Expression and functional characterization of membrane proteins
Gul, Nadia

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Chapter 3

Functional reconstitution and osmoregulatory properties of the ProU ABC transporter from *Escherichia coli*.

Nadia Gul and Bert Poolman.

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1. Abstract
The ATP-binding cassette (ABC) transporter ProU from *Escherichia coli* translocates a wide range of compatible solutes and contributes to the regulation of cell volume, which is particularly important when the osmolality of the environment fluctuates. We have purified the components of ProU, i.e., the substrate-binding protein ProX, the nucleotide-binding protein ProV and the transmembrane protein ProW, and reconstituted the full transporter complex in liposomes. We engineered a lipid anchor to ProX for surface tethering of this protein to ProVW-containing proteoliposomes. We show that glycine betaine binds to ProX with high-affinity and is transported via ProXVW in an ATP-dependent manner. The activity ProU is salt and anionic lipid-dependent and mimics the ionic strength-gating of transport of the homologous OpuA system.

2. Introduction
Osmotic stress is an important environmental parameter that influences the growth and adaptation of (micro)organisms in a given habitat. Microbial cells employ different strategies to overcome the ‘osmotic challenges' in their environment. For instance, under hyperosmotic conditions, bacteria can accumulate osmoprotective compounds or compatible solutes, such as choline, proline, glycine betaine, carnitine and ectoine (Perroud, Le Rudulier 1985, Grothe et al. 1986, Wood et al. 2001). In addition to these zwitterionic compounds, organisms may transiently accumulate potassium ions together with glutamate or other anionic species as is best documented for *Escherichia coli* (Booth, Higgins 1990, Ogahara et al. 1995). The accumulation of potassium ions is fast but also disturbs ionic homeostasis of the cell, and over time the potassium and counter ion(s) are replaced by neutral or zwitterionic solutes such as trehalose, proline, glycine betaine and others (Lucht, Bremer 1994, Sutherland et al. 1986). The accumulation of compatible solutes can be achieved via de novo biosynthesis or rapid uptake of solutes from the medium; the latter mechanism can be more effective in terms of costs of metabolic energy. Osmoregulatory transport systems have been extensively studied in both Gram-positive and Gram-negative bacteria, e.g., OpuA from *Lactococcus lactis* and *Bacillus subtilis*, BetP from *Corynebacterium glutamicum*, OpuC from *Pseudomonas syringae*, and ProP and ProU from *E. coli* (Kempf, Bremer 1995, Perez et al. 2011a, Mahmood et al. 2006, Karasawa et al. 2011, Culham, Meinecke & Wood 2012, May et al. 1986, Kempf, Gade & Bremer 1997, Horn, Bremer & Schmitt 2003, Cairney, Booth & Higgins 1985). Other biochemically less well-studied systems include: Ota from *Methanosarcina mazei*, and ProP and ProU from *Yersinia enterocolitica* (Schmidt et al. 2007, Annamalai, Venkitanarayanan 2009). The OpuA, OpuC, Ota and ProU systems are all ATP-binding cassette transporters that have a cystathionine-β-synthase (CBS) module fused to the C-terminus of the nucleotide-binding domain. This CBS module has been shown to be critical for the osmoregulation of the OpuA transporter from *L. lactis* and OpuC from *P. syringae* (Biemans-Oldehinkel, Mahmood & Poolman...
2006, Chen, Beattie 2007, Mahmood, Biemans-Oldehinkel & Poolman 2009, Mahmood, Biemans-Oldehinkel & Poolman 2009). A particularly interesting system is ProU from *E. coli* as the osmotic regulation of its expression is known in great detail (Higgins 1992, Lucht, Bremer 1994, Perroud, Le Rudulier 1985). So far, little is known of the regulation of the activity of the protein itself but activation by osmotic stress has been observed (Faatz, Middendorf & Bremer 1988). The *proU* operon includes three open reading frames (ORFs) *proV, proW* and *proX* (Mimmack et al. 1989, Gowrishankar 1989). The *proV* gene encodes the NBD with the C-terminal tandem CBS domains. *ProW* encodes the transmembrane domain (TMD) and has an N-terminal ~100 amino acid extension of unknown function (Haardt, Bremer 1996). *ProX* encodes the substrate-binding protein (SBD), which is localized in the periplasm. *ProX* has been purified to homogeneity and shown to bind glycine betaine with high affinity (*K*<sub>D</sub> of ~1 μM); proline betaine is bound with a *K*<sub>D</sub> of ~5 μM (Barron, Junga & Villarejo 1987, May et al. 1986, Haardt et al. 1995).

Here, we report on the *in vitro* reconstitution and biochemical characterization of the osmoregulatory ABC transporter ProU from *E. coli*. We engineered *ProX* with an N-terminal lipid anchor sequence for efficient tethering of the substrate-binding protein to the surface of proteoliposomes, in which *ProVW* was reconstituted first (figure 1). We show that the ProU system is activated by osmotic stress and that, similar to OpuA, anionic lipids play an important role in the activity and regulation of transport.

### 3. Materials and methods

#### 3.1 Bacterial strains and growth conditions

For expression and purification of ProX, *L. lactis* Opu401 was used as host and transformed with pAMP31proX. Opu401 is a derivative of *L. lactis* NZ9000 with the chromosomal *opuA* genes deleted (Biemans-Oldehinkel, Mahmood & Poolman 2006). *L. lactis* was cultivated semi-aerobically in M17 broth (Difco Laboratories, East Molesey, UK) at 30°C pH 6.5, supplemented with 1% (w/v) glucose plus 5 μg/ml erythromycin, using a 10 litre pH- and temperature-controlled bioreactor. Cells were harvested at OD<sub>600</sub> of ~5, corresponding to the late exponential phase of growth, washed and resuspended in 50 mM potassium phosphate (KPi) pH 7.0 and stored at −80°C after flash-freezing of 40 ml aliquots of cells in liquid nitrogen.

For expression and purification of ProVW, *E. coli* strain MC1061 (Casadaban, Cohen 1980) was used as host and transformed with pBADcLICproVW. The cells were cultivated under vigorous aeration in Luria broth supplemented with 100 μg/ml ampicillin at 37°C and 200 rpm. At A<sub>600</sub> of 0.5–0.6 the temperature was decreased to 25°C, and, after equilibration for 15 min, membrane protein
expression was induced by the addition of L-arabinose in growth medium at a final concentration of 0.001% (w/v). Cultivation was continued for 3 h and cells were harvested, washed and resuspended in 50 mM KPi pH 7.0.

3.2 Construction of pBADcLICproVW and pNZcLICproVW

The proVW genes were amplified using genomic DNA of *E. coli* K12 as template and the following oligonucleotides:

**ProV fwd nLic:** 5′
ATGGTGAGAATTTATATTTTCAAGGTATGGCAATTAAATTAGAAAT

**ProW rev nLic** 5′
TGGGAGGGTGGGATTTTCATTACTTAATGAATGGGCGGGTCAG

**ProV fwd cLic:** 5′
ATGGGTGGTGGATTTGCTATGGCAATTAAATTAGAAAT

**ProW rev nLic** 5′
TTGGAAGTATAAATTTTCCTTAATGAATGGGCGGGTC

Both genes were ligated in the pBADnLIC and pBADcLIC vector and modified at the 3′ end to specify a 10-histidine tag, using a ligation-independent cloning method as described in (Geertsma, Poolman 2007).
Figure 1: Schematic representation of the organization of the subunits in ProU. (A) The engineered lipid-anchored ProX, the membrane-embedded TMD ProW, and the NBD ProV are shown. (B) Schematic of proteoliposomes with ProVW in two orientations, ProX tethered to the external surface, and the ATP-regenerating system (ARS) in the vesicle lumen. Only right-side-out ProVW plus ProX will facilitate the uptake of glycine betaine.

3.3 Construction of pAMP31-proX and pAMP31-GGGSAGGGS-proX

The wild-type ProX is produced with a signal sequence for translocation of the protein into the *E. coli* periplasm. To provide ProX with a lipid anchor, the *proX* gene was cloned in pAMP31, downstream of the *oppA* signal sequence and upstream of a sequence specifying a C-terminal six-histidine tag (Picon et al. 2000). When expressed in *L. lactis*, the OppA signal sequence is cleaved at position 23 and the N-terminal Cys residue is lipid modified (Hutchings et al. 2009). For this purpose, a unique *XhoI* site was introduced at position 88 of the *oppA* gene in pAMP31OppA, which allowed replacing the *oppA* gene of
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pAMP31 for proX. The proX gene was amplified by PCR, using genomic DNA of *E. coli* K12 as template and the oligonucleotides ProX fwd (with *Xho*I site) and ProX rev (with *Bam*HI) (Table 1). The PCR product was digested with *Xho*I and *Bam*HI and ligated into the vector pAMP31 (cut with *Xho*I and *Bam*HI). The resulting plasmid was named pAMP31-proX and transformed into *L. lactis* Opu401 by electroporation.

To increase the space between the lipid anchor and the ProX protein, we introduced a flexible linker (sequence GGGSAGGGS) by inserting the corresponding synthetic DNA fragment into a unique *Bgl*II site that was created just after the proX signal sequence. Thus, pAMP31-proX-*Bgl*II was digested with *Xho*I and *Bgl*II, and the synthetic DNA with complementary ends was introduced. The synthetic DNA was prepared from complementary oligonucleotides ProX (GGGSAGGGS) fwd and rev; Table 1); the oligonucleotides were annealed by slowly decreasing the temperature from 94–40°C (over 3 min). The resulting plasmid, pAMP31-GGGSAGGGS-proX, was transformed into *L. lactis* Opu401 by electroporation. Initial experiments showed that the GGGSAGGGS linker is required for substrate transfer from ProX to ProVW, and in the experiments described below pAMP31-GGGSAGGGS-proX was used to produce lipid-anchored ProX (figure 1).

Table 1: Primers used for construction pAMP31+GGGSAGGGS+proX. Restriction sites are indicated in bold and underlined. Nucleotides different from those in wild type ProX are indicated in italics.

<table>
<thead>
<tr>
<th>Primer</th>
<th>sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>ProX fwd</td>
<td>ATGCCTCGAGTATGCGACATAGCGTACTTTTTTGC</td>
</tr>
<tr>
<td>ProX rev</td>
<td>GCATGGATCCCTTCTGCCTGCGCAGCGCCTTC</td>
</tr>
<tr>
<td>ProX <em>Bgl</em>II fwd</td>
<td>CATGCAAGATCTGGCGGCAAAGGCATTACTGTATCCAG</td>
</tr>
<tr>
<td>ProX <em>Bgl</em>II rev</td>
<td>CATGCAAGATCTGGCGGCAAAGGTATTGTGTAGAGATAAGCG</td>
</tr>
<tr>
<td>ProX(GGGSAGGGS)fwd</td>
<td>TCGAGTGGTGAGGAGTTCCAGCTGGAGGTTGGATCAGCA</td>
</tr>
<tr>
<td>ProX(GGGSAGGGS)rev</td>
<td>GATCTGCTGATCCACCTCCAGCTGAACCTCCACCCAC</td>
</tr>
</tbody>
</table>
3.4 Co-transformation with pNZcLICproVW and pAMP31-GGGSAGGGS-proX

*L. lactis* Opu401 was transformed with pNZcLICproVW and pAMP31-GGGSAGGGS-proX for the simultaneous expression of ProVW and ProX. The mechanism of replication (rolling circle versus theta replication) and antibiotic resistance markers (chloroamphenicol versus erythromycin) of these plasmids differ, which makes them suitable for co-expression studies. 100 ng of each of the plasmids was used for electroporation, after which the cells were plated on GM17 agar, supplemented with 2.5 μg/ml of chloroamphenicol and 2.5 μg/ml erythromycin for plasmid selection.

3.5 *In vivo* uptake experiments

For *in vivo* glycine betaine uptake, *L. lactis* Opu401, containing pNZcLICproVW and pAMP31-GGGSAGGGS-proX, was grown in GM17 or GCDM supplemented with 2.5 μg/ml chloroamphenicol plus 2.5 μg/ml erythromycin. At OD₆₀₀ ~ 0.5, nisin A was added at a final concentration of 0.01% (v/v) to induce the expression of *proVW*. The expression of *proX* was driven from the constitutive lactococcal promoter *P₃₂* (van der Vossen, van der Lelie & Venema 1987). After 2 h of nisin A induction, the cells were washed twice and resuspended to OD₆₀₀ of ~ 12.5 (which corresponds to ~ 2.5 mg/ml protein) in 50 mM HEPES-methylglucamine pH 7.3. For the uptake assays, the cells were diluted to 0.2 mg/ml of total cell protein and pre-energized with 10 mM glucose for 5 min at 30°C. At t = 5 min, 50 mM HEPES-methylglucamine pH 7.3 supplemented with varying amounts of sucrose (0–600 mM) plus [¹⁴C]glycine betaine (final concentration of 1 mM; the total reaction mixture was 500 μl) was added, and uptake was monitored over time. To prevent protein synthesis, 25 μg/ml chloroamphenicol plus 25 μg/ml erythromycin was present during the assay. At given time intervals, 80 μl samples were withdrawn and diluted with 2 ml ice-cold assay buffer of equal osmolality and immediately filtered through 0.45 μm cellulose nitrate filters under high vacuum. After washing with 50 mM HEPES-methylglucamine pH 7.3 with the appropriate sucrose concentration, the filters were dried and dissolved in 2 ml scintillation liquid and radioactivity was determined in a scintillation counter.

3.6 Membrane vesicle preparation

Membrane vesicles were prepared by lysing the cells in a cell disruption system (high pressure Constant Systems cell disrupters, UK) at a pressure of 39,000 psi for *L. lactis* and 25,000 psi for *E. coli* at 4°C in 50 mM potassium phosphate (KPi) with 1 mM MgSO₄, 1 mM PMSF, 100 μg/ml DNase plus 100 μg/ml RNase. Subsequently, differential (ultra) centrifugation steps were carried out to separate membrane vesicles from cell debris and cytosolic fractions.


3.7 Purification of ProVW

For purification of ProVW, membrane vesicles (10 mg of total protein) of *E. coli* MC1061/pBADcLICproVW were solubilized with 1% (w/v) dodecyl-β-D-maltoside (DDM) for 45 min at 4°C; the membranes at 5 mg/ml of protein were present in buffer A (50 mM KPi, 200 mM KCl plus 20% glycerol [pH 7.0]) plus 1 mM dithiothreitol (DTT). Following ultracentrifugation at 347,000 g for 15 min, the solubilized material was incubated with Ni-Sepharose (GE Healthcare) (20 mg of protein/ml resin) for 1 h at 4°C, i.e., after the addition of 15 mM imidazole plus 1 mM DTT. Next, the resin was washed with 20 column volumes of buffer A with 0.05% DDM plus 50 mM imidazole. ProVW was eluted in buffer A with 0.05% DDM plus 200 mM imidazole.

3.8 Purification of proX

For purification of lipid-anchored ProX, membrane vesicles of *L. lactis* Opu401/pAMP31-proX were solubilized with 1% DDM for 45 min in buffer B (25 mM KPi, 100 mM KCl plus 10% glycerol [pH 7.0]) at a final concentration of 5 mg/ml of protein. Following ultracentrifugation at 347,000 g for 15 min, the solubilized material was incubated with Ni-Sepharose (20 mg of protein/ml resin) for 1 h at 4°C in the presence of 15 mM imidazole. Next, the resin was washed with 20 column volumes of buffer A with 0.05% DDM plus 50 mM imidazole. The protein was eluted in buffer B with 0.05% DDM plus 200 mM imidazole. Next, the buffer was exchanged to buffer B with 0.05% DDM plus 15 mM imidazole for membrane reconstitution or 100 mM KPi plus 0.05% DDM for fluorescence measurements, using a NAP10 column (GE Healthcare).

3.9 Membrane reconstitution of ProVW and surface tethering of lipid-anchored ProX

For membrane reconstitution of ProVW, preformed liposomes composed of *E. coli* polar lipids and L-α-phosphatidyl choline from egg yolk in a ratio of 3:1 (w/w) or synthetic lipid mixtures composed of dioleoyl-phosphatidylethanolamine (DOPE), dioleoyl-phosphatidylglycerol (DOPG) and dioleoyl-phosphatidylcholine (DOPC), at mole ratios specified in the Figure legends, were used. Liposomes were extruded through a 400 nm pore diameter (400nm pore diameter) polycarbonate filter, and, subsequently, detergent-denaturated by adding increasing amounts of Triton X-100; typically, 70–100 μl of 10% (v/v) of Triton X-100 was used for 2.5 ml of liposomes at a lipid concentration 4 mg/ml (Geertsma et al. 2008b). Purified ProVW was incorporated in the Triton X-100-denaturated liposomes at a protein-to-lipid ratio (w/w) of 1:100. Subsequently, Biobeads™ were added in steps to remove the detergent (Geertsma et al. 2008b); 1 mM DTT was present in all steps of the purification and membrane reconstitution. For membrane tethering of lipid-anchored ProX, proteoliposomes containing ProVW were extruded through a 400 nm polycarbonate filter, diluted to 4 mg/ml in 50 mM KPi pH 7.0 and mixed with
purified ProX at a protein-to-lipid ratio of 1:50 (w/w). Residual detergent was removed by incubation of the (proteo)liposomes with Biobeads. The ATP regenerating system (ARS), composed of 10 mM Mg-ATP, 24 mM phosphocreatine plus 2.4 mg/ml creatine kinase in 50 mM KPi, pH 7.0, was included in the proteoliposomes by two cycles of freezing in liquid nitrogen and thawing at room temperature (Geertsma et al. 2008b, Doeven et al. 2004) (figure 1).

3.10 Glycine betaine uptake in proteoliposomes

For ATP-driven uptake of glycine betaine into proteoliposomes, the membranes with ARS inside were extruded and resuspended in 100 mM KPi pH 7.0, and, prior to the start of the assay, they were diluted to a final lipid concentration of 5 mg/ml. The assay buffer was composed of 100 mM KPi pH 7.0 plus 0–300 mM KCl. Following 2 min of equilibration at 30°C, the transport reaction was started by the addition of 40 μM [14C]glycine betaine. At given time intervals, 25 μl samples were withdrawn and diluted with 2 ml ice-cold assay buffer (isotonic with the reaction conditions) and immediately filtered through 0.45 μm cellulose nitrate filter. Following washing of the filter with another 2 ml of ice-cold assay buffer, the radioactivity was determined in a scintillation counter.

3.11 Glycine betaine binding

Binding of radio labeled glycine betaine was measured by the precipitation method (Richarme, Kepes 1983), which is based on the principle that, upon salting-out of the protein by ammonium-sulfate, the substrate remains trapped in the ligand-binding site. Binding assays were carried out in a volume of 100 μl with 100 μg/ml of purified ProX in 100 mM KPi, pH 7.0, plus 0.05% (w/v) DDM. Following equilibration for 2 min at 30°C, tracer amounts of [3H]-glycine betaine plus varying amounts of non-labeled glycine betaine were added to the assay mixture, yielding final concentrations of glycine betaine ranging from 0.1–20 μM. The binding reaction was quenched after 2 min by dilution of the sample into 2 ml ice-cold 50% (w/v) ammonium sulfate solution. The mixture was filtered rapidly through 0.45 μm pore-size cellulose nitrate filters. The filters were washed once with 2 ml ammonium sulfate solution. Subsequently, the filters were placed in an open plastic vial and dried overnight at room temperature. The radioactivity on the filters was measured via liquid-scintillation counting, using emulsifier plus scintillation liquid (Perkin Elmer).

3.12 Immunodetection

Protein samples were analyzed by 12.5% SDS-PAGE electrophoresis, semi-dry electroblotting and immunodetection with a primary antibody directed against a hexa-His tag (Amersham Pharmacia Biotech). Subsequently, the gels were
submitted to Chemiluminescence detection, using the Western-Light kit (Tropix Inc).

3.13 Miscellaneous
Radiolabeled [N-methyl-\(^{14}\)C] choline chloride (55 mCi/mmol) was obtained from Amersham Biosciences, NJ, USA. The [N-methyl-\(^{14}\)C] choline chloride was used as precursor for the synthesis of [N-methyl-\(^{14}\)C] glycine betaine as described (Landfald, Strom 1986). Radiolabeled [\(^3\)H]-glycine betaine was prepared via a conversion of [\(^3\)H]-choline chloride (Amersham, specific activity: 2.01*10\(^6\) MBq/mmol) to glycine betaine. Creatine kinase and creatine phosphate were obtained from Sigma Chemical Co. All other chemicals were of reagent grade and purchased from commercial sources, unless specified otherwise.
Figure 2: In vivo glycine betaine uptake
(A) Expression levels of ProX and ProW in L. lactis Opu401 co-transformed with pNZcLICproVW and pAMP31-GGGSAGGGS-proX and grown in GM17 and GCDM.
(B) Rate of glycine betaine uptake as a function of sucrose concentration for GCDM- (●) and GM17-grown (○) cells of L. lactis expressing ProVW and ProX. L. lactis Opu401 cells, co-transformed with pNZcLICproVW and pAMP31-GGGSAGGGS-proX, were grown semi-aerobically at 30°C in GM17 and GCDM with 2.5 µg/ml chloroamphenicol plus 2.5 µg/ml erythromycin. Induction with 0.01% (v/v) nisinA was done for 2h, prior to harvesting of the cells for the transport assays. Uptake of [14C]glycine betaine (1 mM, final concentration) was assayed in 50 mM HEPES-methylglucamine pH 7.0 in the presence of 10 mM glucose plus 25 µg/ml chloroamphenicol and erythromycin; the final cell concentration in the assay buffer was 0.5 mg/ml. The initial rates of uptake were determined from the linear parts of the progress curves; measurements were done in duplicate.
4. Results and discussion

4.1 Cloning and functional expression of ProVW and ProX

The osmoregulatory ABC transporter ProU is composed of an ATPase subunit ProV, a TMD subunit ProW and the substrate-binding protein ProX (Gowrishankar 1989). ProX is located in the periplasm of *E. coli*, unlike homologues in Gram-positive bacteria or Archaea where the protein is either fused to a TMD or tethered to the membrane via lipid anchor (Gilson et al. 1988, van der Heide, Poolman 2000). To unleash the properties and functional characterization of ProU in *vitro*, we cloned and expressed *proVW* and *proX* separately to maximize protein yield. For functional complementation *in vivo*, the three proteins were expressed simultaneously in a *L. lactis* strain defective in glycine betaine uptake. The immunoblots of ProX and ProW expressed in *L. lactis*, grown in GCDM and GM17, are shown in figure 2A; ProV is not visible because it lacks a Histag. Figure 2B shows that the rate of glycine betaine uptake via ProU expressed in *L. lactis* increases with the extent of osmotic stress, which is similar to what has been observed when the endogenous OpuA transporter is expressed (Biemans-Oldehinkel, Mahmood & Poolman 2006). However, the absolute rates of glycine betaine uptake via OpuA are approximately 5-fold higher than for ProU, which at least in part reflects a higher expression level of the OpuA transporter (data not shown).

To optimize the expression of ProVW in *E. coli* MC1061/pBADcLICproVW, various inducer (L-arabinose) concentrations, induction times and temperatures were tested. Good expression of ProVW (and avoiding the formation of inclusion bodies; see also (Geertsma et al. 2008a) was obtained with 0.001% L-arabinose at an induction temperature of 30°C and cells grown in LB medium supplemented with 100 μg/ml ampicillin. L-arabinose was added at an OD$_{600}$ ~ 0.5 and the cells were harvested 3 h later. Inside-out membrane vesicles were isolated as described in Materials and methods, and the expression levels were quantified by immunoblotting, using an anti-Histag antibody. On SDS-PAGE, ProV and ProW display molecular masses around 44 and 37 kDa, respectively (figure 3A).

ProX was expressed in *L. lactis* with the following modifications: (i) The endogenous signal sequence was replaced for the one from OppA, which allows efficient lipid modification of the N-terminus of the mature protein; (ii) a flexible linker (GGGSAGGGS) was introduced N-terminal to the mature ProX sequence (the first amino acids are indicated below: ADLPGKGITV) and C-terminal to the cleavage site for the signal sequence (underlined) and a short linker (italics) originating from the OppA protein, yielding: MNKLKVTLLASSVLAATLLSA C GSNQSSS GGGSAGGGSADLPGBKTV as N-terminal sequence; and (iii) a six-histag preceded by a Factor XA cleavage site was engineered at the C-terminus of ProX (added sequence: GSIEGRHHHHHH). As anticipated for lipid-anchored ProX, the protein was found in the membrane fraction of *L. lactis* cells; the identity of the protein was
confirmed by mass spectrometry. The lipid anchor has the advantage that ProX can be tethered to the membrane of liposomes reconstituted with ProVW (figure 1B). As the interaction of substrate-binding proteins with the translocator subunits of Type I ABC transporters is generally of low affinity $K_m > 50 \mu M$ (Doeven et al. 2004, Prossnitz, Gee & Ames 1989, Dean et al. 1992), membrane tethering offers enormous advantages for the functional reconstitution of the protein complex and monitoring of the transport reaction.

**Figure 3**: Expression of ProVW and proX. SDS-PAGE (12% gel) stained with coomassie brilliant blue. Lane 1: Membrane vesicles of *E. coli* harboring pBADcLICproVW; lane 2: Purified ProVW; lane 3: Membrane vesicles of *L. lactis* harboring pAMP31+GGGSAGGGS+proX; lane 4: purified proX; lane 5: proteoliposomes with ProVW; and lane 6: proteoliposomes with ProVW and proX.
4.2 Glycine betaine binding to ProX

Previous studies showed that genuine ProX binds glycine betaine with a $K_D$ of $\sim 1 \mu M$ (Barron, Junga & Villarejo 1987, Haardt et al. 1995). To test whether or not the lipid anchor has an effect on the binding of glycine betaine by ProX, we used the protein precipitation method of (Richarme, Kepes 1983). Figure 4 shows the binding isotherm from which we estimate a $K_D$ of 1–2 $\mu M$. Thus, we conclude that the lipid anchor and His-tag do not affect the functionality of ProX.

Figure 4: Glycine betaine binding to ProX
Glycine betaine binding to purified ProX with lipid anchor and flexible inker. The lipid-anchored ProX was kept soluble by including of 0.05% (w/v) DDM in the assay buffer (100 mM KPI, pH 7.0). See Materials & Methods for further details.

4.3 Purification and membrane reconstitution of ProVW-ProX system

For further characterization of the ProU system, large scale fermentations were performed and ProVW and ProX were purified (figure 3A). As ProV contains 1 cysteine residue in the CBS2 domain and ProW contains 1 cysteine residue in transmembrane helix III, 1 mM of freshly prepared 1 mM DTT was present in each step of membrane vesicle solubilization, protein purification and membrane
reconstitution. In fact, ProU lost activity when DTT was omitted in these steps (data not shown). The reconstitution of ProVW and ProX was carried out in three steps: (i) Incorporation of ProVW into liposomes; (ii) membrane-tethering of ProX; and (iii) enclosing in the proteoliposome lumen of an ATP-regenerating system consisting of creatine-phosphate, creatine kinase and Mg-ATP. The procedure is in essence the same as previously described for the OppBCDF-OppA system (Doeven et al. 2004) and further specified in (Geertsma et al. 2008b), except that in the course of the project, the order of step (ii) and (iii) was reversed. We noted that a higher transport activity, presumably corresponding to a higher reconstitution efficiency, was observed when ProX was tethered to the membrane after the incorporation of the ATP-regenerating system.

4.4 Osmotic activation of membrane-reconstituted ProU

Previous studies have shown that the expression of *proU* is regulated by the osmotic conditions of the medium (May et al. 1986, Lucht, Bremer 1994, Perroud, Le Rudulier 1985). Furthermore, there are indications that the activity of ProU is also affected by osmotic stress (e.g., figure 4 in (Faatz, Middendorf & Bremer 1988). To circumstantiate these findings, we determined the regulation of activity of ProU reconstituted in proteoliposomes. To discriminate binding of glycine betaine to surface-tethered ProX from uptake via ProU, we performed a chase experiment with unlabeled glycine betaine. ProX-bound [\(^{14}\text{C}\)]glycine betaine will be chased upon the addition of an excess of unlabeled glycine betaine, whereas accumulated [\(^{14}\text{C}\)]glycine betaine will not be chased when the transport reaction is unidirectional. Indeed, a more than 100-fold excess of unlabeled glycine betaine to proteoliposomes that had taken up [\(^{14}\text{C}\)]glycine betaine for 3 min (figure 5A) did not result in efflux. Also, the presence of external Mg-ATP did not elicit \(^{14}\text{C}\)-glycine betaine efflux, demonstrating that inside-out-oriented ProVW (without luminal ProX) does not facilitate transport. Figure 5B shows that the rate of glycine betaine is stimulated significantly when the external salt concentration is raised by 300 mM KCl; the basal assay medium is composed of 100 mM KPi, pH 7.0. In this experiment, we monitored the uptake reaction in the absence and presence of 5 mM glycine betaine in the stop buffer. In the absence of glycine betaine in the stop buffer, the offset at the y-axis represents the amount of [\(^{14}\text{C}\)]glycine betaine bound to ProX. The actual uptake (rate) is obtained by subtracting the bound glycine betaine or by including 5 mM glycine betaine in the stop buffer. Figure 5B shows that rapid binding is followed by true transport (accumulation) of [\(^{13}\text{C}\)]glycine betaine. The rate of uptake increases with the salt concentration and a sigmoidal dependence was observed (figure 5C), similar to the activation of OpuA (van der Heide, Stuart & Poolman 2001).
4.5 Effect of anionic lipids on ProU activity

The experiments described heretofore were carried out with ProU reconstituted in liposomes composed of *E. coli* polar lipids plus L-α-phosphatidyl choline from egg yolk in a ratio of 3:1 (w/w). The fraction of phosphatidylethanolamine in *E. coli* can be up to 75%; the remaining lipids are mostly anionic, i.e., phosphatidylglycerol (PG) and cardiolipin, typically at 20 and 5 mole% (Cronan 2003). The relative amounts of PE, PG and cardiolipin vary with osmotic stress as shown by (Romantsov, Guan & Wood 2009). As osmotic (ionic) regulation of OpuA has been shown to be critically dependent on the fraction of anionic lipids (Biemans-Oldehinkel, Mahmood & Poolman 2006), we reconstituted ProU in 50 mole% of DOPE (zwitterionic, non-bilayer-forming lipid) plus varying ratios of DOPG (anionic, bilayer-forming lipid) and DOPC (zwitterionic, bilayer-forming lipid). Figure 6 shows that ProU activity was not affected by the osmotic stress, i.e., KCl concentration in the range of 0–300 mM in assay buffer composed of 100 mM KPi, pH7.0 (above 300 mM KCl the activity decreased) when DOPG was absent or present at 12 mole%. Importantly, osmotic stress-activated activity was observed at 25 mole% of DOPG and even more so at 38 mole%. The activation was instantaneous, that is, within seconds following the addition of threshold levels of KCl. Although the fraction of anionic lipids in *E. coli* can vary somewhat (Romantsov, Guan & Wood 2009), such long-term adaptation is probably not relevant for osmoregulation via ProU. Instead, the anionic lipids are needed to make the ionic-gating mechanism of ProU effective. The above findings are reminiscent of what we observed for OpuA from *L. lactis* (Biemans-Oldehinkel, Mahmood & Poolman 2006), and they suggest that electrostatic interactions between the anionic membrane surface and a protein domain are intrinsic to the osmotic gating of the transporter. Moreover, with 38 mole% of DOPG, maximal ProU activity was observed at 300–400 mM KCl added to the basal assay medium, which is well in the range where the transporter needs to be active in vivo. For instance, the steady state levels of K$^+$ ions increase from 0.15 M to more than 0.5 M when the osmolarity of the medium is increased from 0.1–1.2 Osm (Epstein, Schultz 1965, Sutherland et al. 1986, Cayley et al. 1991).
Functional reconstitution of ProU in proteoliposomes

Figure 5: Glycine betaine uptake in proteoliposomes
ProU was reconstituted in liposomes composed of E. coli polar lipids and L-α-phosphatidylcholine from egg yolk in a ratio of 3:1 (w/w); the final ProVW concentration was 0.05 mg/ml.

(A) $[^{14}\text{C}]$-glycine betaine uptake (40 µM, final concentration) assayed in 100 mM KPi, pH 7.0 plus 300 mM KCl with an ATP-regenerating system enclosed in the vesicle lumen, composed of E. coli polar lipids plus L-α-phosphatidylcholine from egg yolk in a ratio of 3:1 (w/w). Time course of uptake (●); after 3 min of uptake 5 mM Mg-ATP (▼) or 5 mM glycine betaine (○) was added to the assay mixture.

(B) $[^{14}\text{C}]$-glycine betaine uptake (40 µM, final concentration) assayed in 100 mM KPi, pH 7.0, under iso-osmotic (△, ▲) and hyperosmotic (◇, ●) conditions (100 mM KPi + 300 mM KCl); with (open symbols) and without (closed symbols) 5 mM glycine betaine in the stop buffer.

(C) Rate of glycine betaine uptake as a function of KCl concentration, added to 100 mM KPi, pH 7.0 assay buffer.
Chapter 3

Figure 6: Effect of anionic lipids on glycine betaine uptake
ProU was reconstituted in liposomes composed of 50 mol% DOPE and 0 (■), 12 (▲), 25 (▼), 38 (○), 50 (●) mol% DOPG, plus 50, 38, 25, 12 and 0 mol% DOPC. The ATP-regenerating system was enclosed in the vesicle lumen; the final ProVW concentration in the assay medium was 0.05 mg/ml. The rate of [14C]-glycine betaine uptake (40 µM, final concentration) was assayed in 100 mM KPi, pH 7.0, and different concentrations of KCl.

5. Conclusion
The glycine betaine transporter ProU from *E. coli* was functionally expressed (in *L. lactis* and *E. coli*), and the components were purified and reconstituted in proteoliposomes. Lipid-tethering of ProX allowed us to observe glycine betaine uptake in ProVW-containing proteoliposomes. The uptake of glycine betaine in whole cells and in proteoliposomes is regulated by osmotic stress, presumably via an increase in the internal salt concentration as observed for OpuA from *L. lactis* (ionic strength-gating) and BetP from *C. glutamicum* (K⁺-gating). We have not explicitly demonstrated salt-gating of ProU but the parallels with the OpuA system are striking, i.e., for both systems the fraction of anionic lipids is critical for osmoregulated transport, the activity increases with the salt concentration, and both have a CBS module linked to the NBD that has been implicated in osmosensing. The salt dependencies of ProU and OpuA differ somewhat as do the intracellular concentrations of K⁺ and the fractions (and types) of anionic lipids in the corresponding parent cells, *E. coli* and *L. lactis*, respectively. We speculate that the mechanism of osmotic regulation of ProU in *E. coli* is in essence similar to what has been observed for OpuA in *L. lactis*. The two transporters have likely evolved
to match the physicochemical conditions (internal ionic strength, K⁺ concentration, water activity, lipid composition) that *E. coli* and *L. lactis* face. We are currently exploring the possibility to crystallize the OpuA and ProU proteins to gain further insight in the mechanism of transport and regulation of these ABC transporters.

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