Venn diagrams presented in Figure 3D & 3E show the overlap between the different transcriptome (Figure 3D) and proteome (Figure 3E) datasets. The clustering of strains overproducing the three membrane proteins was also observed in the proteome data.

The total number of genes/proteins that were significantly differentially expressed upon membrane protein overproduction is given in Table 1. The stress response was most pronounced for cells overproducing OpuA(H223A), which can be explained by the fact that this protein is produced to much higher levels than PS1A9 or SsSUT1. We also note (see Fig. 1A) that growth of L. lactis was inhibited upon induction of the synthesis of OpuA(H223A), PS1A9 and SsSUT1, and that a large part of the response may reflect adaptation to the lower growth rate. The most notable differences in the physiological response of L. lactis upon production of the membrane proteins OpuA(H223A), PS1A9 and SsSUT1 and the soluble protein OpuAC are endogenous and can, de facto, be said to be up- or downregulated. As the other target genes are not present in the control strain, the obtained ratios serve only as qualitative controls for the production of the target proteins in each experiment. Only native L. lactis genes were represented on the DNA microarray slides and therefore only data on the overexpression of opuA was obtained in this way. That the expression fold change was negative only signifies that the mRNA control strain NZ9000 [pNZopuAmRNA] overexpressed the opuA transcript to a higher extent than NZ9000 [pNZOpuA(H223A)] due to the lack of additional burden of overproducing the OpuA protein. A global quantification of the statistically significant observations is presented in Table 1. More changes were observed at the level of the transcriptome than at the proteome level, which is commonly observed in omics studies [34,35]. The correlation between the transcriptome and proteome datasets are represented as Venn diagrams and the Pearson coefficients are given in Figure 3C, showing that for about ~25% to 40% of the identified differentially expressed proteins the corresponding genes in the transcriptome were identified to be differentially regulated. The Venn diagrams presented in Figure 3D & 3E show the overlap between the different transcriptome (Figure 3D) and proteome (Figure 3E) datasets. The clustering of strains overproducing the three membrane proteins was also observed in the proteome data.

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1) Cell envelope stress response and protein translocation. A number of genes/proteins that were upregulated in response to the overproduction of proteins, such as fisH, oxaA2, ppiB, pauL and telD, form part of the CesSR regulon controlled by the two-component system specified by cesSR. The genes cesS (llmg_1649) and cesR (llmg_1648) are also both upregulated under the protein production stress applied, orchestrating a cell envelope stress response [36,37] (Figure 4B). Some of these proteins, notably FisH and OxaA2, are known to play crucial roles in membrane protein biogenesis [30]. Their upregulation may increase the capacity to remove misfolded protein (FisH, a membrane-bound cell division pro tease), while allowing a more efficient insertion or folding of proteins into the membrane (OxaA2). The differential expression of all of these genes was seen with all three overproduced membrane proteins but not for OpuAC (Table S2), indicating that the integral membrane proteins induce a CesSR response.

2) General stress response. The proteome and transcriptome data show that, specially for OpuA and PS1A9, recombinant membrane protein production in L. lactis also evoked a more general stress response, including the upregulation of hrcA-gppE-dnaK, dnaJ, groES-groEL, clpP, clpB, clpE and clpC (Figure 4D; in most cases the corresponding proteins were upregulated similarly). SsSUT1, on the other hand, only evoked a change (upregulation) in the levels of hrcA, clpB and clpE (and in most cases the corresponding proteins). This response is a clear indication that the pool of mis-folded proteins is increased. Possibly, some of the produced protein molecules might not have correctly assembled in the membrane, e.g. due to an overload of the membrane targeting/insertion process. The identification of OpuA, the nucleotide-binding subunit of the ABC transporter OpuA, in the soluble proteome is consistent with a somewhat higher production of OpuAA, from the first gene of the operon, as compared to OpuABC. Possibly, the surplus OpuAA cannot associate with the membrane component and ends up in the cytoplasm (Table S1). Contrary to what was observed with the membrane proteins, in case of OpuAC overproduction none of the genes/proteins concerned with the general stress were differentially regulated (except for clpE at the transcriptome level; Table S2).

3) Protein synthesis. A severe effect on the cell’s translational machinery was observed upon production of all 4 recombinant proteins, as was the case for ribosomal proteins, both at the level of the transcriptome and the proteome (Figure 4E & Table S2). Translation and the tRNA synthetase proteins/genes show more mixed effects, which were not specific for any of the samples (Figure 4E & Table S2). Almost all of the differentially-regulated ribosomal proteins were observed in the membrane proteome (despite the urea/EDTA and cholate treatment) but not in the soluble proteome, which is consistent with an interaction of ribosomes with the SEC translocon and a coupled translation and membrane targeting/insertion process.
4) Amino acid biosynthesis. The levels of transcripts encoding for enzymes involved in amino acid biosynthesis, especially for methionine (MetC) and cysteine (CysD/K), were downregulated, but these effects were not seen at the proteome level (Figure 5A). In addition, the transcripts for the cytosolic peptidases (PepC, PepF and PepXP) were increased upon recombinant protein expression. The expression of most of these genes is controlled by a global nitrogen metabolism regulator, CodY, whose repressing effect is relieved when the intracellular concentration of branched-chain amino acids becomes limiting [39]. Thus, the differential regulation of the peptidase genes is consistent with a limitation in (branched-chain) amino acids in OpuA(H223A)-producing cells (see also [17]). Notably, the genes for the di-peptide and oligo-peptide transporters (dpp, dppT and opp), which are also regulated by CodY, were downregulated upon overexpression of the membrane proteins but they were upregulated upon OpuAC overproduction (Figure 5A & Table S2). This suggests that for membrane proteins the possible CodY-mediated upregulation may be overruled by the specific cell envelope stress response described above.

5) Nucleotide metabolism. A sharp downregulation of genes involved in the synthesis of nucleotides (purines and pyrimidines) via the de novo and salvage pathways was observed in all samples. Furthermore, genes encoding proteins involved in synthesis of deoxy-ribonucleotides (nrpF, nrdE, nrdF and llmg_0281) were downregulated, but most of these changes were not observed at the protein level (Figure 5B). The responses of nucleotide metabolism to a variety of environmental stresses has been

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**Figure 3. Analysis of proteome and transcriptome data.** Principal component analysis (PCA) of transcriptome (A) and proteome data obtained from 2D gel spot maps (B) of cells producing OpuA (H223A) (full triangles) and compared with the opuAmRNA control (open diamonds). The 2D gels were analyzed and spot volumes were extracted using Decyder 6.5 (GE Healthcare, Uppsala, Sweden). Transcriptome PCA (full squares) was obtained from the ratio data of the two populations of cells. (C) Overlap in the transcriptome (yellow) and proteome data (blue). The number of overlapping genes/proteins are indicated in the overlap of both circles and the associated Pearson correlation coefficients are given. Overlap of significant differences in the transcriptome (D) and proteome (E) datasets obtained for OpuA(H223A) (Green), PS1A9 (purple), StSUT1 (pink). doi:10.1371/journal.pone.0024060.g003
observed previously and is most likely not specific for membrane protein production. Most likely, it relates to the reduced growth rate of *L. lactis*, which has previously been coupled to a downregulation of *pyr* and *pur* genes [40–42]. Contrary to what was observed for the membrane proteins, production of OpuAC did not lead to differential regulation of the genes/proteins concerned with nucleotide metabolism (except for *pbnX* at the transcriptome level).

6) **Carbon and energy metabolism.** The levels of transcripts encoding glycolytic enzymes and pyruvate-dissipating enzymes were decreased in the OpuA(H223A), PS1A9 and StSUT1 producing cells. Similar observations were made at the proteome level (Figure 5C). Like for nucleotide metabolism, the trends correlate with the decrease in growth rate and may reflect the lower need for metabolic energy. The transcript levels for the subunits of F1F0-ATPase (*atpA, atpB, atpD* and *atpH*) were higher in cells overproducing OpuA(H223A), even though AtpD and AtpA were downregulated at the protein level. In strains overproducing PS1A9 or StSUT1, AtpD and AtpH were downregulated at the protein level (Figure 5D). Except for *pilD* at transcriptome level, none of the genes were differentially regulated in OpuAC producing cells (Table S2).

7) **Cell envelope biosynthesis.** Proteins and transcript levels for the enzymes involved in the biosynthesis of the peptidoglycan layer were upregulated, independent of the overproduced protein. With respect to lipid synthesis the cells do not seem stressed, because an increase in cardiolipin at the expense of phosphatidylglycerol is often linked to stress conditions and/or a reduced growth rate [43], however the transcriptomic and proteomic data was obtained by 2D DIGE. Only statistically significant p-values are indicated. The color scheme is a measure of the depicted fold changes (green for upregulation and red for downregulation; the color intensity is proportional to the fold change). Proteins/genes that were either significantly not differentially regulated or not identified are indicated in black.

Table 1. Number of up- and downregulated genes/proteins in *L. lactis* strains.

| Strain comparison               | Microarrays | 2D DIGE | ITRAQ |
|---------------------------------|-------------|---------|-------|
| pNZopuAm (mRNA) versus pNZopuA(H223A) | 164 209 12 9 7 92 |         |       |
| pNZ8048 versus pNZPS119          | 154 167 8 19 10 23 |         |       |
| pNZ8048 versus pNZSUT1           | 145 143 4 12 31 44 |         |       |

*L. lactis* NZ9000, carrying the indicated plasmids, was grown as described in Materials and Methods and numbers of differentially expressed genes and proteins were determined.

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Discussion

The major hurdle in the structural analysis of membrane proteins is their overproduction in a functional state. Producing membrane proteins requires coordination of several processes, such as transcription, translation, targeting, membrane insertion, folding, and, in many cases, post-translational modifications. A thorough analysis of the host cell response to and mechanistic information about membrane protein biogenesis will aid in the design of strategies to optimize the recombinant production of these proteins. Here, we report on the physiological response of *L. lactis* NZ9000 to apparent stress(es) evoked by the synthesis of a number of integral membrane proteins, using a water-soluble, cytosolically expressed protein as a reference. We show that, although OpuA is expressed to much higher levels than PS1A9 and StSUT1, the response of the cells to the stress of producing these proteins is very similar, which was not anticipated. At this point, we do not know whether a lower expression is due to a lower growth rate or a lower growth rate is caused by the expression of the particular protein. As the amount of PS1A9 and StSUT1 synthesized is very low, it is unlikely that the growth inhibition is due to diversion of nutrients towards the synthesis of recombinant protein. What, then, causes the inhibition of growth? In *E. coli* and *B. subtilis* downregulation of genes involved in transcription and translation, *e.g.*, RNA synthetases, similar to what is observed here in *L. lactis*, is invoked by the synthesis of ppGpp via the ribosome-associated RelA protein. The accumulation of ppGpp acts as an alarmone, repressing the transcription of various genes essential for cell growth. This phenomenon is called the stringent stress response [44,45]. The stringent stress response in *L. lactis* has been identified in for instance acid-stressed cells [46,47] and might play a role here.

The production of recombinant membrane protein was accompanied by a general stress response, as the heat shock proteins DnaK, GroEL, DnaJ and GrpE were highly upregulated in case OpuA and PS1A9. The expression of these genes is controlled by HrcA, a regulatory protein that binds to the CIRCE sequence present in the upstream region of these genes [48,49]. In addition, the protease ClpE and the chaperones ClpB, ClpE and ClpX, whose expression is regulated by CtsR [50], were also upregulated. The response was observed for a broad range of stresses such as acid, heat and osmotic challenges [41,51]. The response is triggered by the accumulation of misfolded protein,
Figure 5. Quantification of mRNA and protein differences in cells producing recombinant proteins. For details on headings, statistics and color scheme, see legend to Figure 4.
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suggesting that not all OpuA(H223A) and PS1Δ9 (and possibly also StSUT1) is correctly folded. Alternatively, it might be that the stress results from an increased population of generic unfolded proteins, due to depletion of the folding machinery by the overproduced recombinant proteins. Notably, none of the proteins concerned with general stress were differentially regulated in case of OpuAC overexpression.

Arguably the most important finding of our work is the CesSR-mediated response, resulting in the differential regulation of a wide variety of genes, many of which have a (putative) central role in maintaining cell envelope integrity and various membrane functions, such as LmrA (a multidrug resistance ABC transporter), RmaB (a transcriptional regulator of the MarR family), SpxB (transcriptional regulator), OxaA2 (membrane insertase/foldase) or FtsH (AAA-type ATP-dependent membrane-bound metalloprotease) (Figure 4B). CesSR is a two-component regulatory system (TCS) that orchestrates a cell envelope stress response [36]. Although it is not known what is actually sensed by CesS, the histidine-kinase and sensor component of the system, the system responds to the presence of bacteriocins [36] and lysozyme [37]. We show here that CesSR also responds to stress evoked by the overproduction of a variety of membrane proteins, and of a secretory protein. This may occur via the association of misfolded hydrophobic protein with the cytoplasmic membrane. In fact, by using GFP as a quality control indicator of correctly folded protein, we have observed that misfolded membrane proteins in L. lactis do not end up in electron-dense inclusion bodies (as frequently observed in E. coli) but, rather, are associated with the membrane lipid fraction of the cell [22]. An important role for FtsH seems obvious under those circumstances. Most genes from the CesSR regulon code putative membrane proteins or proteins acting on the cytoplasmic membrane, clearly indicating the specificity of this response. The influence of CesSR and members of this regulon on membrane protein production is described in the accompanying paper [32].

In conclusion: By using a combined proteomic and transcriptomic approach we were able to determine the physiological response of L. lactis to membrane protein overproduction (Figure 6). Intriguingly, the extent of the stress responses was not proportional to the production levels and/or activity of the overproduced protein. The observations on the general and cell envelope stress are also in agreement with a recent study in which the human ABC chloride channel CFTR was expressed in L. lactis [32]. In addition, we monitored the effect of overexpression of OpuAC, the glycine betaine-binding domain of OpuA, as a water-soluble cytoplasmic protein, but did not observe the general and cell envelope stress response. Overproduction of various other cytoplasmic proteins also did not trigger a CesSR response (Anne de Jong, personal communication). Several of the other responses, i.e. of energy, nucleotide and amino acid metabolism and of protein synthesis, seem to occur under a variety of stresses and are not related to the production of membrane proteins per se [17,41,53,54]. Our work points to a specific response in L. lactis towards membrane protein production, of which the CesSR-mediated one seems most relevant for producing well folded protein (see the accompanying paper [32], in which knowledge on the players in the CesSR response was used to engineer L. lactis to produce more membrane protein).

In contrast to E. coli, overproduction of membrane/secretory proteins in L. lactis does not elicit a major change in the expression of the Sec protein translocation machinery. On the basis of comparisons with published studies on L. lactis physiology and
stress response [17,41,53,54], we conclude that many differences in the expression of genes in for instance carbon, energy and nucleotide metabolism are due to the decreased growth rate. Under these conditions, the cell may divert nutrients towards the synthesis of recombinant proteins and/or have too limited a capacity to import amino acids in case of nitrogen metabolism. Finally, it is evident that upon membrane protein production the cell is sensing various changes that elicit a response in the transcription machinery, which is not reflected in the proteome. The proteome may thus be a better indicator for design and engineering of cells with the aim to overcome expression bottlenecks.

Materials and Methods

Bacterial strains and growth conditions

_Lactococcus lactis_ NZ9000 [28] derivatives were grown at 30°C in M17 broth (Difco, Detroit, MI, USA) containing 1% glucose (w/v) and 5 μg/ml chloramphenicol (GM17-Cm5). _L. lactis_ Opu401 is a derivative of _L. lactis_ NZ9000 in which the chromosomal _opuA_ genes have been deleted by double cross-over recombination [31].

Plasmid construction

DNA manipulations were done according to standard procedures. Plasmids and primer sequences used in this study are listed in Table 2 and Table S3, respectively. The _opuA_ mutants, either specifying an inactive version of OpuA or a non-translatable transcript, were constructed by site-directed mutagenesis on pNZOpuA [55]. Ligation-Independent Cloning of PS1A9 and StSUT1 in pRecLIC and the subsequent conversion of the plasmids into lactococcal expression vectors by Vector Backbone Exchange (VBEx) were performed as described [11]. All proteins were produced with a C-terminal his-tag for rapid screening of expression. Preparation of electrocompetent cells and electroporation into _L. lactis_ was performed as described [56].

### Protein production and immunodetection

For protein (over)production, cells were grown until OD$_{600}$=0.5 and induced with $10^{-3}$ volume of filtered culture supematant of the nisin-A producing strain _L. lactis_ NZ9700, containing 10 μg/mL of nisin A. Cells were allowed to grow for the required amount of time and harvested by centrifugation. Sample volumes were normalized on the basis of OD so that an equivalent amount of whole-cell protein was taken for all samples. Cells were washed once with 100 mM potassium phosphate (KPi), pH 7.0, and resuspended in 1 mL of ice-cold 100 mM KPi, pH 7.0, 10% glycerol (w/v), 1 mM MgSO$_4$, 1 mM PMSF and trace amounts of DNase I. After the addition of 300 mg of glass beads (~100 μm diameter), cells were lysed by three rounds of bead beating in a Fastprep machine for 20 seconds (speed 6.0) with cooling intervals of 5 min on ice in between. Unbroken cells and cell debris were removed by centrifugation at 16,100 × g for 30 min and membrane fragments were collected by centrifugation at 267,000 × g for 20 min. Protein samples were resolved on 12.5% SDS-PAGE gels and detected by immunodetection with an anti-histidine tag primary monoclonal antibody (GE Healthcare, Uppsala, Sweden). Chemiluminescence detection was done using the Western-light kit with CSPD (Tropix Inc, Bedford, MA, USA) as the substrate and imaging with the LAS-3000 imaging system (Fujifilm, Minatoku, Tokyo, Japan).

Reverse transcriptase-quantitative-PCR

An equivalent of 10 OD$_{600}$ units [OD$_{600}$ × V (ml)] of _L. lactis_ cells were harvested by centrifugation and cell pellets were kept at −80°C until further processing. Cells were washed with DEPC treated T$_{10}$E$_{1}$ buffer and resuspended in 500 μL T$_{10}$E$_{1}$ (10 mM Tris-HCl pH 8.0, 1 mM Na$_2$-EDTA) and transferred to 2 ml screw-cap tubes. To this cell suspension, 50 μl 10% SDS (w/v), 500 μl phenol/chloroform, 500 mg glass beads (50–105 μm of diameter) and 175 μl macaloid suspension (Bentone MA, Highstown, NJ) were added. All reagents used for RNA work were

### Table 2. Bacterial strains and plasmids used in this study.

| Description* | Mol Wt (kDa) | No. of TMDs$^b$ | % membrane segment to extramembranous loops | Source |
|---------------|-------------|----------------|---------------------------------------------|--------|
| **Strains**   |             |                |                                             |        |
| _L. lactis_ MG1363 | _L. lactis_ subsp. cremoris, plasmid-free derivative of NCD0712 | 54.8 | 12 | 51.16 | This work |
| _L. lactis_ NZ9000 | _L. lactis_ MG1363 Δ pepN::nisRK | 54.8 | 12 | 51.16 | This work |
| _L. lactis_ Opu401 | _L. lactis_ NZ9000 Δ opuAABC | 54.8 | 12 | 51.16 | This work |
| **Plasmids**  |             |                |                                             |        |
| pNZ8048      | Cm$^r$, Expression vector with nisin A-inducible promoter P$_{nisA}$ | 28.9 | 0 | 0 | [29] |
| pNZOpuA      | pNZ8048 containing opuA and opuABC | 107.8 | 7 | 15.3 | [55] |
| pNZOpuA(H223A) | pNZOpuA derivative; specifying OpuA with the His at position 223 replaced by Ala | 107.8 | 7 | 15.3 | This work |
| pNZOpuA(mRNA) | pNZOpuA derivative carrying opuA with ATG codons at positions 1, 1224 and 1344 replaced by TAA; the numbering refers to the position of nucleotide in the gene sequence | 49.3 | 9 | 45.2 | This work |
| pNZPS1A9     | pNZClic derivative specifying PS1A9 with a hexa-histidine tag on the C-terminus | 54.8 | 12 | 51.16 | This work |
| pNZStSUT1    | pNZClic derivative specifying StSUT1 with a hexa-histidine tag on the C-terminus | 28.9 | 0 | 0 | [29] |

*Cm$^r$, chloramphenicol resistance.

$^b$TMD’s: Transmembrane domains.

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treated with diethylpyrocarbonate (DEPC) Sigma-Aldrich, St. Louis, MO). The macoloid suspension was prepared as follows: 2 g Macoloid was added to 100 ml T_{10}E_{1}, boiled for 5 min, then cooled to room temperature and sonicated for short periods of time until a gel was formed; the gel was spun down in a microcentrifuge and resuspended in 50 ml T_{10}E_{1} (pH 8.0). Cells were disrupted by bead-beating twice for 45 sec in a Mini-BeadBeater (Biospec Products, Bartlesville, OK) with a 1-min cooling interval on ice. The cell lysate was cleared by centrifugation and 500 μl supernatant was extracted with 500 μl phenol/chloroform, and subsequently with 500 μl chloroform.

Total RNA was isolated from the water phase using the High Pure RNA Isolation Kit (Roche Molecular Biochemicals, Mannheim, Germany), according to the manufacturer’s protocol. RNA quality was verified with an Agilent Bioanalyzer 2100 using RNA 6000 LabChips (Agilent Technologies Netherlands BV, Amstelveen, the Netherlands) and RNA concentration was determined spectrophotometrically with a Nanodrop ND1000 (NanoDrop Technologies, Wilmington, DE). Copy DNA (cDNA) was synthesized using Superscript III Reverse Transcriptase (Invitrogen, Carlsbad, CA) and the quantification was done with Maxima SYBR Green qPCR Master Mix (Fermentas, Burlington, Canada), according to the suppliers instructions and using an optical iCycler (BioRad, Hercules, California, USA). The forward and reverse primers used for the qPCR were spaced 100 bp apart, around the 5′-UTR.

Preparation of cell samples for transcriptomics and proteomics

To assure true biological replicates (triplicates), _L. lactis_ cells transformed with appropriate plasmids were streaked onto GM17-Cm5 agar plates and single colonies were used to start pre-inoculums in GM17-Cm5 medium. After 8–10 h of growth of the pre-inoculums, several dilutions were prepared, ranging from 10^{-2} to 10^{-6} fold, to obtain an overnight culture growing exponentially (i.e., OD_{600} = 0.2–0.4). This culture was used to inoculate 2.5 L of fresh GM17-Cm5 medium (1/100 dilution). The culture was stirred at 200 rpm and the pH was maintained at 6.8 by automatic determination of 10 mg/ml (based on protein concentration determination using Bradford reagent, with bovine serum albumin as calibration standard) in a volume of 200–500 μl (initial volume) and kept on ice. An equal volume of buffer A supplemented with 10 M urea plus 10 mM K-EDTA pH 8.0 was added slowly, while being stirred and the solution was incubated on ice for 20 min. The ‘stripped’ membrane vesicles were collected by centrifugation at 272,000 × g for 1 h at 4°C and resuspended in the initial volume with buffer A. An equal volume of buffer A supplemented with 14C-glycine betaine to a final concentration of 15 mM; the transport reaction was stopped at given times by the addition of 2 ml of ice cold stop buffer (650 mM sucrose in 50 mM K-HEPES pH 7.3), followed by filtration through 450 nm pore size nitrocellulose filters. The filters were washed twice with 2 ml of stop buffer and subsequently transferred into vials containing 2 ml of scintillation fluid. The radio activity was measured using a TriCarb-2800 TR liquid scintillation analyzer (PerkinElmer, Massachusetts, USA).

For mRNA isolation, an equivalent of 10 OD_{600} units (OD_{600} = 0.5) of _L. lactis_ cells was harvested at the various time points. In case of OpuA(H223A) and its control (OpuAmRNA), cells were harvested 0, 2, 8, 16, 32 and 64 min after the addition of inducer. In case of PS1Δ9, SsSU1 and OpuAC and the corresponding control (pNZ8048), only the cells from the 64 min time point were used. Cells were resuspended in ice-cold 500 μl T_{10}E_{1}, transferred to 2 ml screw-cap tubes, immediately frozen in liquid nitrogen and kept at −80°C until further processing.

For the proteome analyses of cells expressing OpuA(H223A) and OpuAmRNA, one liter of culture sample was rapidly withdrawn at 0, 16 and 64 min after the addition of inducer; for the analyses of cells expressing PS1Δ9, SsSU1 or OpuAC or carrying the control plasmid pNZ8048, only the 64 min time point was used. To inhibit protein synthesis, chloramphenicol (100 μg/ml, final concentration) was added immediately upon sampling. Cells were harvested by centrifugation at 6281 × g for 15 min at 4°C. The cell pellet was washed once with ice-cold 100 mM KPi, pH 7.0, resuspended in 5 ml of the same buffer, frozen in liquid nitrogen and stored at −80°C.

### Glycine betaine transport assay

Whole-cell glycine betaine transport was measured essentially as described before [58]. Cells were grown in GM17 to an OD_{600} = 0.3 and induced for 1 h with 100 ng/L nisin A (final concentration), harvested (at OD 600 = 0.1) and washed with 50 mM K-HEPES pH 7.3, concentrated to OD600 of 5.0 into 50 mM K-HEPES pH 7.3 plus 650 mM sucrose and 10 mM glucose and pre-energized for 5 min by stirring at 200 rpm and the pH was maintained at 6.8 by automatic determination using Bradford reagent, with bovine serum albumin as calibration standard) in a volume of 200–500 μl (initial volume) and kept on ice. An equal volume of buffer A supplemented with 10 M urea plus 10 mM K-EDTA pH 8.0 was added slowly, while being stirred and the solution was incubated on ice for 20 min. The transport reaction was stopped at given times by the addition of 2 ml of ice cold stop buffer (650 mM sucrose in 50 mM K-HEPES pH 7.3), followed by filtration through 450 nm pore size nitrocellulose filters. The filters were washed twice with 2 ml of stop buffer and subsequently transferred into vials containing 2 ml of scintillation fluid. The radio activity was measured using a TriCarb-2800 TR liquid scintillation analyzer (PerkinElmer, Massachusetts, USA).

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12% cholic acid (w/v) was added slowly to the membrane vesicles, while being stirred and the solution was incubated for 20 min on ice. The membrane vesicles were collected by centrifugation at 272,000 × g for 1 h at 4 °C and resuspended in the initial volume with buffer A. Protein concentrations were on average around 1 mg/ml after the two extraction steps.

Membrane proteome identification

The extracted membrane vesicles were digested with trypsin, the obtained peptides were labeled with iTRAQ reagents, the labeled peptide mixtures were separated by strong cation exchange (SCX) and reverse phase nano-liquid chromatography (RP-nLC) and subsequently identified/quantified by MALDI-MS/MSMS according to published methods [52]. A detailed description of the experimental setup and conditions is presented in the Text S1. The cut-offs that were used to define whether the observed differences in protein levels are significant were kept more strict than the cut-offs defined by [52], that is, differences in relative protein ratios were considered to be significant if both ratios of the biological replicates had a p-value below 0.01 and an average intensity fold change greater than 1.5, or lower than -1.5.

DNA-microarray experiments

Total RNA isolation and cDNA synthesis was performed as described above for the RT-qPCR, except that amino allyl-modified dUTPs were used in the nucleotide mix for cDNA synthesis. Indirect Cy-3/Cy-5 labelling of cDNA was performed according to supplier’s instructions (Amersham Biosciences, Piscataway, NJ). Hybridisation of Cy-labelled cDNA was performed during 16 h at 45 °C in a microarray hybridisation incubator ISO20 (Grant Boeckel, Cambridge, UK) in Ambion Slidehyb #1 hybridisation buffer (Ambion Biosystems, Foster City, CA). SuperAmine glass slides (ArrayIt, Sunnyvale, CA) spotted with duplicates of around 2500 ORF amplicons of L. lactis subsp. cremoris MG1363 [33] were used. Slides were scanned using a GenePix Autoloader 4200AL scanner (Molecular Devices Corporation, Sunnyvale, CA). DNA microarray data from biological replicates were obtained through dye-swaps, to discard possible differences between the Cy-3 and Cy-5 labeling data from biological replicates were obtained through dye-swaps, to discard possible differences between the Cy-3 and Cy-5 labeling data from biological replicates were obtained through dye-swaps, to discard possible differences between the Cy-3 and Cy-5 labeling data from biological replicates were obtained through dye-swaps, to discard possible differences between the Cy-3 and Cy-5 labeling data from biological replicates were obtained through dye-swaps, to discard possible differences between the Cy-3 and Cy-5 labeling data from biological replicates were obtained through dye-swaps, to discard possible differences between the Cy-3 and Cy-5 labeling data from biological replicates were obtained through dye-swaps, to discard possible differences between the Cy-3 and Cy-5 labeling data from biological replicates were obtained through dye-swaps, to discard possible differences between the Cy-3 and Cy-5 labeling.
5. Katzen F, Peterson TC, Kudlicki W (2009) Membrane protein expression: no cells required. Trends Biotechnol 27: 455–460.
6. Mierau I, Klarebezem M (2005) 10 years of the nisin-controlled gene expression system (NICE) in Lactococcus lactis. Appl Microbiol Biotechnol 69: 747–755.
7. Surade S, Klein M, Stolt-Bergner PC, Muenke C, Roy A, et al. (2006) Comparative analysis and “expression space” coverage of the production of prokaryotic membrane proteins for structural genomics. Protein Sci 15: 2178–2189.
8. Quick M, Javitch JA (2007) Monitoring the function of membrane transport proteins in detergent-solubilized form. Proc Natl Acad Sci U S A 104: 3603–3608.
9. Morello E, Bermudez-Huumaran LG, Lluit D, Sole V, Miraglio N, et al. (2008) Lactococcus lactis, an efficient cell factory for recombinant protein production and secretion. J Mol Microbiol Biotechnol 14: 45–58.
10. Mulligan C, Geertsma ER, Severi E, Kelly DJ, Poolman B, et al. (2009) The sfpA-sppM-binding protein imposes directionality on an electrochemical sodium gradient-driven TRAP transporter. Proc Natl Acad Sci U S A 106: 1778–1785.
11. Geertsma ER, Poolman B (2007) High-throughput cloning and expression in Lactococcus lactis as host for overproduction of functional membrane proteins. Biochim Biophys Acta 1760: 97–108.
12. de Ruyter PG, Kuipers OP, de Vos WM (1996) Over-expression of proteins in Escherichia coli. Bio/Technology 14: 1729–1740.
13. Perret LM, McEwan M, Clark D, Henry J, Tang S, et al. (2010) Amino acid accumulation limits the overexpression of proteins in Lactococcus lactis. PLoS ONE 5: e10317.
14. Macrina L, Rees AT, Rees DC (2008) The funnel approach to the high-throughput cloning and expression in Lactococcus lactis. Proteomics 8: 10.
15. Monne M, Chan KW, Slotboom DJ, Kunji ER (2005) Functional expression of Lactococcus lactis. Appl Environ Microbiol 70: 5398–5406.
16. Kuipers OP, de Ruyter PG, Kleerebezem M, de Vos WM (1998) Quorum sensing by the lacI-lacX operon: evidence for a two-component system CesSR. Mol Microbiol 30: 45–55.
17. Eymann C, Homuth G, Scharf C, Hecker M (2002) Bacillus subtilis functional genomics: global characterization of the stringent response and proteome analysis. J Bacteriol 184: 2500–2502.
18. Varmanen P, Ingmer H, Vogensen FK (2000) ctsR of Lactococcus lactis subsp. cremoris Grown with Glycine in Osmotically Stabilized media. Appl Environ Microbiol 66: 15–21.
19. van Hijum SA, de Jong A, Buist G, Kok J, Kuipers OP (2003) UniFrag and Comparative analysis and “expression space” coverage of the production of Lactococcus lactis. Proteomics 3: 786–797.
20. van Hijum SA, de Jong A, Barea R, van Strijp S, van Hijum SA (2005) Comparative genomic and proteomic analyses of a systematically perturbed metabolic network. Science 309: 539–542.
64. Kanehisa M, Goto S, Hattori M, oki-Kinoshita KF, Itoh M, et al. (2006) From genomics to chemical genomics: new developments in KEGG. Nucleic Acids Res 34: D354–D357.
65. Kanehisa M, Araki M, Goto S, Hattori M, Hirakawa M, et al. (2008) KEGG for linking genomes to life and the environment. Nucleic Acids Res 36: D480–D484.
66. Sturn A, Quackenbush J, Trajanoski Z (2002) Genesis: cluster analysis of microarray data. Bioinformatics 18: 207–208.
67. Gasson MJ (1983) Plasmid complements of Streptococcus lactis NCDO 712 and other lactic streptococci after protoplast-induced curing. J Bacteriol 154: 1–9.
68. Kleerebezem M, Beerthuyzen MM, Vaughan EE, de Vos WM, Kuipers OP (1997) Controlled gene expression systems for lactic acid bacteria: transferable nisin-inducible expression cassettes for Lactococcus, Leuconostoc, and Lactobacillus spp. Appl Environ Microbiol 63: 4581–4584.