Engineering of Ion Sensing by the Cystathionine $\beta$-Synthase Module of the ABC Transporter OpuA*

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We have previously shown that the C-terminal cystathionine $\beta$-synthase (CBS) domains of the nucleotide-binding domains of the ABC transporter OpuA, in conjunction with an anionic membrane surface function, act as sensor of internal ionic strength ($I_m$). Here, we show that a surface-exposed cationic region in the CBS module domain is critical for ion sensing. The consecutive substitution of up to five cationic residues led to a gradual decrease of the ionic strength dependence of transport. In fact, a 5-fold mutant was essentially independent of salt in the range from 0 to 250 mM KCl (or NaCl), supplemented to medium of 30 mM potassium phosphate. Importantly, the threshold temperature for transport was lowered by 5–7 °C and the temperature coefficient $Q_{10}$ was lowered from 8 to 1.5 in the 5-fold mutant, indicating that large conformational changes are accompanying the CBS-mediated regulation of transport. Furthermore, by replacing the anionic C-terminal tail residues that extend the CBS module with histidines, the transport of OpuA became pH-dependent, presumably by additional charge interactions of the histidine residues with the membrane. The pH dependence was not observed at high ionic strength. Altogether the analyses of the CBS mutants support the notion that the osmotic regulation of OpuA involves a simple biophysical switching mechanism, in which nonspecific electrostatic interactions of a protein module with the membrane are sufficient to lock the transporter in the inactive state.

In their natural habitats microorganisms are often exposed to changes in the concentration of solutes in the environment (1). A sudden increase in the medium osmolality results in loss of water from the cell, loss of turgor, a decrease in cell volume, and an increase in intracellular osmolyte concentration. Osmoregulatory transporters such as OpuA in Lactococcus lactis, ProP in Escherichia coli, and BetP in Corynebacterium glutamicum diminish the consequences of the osmotic stress by mediating the uptake of compatible solutes upon an increase in extracellular osmolality (2–4). For the ATP-binding cassette (ABC)5 transporter OpuA, it has been shown that the system, reconstituted in proteoliposomes, is activated by increased concentrations of luminal ions (increased internal ionic strength) (2, 5, 6). This activation is instantaneous both in vivo and in vitro and only requires threshold levels of ionic osmolytes. Moreover, the ionic threshold for activation is highly dependent of the ionic lipid content (charge density) of the membrane and requires the presence of so-called cystathionine $\beta$-synthase (CBS) domains, suggesting that the ionic signal is transduced to the transporter via critical interactions of the protein with membrane lipids.

The ABC transporter OpuA consists of two identical nucleotide-binding domains (NBD) fused to CBS domains and two identical substrate-binding domains fused to transmembrane domains. The NBD-CBS and substrate-binding domain-transmembrane domain subunits are named OpuAA and OpuABC, respectively. Two tandem CBS domains are linked to the C-terminal end of the NBD; each domain (CBS1 and CBS2) has a $\beta$-a-$\beta$-a secondary structure (5) (Fig. 1A). The CBS domains are widely distributed in most if not all species of life but their function is largely unknown. Most of the CBS domains are found as tandem repeats but data base searches have also revealed tetra-repeat units (5). The crystal structures of several tandem CBS domains have been elucidated (7–9, 32), and in a number of cases it has been shown that two tandem CBS domains form dimeric structures with a total of four CBS domains per structural module (hereafter referred to as CBS module). The crystal structures of the full-length MgtE Mg$^{2+}$ transporter confirm the dimeric configuration and show that the CBS domains undergo large conformational changes upon Mg$^{2+}$ binding or release (10, 11). In general, ABC transporters are functional as dimers, which implies that two tandem CBS domains are present in the OpuA complex. Preliminary experiments with disulfides engineered at the interface of two tandem CBS domains in OpuA suggest that large structural rearrangements (association-dissociation of the interfaces) play a determining role in the ionic strength-regulated transport. Finally, a subset of CBS-containing proteins has a C-terminal extension, which in OpuA is highly anionic (sequence: ADIDPD-EDEVEEEIKEEENK) and modulates the ion sensing activity (6).

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5 The abbreviations used are: ABC, ATP-binding cassette; CBS, cystathionine $\beta$-synthase; NBD, nucleotide-binding domain; DOPE, dioleoyl-phosphatidylethanolamine; DOPG, dioleoyl-phosphatidylglycerol; DOPC, dioleoyl-phosphatidylcholine; PIP$_2$, phosphatidylinositol bisphosphate; MARCKS, myristoylated alanine-rich protein kinase C substrate; PA, phosphatidic acid; LPA, lysophosphatidic acid.
Ion Sensing by the CBS Module

In this study, we have engineered the surface-exposed cationic residues of the CBS module and the C-terminal anionic tail of OpuA (Fig. 1B). The ionic strength and lipid dependence of the OpuA mutants were determined in vivo and in vitro. We show that substitution of five cationic residues for neutral amino acids is sufficient to inactivate the ionic strength sensor and convert OpuA into a constitutively active transporter. Moreover, by substituting six anionic plus four neutral residues of the C-terminal anionic tail for histidines, the transport reaction becomes strongly pH-dependent.

MATERIALS AND METHODS

Plasmid Construction

Site-specific mutations in the CBS part of the opuAA gene were made via the megaprimer approach. Briefly, specific megaprimer oligonucleotides were obtained by using a specific forward oligonucleotide (Table 1) and the general reverse primer EOL0404 5′→3′ and plasmid pNZOpuA(PstI)His as template. pNZOpuA(PstI)His was created previously (5) and contains an additional PstI site at the 3′ end of opuAA to facilitate the swapping of a 798-bp Clal-PstI fragment. The corresponding protein, hereafter referred to as OpuA(parental), has two mutations at the very end of the C terminus (Table 1), but its activity is identical to that of wild type OpuA. The megaprimer was used as reverse primer in a second amplification reaction with the forward primer EOL0606 5′→3′. After digestion with Clal and PstI, the parental gene fragment of pNZOpuA-(PstI)His was exchanged for the mutated fragment, yielding the specific CBS domain mutant (Table 1). To create OpuA[K1], the plasmid carrying opuA(parental) was used as template; for OpuA[K2], the plasmid carrying opuA[K1] was used, and so forth.

To replace part of the negatively charged C terminus of OpuAA with histidines, pNZOpuA(D12)His was digested with PstI and an annealed oligonucleotide, encoding 10 histidine residues and containing PstI compatible ends, was inserted, resulting in pNZOpuA(insHis)His. pNZOpuA(D12)His lacks the main part of the C-terminal anionic tail of OpuAA (5). Primers used to create the annealed oligonucleotide, encoding 10 histidine residues and containing PstI compatible ends, was inserted, resulting in pNZOpuA(insHis)His. pNZOpuA(D12)His lacks the main part of the C-terminal anionic tail of OpuAA (5). Primers used to create the annealed oligonucleotide, encoding 10 histidine residues and containing PstI compatible ends, was inserted, resulting in pNZOpuA(insHis)His. pNZOpuA(D12)His lacks the main part of the C-terminal anionic tail of OpuAA (5). Primers used to create the annealed oligonucleotide, encoding 10 histidine residues and containing PstI compatible ends, was inserted, resulting in pNZOpuA(insHis)His. pNZOpuA(D12)His lacks the main part of the C-terminal anionic tail of OpuAA (5).

Bacterial Strains, Growth Conditions, and Vesicle Preparation

L. lactis strain Opu401 (4) was cultivated seminaerobically at 30°C in a medium containing 2% (w/v) gisetex LS (Strik BV, Eemnes, NL) and 65 mM sodium phosphate, pH 6.5, supplemented with 1.0% (w/v) glucose and 5 μg/ml chloramphenicol when carrying pNZOpuAHis or derivatives. For isolation of membrane vesicles, cells were grown in a 2-liter pH-regulated bioreactor to an A_{600} of 2, after which transcription from the nisA promoter was switched on by the addition of 0.1% (v/v) culture supernatant of the nisin A-producing strain NZ9700. The cells were harvested and membrane vesicles were prepared according to standard procedures (12). The protein concentration was determined using the DC Protein Assay (Bio-Rad).

Synthesis of Hybrid Membranes

Membrane vesicles of the L. lactis Opu401/pNZOpuA derivative were isolated as described (13), and fused 1:10 with preformed liposomes, composed of dioleoyl-phosphatidylethanolamine (DOPE), dioleoyl-phosphatidylglycerol (DOPG), and dioleoyl-phosphatidylcholine (DOPC). The hybrid membranes were obtained by two freeze/thaw cycles, followed by extrusion through a polycarbonate filter (200 nm pore size); for details on the preparation of liposomes and the freeze-thaw-extrusion steps, we refer to Geertsma et al. (14). The fraction of DOPE was always 50 mol % and the mole fractions of DOPG and DOPC were varied reciprocally; for most experiments we used either 38 mol % DOPG, 12 mol % DOPC or 12 mol % DOPG, 38...
Ion Sensing by the CBS Module

mol % DOPC. The synthetic lipids were obtained from Avanti Polar Lipids, AL.

Purification and Membrane Reconstitution of OpuA

Purified OpuA and mutant derivatives were reconstituted in liposomes composed of synthetic lipids (see “Synthesis of Hybrid Membranes”), essentially as described previously (14). The final protein to lipid ratio was 1:100 (w/w), unless stated otherwise.

Transport Assays

In Vivo Uptake—Opu401 cells carrying the pNZOpuA-derived plasmids were grown in M17 supplemented with 0.5% glucose and 5 μg/ml chloramphenicol to an A600 of 0.8. For induction of the opuA genes, 2 × 10^{-4} % (v/v) culture supernatant of the nisin A producing strain NZ9700 was used. After induction, the cells were washed twice with ice-cold 50 mM K-HEPES, pH 7.3. Prior to initiation of transport, cells were pre-energized for 5 min with 10 mM glucose at 30 °C. Uptake of [14C]glycine betaine was assayed in 50 mM K-HEPES, pH 7.3, supplemented with 50 μg/ml chloramphenicol and 10 mM glucose, with or without sucrose as indicated. The concentration of cells in the uptake assay was ~0.4 mg of total protein/ml.

Hybrid Membranes—For ATP-driven uptake of glycine betaine in hybrid membranes, the ATP-regenerating system was enclosed during the fusion of membrane vesicles with liposomes (14); the ATP-regenerating system corresponds to 10 mM Na2-ATP + 10 mM MgSO4 + 24 mM Na2-creatine phosphate + 2.4 mg/ml creatine kinase in 50 mM potassium P0, pH 7.0. The fused and loaded hybrid membranes were washed twice with 90 mM potassium P0, pH 7.0 (osmolality of approximately 240 mosmol/kg), and resuspended in the same buffer to a concentration of 60 mg of lipids/ml. For osmotically activated transport, the hybrid membranes were diluted to a lipid concentration of 4.5–12 mg/ml into assay buffer (90 mM potassium P0, pH 7.0 unless specified otherwise), supplemented with different concentrations of KCl. Following incubation for 2 min at 30 °C, the transport reaction was initiated by the addition of [14C]glycine betaine (Amersham Biosciences) to a final concentration of 31 μM (more than 10-fold above the K0.5 for transport and K0.5 for binding) (13). At given time intervals, 40-μl samples were taken and diluted with 2 ml of ice-cold isotonic assay buffer. The samples were filtered rapidly through 0.45-μm pore-size cellulose nitrate filters (Schleicher & Schuell) and washed twice with 2 ml of assay buffer. The radioactivity on the filters was determined by liquid scintillation counting.

Proteoliposomes—ATP-driven uptake of glycine betaine by right-side in reconstituted OpuA was performed as described by van der Heide and Poolman (12), with some modifications. Briefly, proteoliposomes were loaded with 9 mM Mg-ATP (prepared from 9 mM MgSO4 plus 9 mM Na2-ATP), 10 mM potassium P0, pH 7.0, unless stated otherwise. The inclusion of the Mg-ATP/potassium P0 was done via two cycles of freeze-thawing. After extrusion of the proteoliposomes through a polycarbonate filter (200 nm pore size), the proteoliposomes loaded with Mg-ATP were washed twice with 30 mM potassium P0, pH 7.0, and resuspended in the same buffer unless specified otherwise (30 mM potassium P0, pH 7.0 is isoosmolar with the luminal contents and corresponds to an osmolality of ~70 mosmol/kg as determined by freezing point depression) to a concentration of 80 mg of lipids/ml.

For osmotically activated transport, the proteoliposomes were diluted to a lipid concentration of 5 mg/ml into assay buffer (30 mM potassium P0, pH 7.0, supplemented with different concentrations of KCl). Following incubation for 2 min at 30 °C, the transport reaction was initiated by the addition of [14C]glycine betaine (Amersham Biosciences) to a final concentration of 42.5 μM. At given time intervals, 40-μl samples were taken and diluted with 2 ml of ice-cold isotonic assay buffer and processed further as described under “Hybrid Membranes.”

The initial rates of uptake were calculated from the linear part of the transport curves, using linear regression. Radiolabeled N-[methyl-14C]choline chloride (55 mCi/mmol) was from Amersham Biosciences, and was converted to N-[methyl-14C]glycine betaine as described (15).

RESULTS

Engineering Strategy

We have previously shown that deletion of the CBS2 domain (OpuAΔ61) is sufficient to abolish most of the ionic strength dependence of OpuA (5). A thorough analysis of the mutant with the entire CBS module deleted (OpuAΔ119) has not been reported. We now show that the activity of OpuAΔ119 is independent of salt in the 0–250 mM KCl range (or NaCl), present in the basal assay medium of 30 mM potassium P0, pH 7.0 (Fig. 2). For comparison, the ionic activation profiles of parental OpuA

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14370 JOURNAL OF BIOLOGICAL CHEMISTRY

VOLUME 284 • NUMBER 21 • MAY 22, 2009
and OpuΔ61 are shown. For OpuΔ61 and OpuΔΔ119, part of the OpuA subunit was lost in the reconstitution process (observed as lower ratio of AA/ABC subunits on SDS-PAGE), which explains at least part of the lower activity (i.e. at 250 mM KCl) of these mutants. To gain further insight into the role of the CBS domains in the mechanism of ionic activation, a series of more subtle mutants were designed and constructed: (i) surface-exposed cationic residues in the CBS module were substituted for neutral amino acids; and (ii) anionic residues in the C-terminal tail of the CBS domains were changed to histidines.

On the basis of a homology-based structure model of the OpuA CBS domains (using a multiple sequence alignment and the crystal structure of the tandem CBS protein Ta0289 from Thermoplasma acidophilum), we identified five cationic residues (Lys<sup>313</sup>, Lys<sup>314</sup>, Arg<sup>315</sup>, Arg<sup>318</sup>, and Lys<sup>377</sup>) as potential candidates for mediating the anionic lipid-dependent ionic strength response. The cationic residues 313–315 and Arg<sup>318</sup> are located in the loop between β2 and β3 of CBS1 and Lys<sup>377</sup> is located in the corresponding loop of CBS2. Single [K1], double [K2], triple [K2R], quadruple [K2R2], and quintuple [K3R2] mutants were constructed and lys and/or arg residues were substituted for neutral amino acids (Table 1). The mutants were made and characterized in two rounds of mutagenesis, i.e. K1, K2, and K2R were constructed first and based on the <i>in vivo</i> and <i>in vitro</i> activity data, K2R2 and K3R2 were designed and analyzed subsequently. Each of the mutants expressed to levels of at least 50% of parental OpuA (e.g. Fig. 3A).

**Assay Strategy**

Transport activity was assessed <i>in vivo</i>, using whole cells of<i> L. lactis</i> osmotically stressed with sucrose, and <i>in vitro</i>, using hybrid membranes or proteoliposomes. Although the internal ionic strength in whole cells can be increased by increasing the osmotic stress, the absolute values are difficult to quantify and low values of internal ionic strength cannot be achieved, that is, without compromising the physiological well being of the organism. In hybrid membranes and proteoliposomes, the internal ionic strength can be set precisely and varied by the addition of salt or sucrose to the external medium. Hybrid membranes offer the advantage that the mutant proteins do not need to be purified, which can be desirable for mutants that are less stable. <i>L. lactis</i> membrane vesicles contain 50–60 mol % anionic lipids (16), and, after a 1 to 10 fusion with liposomes composed of 38 mol % DOPG, the fraction of anionic lipids was ~40 mol %. Full control over lipid and protein composition and ionic strength was obtained in proteoliposomes, in which purified OpuA was incorporated into membranes composed of synthetic lipids.

**Surface-exposed Cationic Residues**

<i>Single, Double, and Triple Mutants—In vivo</i> OpuA[K1], OpuA[K2], and OpuA[K2R] displayed different activation profiles compared with parental OpuA (Fig. 3B). Removing a single cationic residue, OpuA[K1], increased the activity at low osmotic stress, that is, conditions corresponding to lower values of intracellular ionic strength, which is indicative of a lower degree of inhibition by anionic lipids. Removing two cationic residues, OpuA[K2], led to a further increase in activity at low osmotic stress, and by removing all three cationic residues, OpuA[K2R], the transporter became essentially independent of the imposed osmotic stress. The activity of the three mutants at high osmotic stress (0.65 M sucrose in Fig. 3B) was lower than that of the parental transporter, but this point was not studied further <i>in vivo</i>.

Next, membrane vesicles bearing parental OpuA, OpuA[K1], OpuA[K2], or OpuA[K2R] were isolated and fused 1:10 (w/w in lipids) with liposomes containing 38 mol % DOPG, 12 mol % DOPC plus 50% mol % DOPE (Fig. 4). It has been previously

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**TABLE 1**

List of OpuA modifications and primers used in this study

<table>
<thead>
<tr>
<th>Protein</th>
<th>Modifications</th>
<th>Primers/Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>OpuA</td>
<td>Wild type</td>
<td>Van der Heide and Poolman (12)</td>
</tr>
<tr>
<td>OpuA(PstI) (Parental)</td>
<td>E402A, K403A</td>
<td>Biemans-Oldenhinkel et al. (5)</td>
</tr>
<tr>
<td>OpuAΔ12</td>
<td>Δ392–401, E402A, K403A</td>
<td>Biemans-Oldenhinkel et al. (5)</td>
</tr>
<tr>
<td>OpuAΔ61</td>
<td>Deletion of CBS2</td>
<td>Biemans-Oldenhinkel et al. (2006)</td>
</tr>
<tr>
<td>OpuAΔ119</td>
<td>Deletion of CBS1 and 2</td>
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</table>

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**FIGURE 3.** In vivo characterization of CBS surface charge mutants of OpuA. A, visualization of the expression of the CBS mutants in membrane vesicles. Parental OpuA and CBS mutants were expressed in <i>L. lactis</i> Opu401 and 6.5 μg of membrane vesicles was loaded per lane on a 12.5% SDS-PAGE gel. B, in vivo activation profiles of parental OpuA and single (K1), double (K2), and triple (K2R) surface charge mutants. Transport of [<sup>14</sup>C]glycine betaine by wild type OpuA (●), OpuA[K1] (○), OpuA[K2] (▲), and OpuA[K2R] (▲). was assayed in 50 mM K-HEPES, pH 7.3, supplemented with 10 mM glucose, with or without sucrose as indicated.
Ion Sensing by the CBS Module

FIGURE 4. In vitro characterization of CBS surface charge mutants of OpuA in hybrid membranes. Membrane vesicles of L. lactis bearing OpuA mutants were fused with liposomes composed of 50 mol % DOPG, 12 mol % DOPS, and 38 mol % DOPG. The ATP-regenerating system was enclosed inside the hybrid membranes. Uptake of [14C]glycine betaine by parental OpuA (○), OpuA[K1] (△), OpuA[K2] (▲), and OpuA[K2R] (▼) in hybrid membranes was assayed in 90 mM potassium P, pH 7.0, with or without added KCl as indicated on the x axis. Activation profiles in A show the measured values, whereas activation profiles in B represent normalized rates: data were normalized relative to the highest activity in the concentration range of 0 to 0.3 M KCl.

The deregulated activity could be due to a lower degree of inhibition by anionic lipids. To address this point, we determined the anionic lipid dependence of OpuA by comparing proteoliposomes with either 38 (panel C) or 12 (panel D) mol % of DOPG (Fig. 5). The fraction of the zwitterionic lipid DOPC was varied reciprocally with DOPG, and the fraction of DOPE was kept constant at 50%. At low (12 mol %) DOPG, the activities of parental OpuA and OpuA[K3R2] were similar and decreased slightly with increasing ionic strength (Fig. 5D). For comparison, the same batches of protein in liposomes with 38 mol % of DOPG revealed the strong ionic strength-dependence of parental OpuA and the ionic strength-independence of OpuA[K3R2] (Fig. 5C). Taken together, these data strongly suggest that the cationic amino acid residues participate in interaction of the CBS module with the membrane, the more cationic residues, the tighter the binding and thus the more salt one needs to screen the electrostatic interactions.

We also measured the temperature dependence of parental OpuA, OpuA[Δ119], and OpuA[K3R2] to determine the activation energies for (the regulation of) transport. Fig. 6 shows the initial rates of glycine betaine uptake from 7 to 40 °C (panel A, parental OpuA and OpuA[K3R2]) and from 7 to 30 °C (panel B, parental OpuA and OpuA[Δ119]); the corresponding Arrhenius plots are shown in Fig. 6C. Parental OpuA was only active in the presence of salt (I in 0.6 in this particular experiment; closed circles) and above a temperature of 13–14 °C; the low salt concentration at which the system was not active corresponded to I in 0.06 (open circles). The temperature coefficient Q 10 of parental OpuA was about 8 between 15 and 30 °C, yielding an activation energy (E a) of 148 kJ/mol. As anticipated, OpuA[Δ119] was equally active at low and high salt and had a somewhat lower E a of 119 kJ/mol compared with parental OpuA. Importantly, significant activity of OpuA[K3R2] was already observed at temperatures below 10 °C and the activity was relatively insensitive to temperature both at high (I in 0.6) and low (I in 0.06) ionic strength. The Q 10 of OpuA[K3R2] was only 1.5 and corresponded to an E a of 29 kJ/mol. The temperature dependences of the parental OpuA, OpuA[Δ119], and the 5-fold mutant OpuA[K3R2] indicate that deletion of the CBS module or substitution of the surface-exposed cationic residues, both leading to ionic strength-independent transport, have different consequences for the activation energy of the translocation reaction. Depending on the modification made to the CBS module, the transporter can be in a deregulated-low (OpuA[Δ119]) or deregulated-high energy state (OpuA[K3R2]).

C-terminal Tail

It has been previously shown that deleting 12 amino acids of the anionic tail C-terminal of the CBS domains shifts the I in dependence of OpuA to higher values (OpuA[Δ12]) (4), indicating that the anionic tail attenuates the ion sensing function. The role of the anionic C terminus of OpuAA was further explored by substituting 6 of the acidic plus 4 neutral residues of the anionic C terminus for 10 histidines, yielding OpuAΔ12insH119 (Table 1). The pH dependence of transport of parental OpuA, OpuA[Δ12], and OpuA[Δ12insH119] were determined as a function of pH at low (I in 0.06), intermediate (I in 0.3), and high (I in 0.5) internal ionic strength. Assuming an average pK a of about 6.4, the decahistidine sequence will have a charge of +9 to +10 at pH 5.5 and be lower than +1 at pH 7.5. The pH will thus strongly affect the overall charge of the CBS module. To avoid possible interference of the pH on the regen-
DOPG at 1:50 of protein to lipid ratio. Proteoliposomes were loaded with 9 mM MgSO₄, 9 mM Na₂ATP plus 10 mM i

In vitro characterization of CBS surface charge mutants of OpuA proteoliposomes. A, visualization of the proteoliposomes on SDS-PAGE gel. 35 µl of proteoliposomes were treated with 2% (v/v) SDS plus 5× sample buffer (3% (w/v) Tris-HCl, 50% (v/v) glycerol, 0.005% (v/v) bromphenol blue, 1% (v/v) SDS, 3% (v/v) β-mercaptoethanol). 30 µl of the mixture was applied to a 12.5% SDS-PAGE gel and electrophoresed for 90 min at constant current of 15 mA. B, in vitro transport assay, using purified protein reconstituted in proteoliposomes. The proteoliposomal membranes were composed of 50 mol % DOPE, 12 mol % DOPC, and 38 mol % DOPG at 1:50 of protein to lipid ratio. Proteoliposomes were loaded with 9 mM MgSO₄, 9 mM Na₂ATP plus 10 mM potassium Pi, which is equiosmolar with 30 mM potassium Pi. Uptake of [¹⁴C]glycine betaine by parental OpuA ( ), OpuA (K2) ( ), OpuA (K2R) ( ), OpuA (K1) ( ), K2R2 ( ), and OpuA (K3R2) (234). Uptake of [¹⁴C]glycine betaine was assayed in 30 mM potassium Pi, pH 7.0, with or without added KCl (0–250 mM). Normalized rates of transport are shown; the actual uptake rates at 0.25 M KCl were 4.5, 30, 22, 20, 21, and 19 nmol/min per mg of protein for parental OpuA, K1, K2, R2, K2, and K3R2, respectively. C and D, lipid and salt dependence. Panel C, proteoliposomes composed of 50 mol % DOPE, 12 mol % DOPC, and 38 mol % DOPG; parental OpuA ( ) and OpuA (K3R2) ( ). Panel D, proteoliposomes composed of 50 mol % DOPE, 38 mol % DOPC, and 12 mol % DOPG; parental OpuA ( ) and OpuA (K3R2) ( ). Further conditions are as described under B.

Ion Sensing by the CBS Module

In the 5-fold mutant (K3R2) the transport activity was essentially insensitive to pH at pH 5.5 to 7.5. OpuA (K3R2) (234). Uptake of [¹⁴C]glycine betaine was assayed in 30 mM potassium Pi, pH 7.0, with or without added KCl (0–250 mM). Normalized rates of transport are shown; the actual uptake rates at 0.25 M KCl were 4.5, 30, 22, 20, 21, and 19 nmol/min per mg of protein for parental OpuA, K1, K2, R2, K2, and K3R2, respectively. C and D, lipid and salt dependence. Panel C, proteoliposomes composed of 50 mol % DOPE, 12 mol % DOPC, and 38 mol % DOPG; parental OpuA ( ) and OpuA (K3R2) ( ). Panel D, proteoliposomes composed of 50 mol % DOPE, 38 mol % DOPC, and 12 mol % DOPG; parental OpuA ( ) and OpuA (K3R2) ( ). Further conditions are as described under B.

Ion Sensing by the CBS Module

In proteoliposomes with 18 mol % of DOPG, the pH-dependent activation was no longer observed and the pH profiles of OpuA, OpuAΔ12, and OpuAΔ12insHis were very similar at every Iₘ (data not shown), suggesting that pH-dependent switching observed in Fig. 7A (and to a lesser extent Fig. 7B) requires the interaction of the (modified) CBS module with the membrane, either directly or via an additional factor in the core domains of OpuA.

DISCUSSION

We have previously shown that the ABC transporter OpuA is activated when the intracellular ionic strength reaches a threshold value. Below the threshold, the transporter is kept in the inactive “electrostatically locked” state, presumably through interactions of a protein domain with the anionic membrane surface. This mechanism is based on the following observations: (i) the ionic strength needed for activation increases with the fraction of anionic lipids in the membrane; (ii) at 40–60 mol % of DOPG, the transporter is inactive below 100–200 mM salt, which nicely matches the physiological conditions at which the system needs to be “on” or “off”; (iii) OpuA is no longer regulated by ionic strength when the CBS module is deleted; (iv) deletion of the anionic C-terminal tail reinforces the ionic regulation of transport, indicating that this amino acid sequence attenuates the pH-dependent activation suggested in Fig. 7A (and to a lesser extent Fig. 7B) requires the interaction of the (modified) CBS module with the membrane, either directly or via an additional factor in the core domains of OpuA.

Strength when the CBS module is deleted; (iv) deletion of the anionic C-terminal tail reinforces the ionic regulation of transport, indicating that this amino acid sequence attenuates the pH-dependent activation suggested in Fig. 7A (and to a lesser extent Fig. 7B) requires the interaction of the (modified) CBS module with the membrane, either directly or via an additional factor in the core domains of OpuA.
Ion Sensing by the CBS Module

![Graphs showing temperature dependence of transport activity](image)

**FIGURE 6.** Temperature dependence of the transport activity of parental OpuA (panels A and B), OpuA[K3R2] (panel A), and OpuAΔ119 (panel B) at low and high internal ionic strength. Proteoliposomes with parental OpuA (C), OpuA[K3R2] (Δ), and OpuAΔ119 (■) were composed of 50 mol% DOPE, 12 mol% DOPC, and 38 mol% DOPG and were loaded with 9 mM MgSO\(_4\), 9 mM Na\(_2\)ATP plus 10 mM potassium Pi. Uptake of \(^{14}\)Cglycine betaine was assayed in 30 mM potassium Pi, pH 7.0, with (closed symbols) or without (open symbols) 250 mM KCl. Error bars show standard deviations of average rates from 3 independent experiments. Panel C is the Arrhenius transformation of the data from panels A and B.

The observation that OpuAΔ12insHis is active at alkaline pH, whereas parental OpuA and OpuAΔ12 are not (Fig. 7A), suggests that at alkaline pH the mere presence of the decahistidine sequence is sufficient for activation. Upon lowering of the pH, the decahistidine tail reinforces the electrostatic interaction of the CBS module with the membrane, which inactivates OpuAΔ12insHis. We emphasize that the decahistidine sequence is most likely unstructured and modulates the transporter without being an inherent part of the CBS ion-sensing module, in line with the concept of an unspecific electrostatic switching mechanism.

**Activation Energy for Transport**

The temperature dependence of transport revealed an unexpected mechanistic feature of the ionic regulation of OpuA. The activation energy for transport by parental OpuA in the activated state was relatively high (Q\(_{10}\)~ 8), indicative of large conformational transitions in the catalytic cycle of transport. To the best of our knowledge, activation energies for translocation by ABC transporters have not been reported. For secondary transporters, Q\(_{10}\) values of 2–3 have been described, whereas for flux through channel proteins Q\(_{10}\) is <1.5 (17–19). The high Q\(_{10}\) values suggest that the driving force for the transition between different states must have a large enthalpic contribution (e.g. Brownian motion of protein domains). Surprisingly, even though OpuAΔ119 and OpuA[K3R2] are completely deregulated in terms of ionic activation, the temperature dependences and activation energies were very different. OpuAΔ119, lacking the CBS module, had a Q\(_{10}\)~ 5. OpuA[K3R2] had a highly reduced activation energy (Q\(_{10}\)~ 1.5), suggesting that the transporter is in a state where the proposed large conformational transition is bypassed. Thus, OpuAΔ119 and OpuA[K3R2] seem to reflect different energy
states, with the K3R2 mutations trapping the system in an activated state (Fig. 8).

The range of screened electrostatic interactions is defined by Debye length, usually denoted by $1/\kappa$ (20, 21). Because the Debye length is essentially independent of temperature (20, 21), it is not likely that the temperature dependence is related to the electrostatic interaction of the ion-sensing module with the membrane. Thus, at 250 mM KCl plus 30 mM potassium Pi, pH 7.0, in the external medium (yielding an $I_{in} \sim 0.6$), the electrostatic interactions are sufficiently screened and similar at every temperature. The substitution of five cationic residues thus seems to have dual effects: the interaction of the ion sensing module with the membrane is decreased (transport has become independent of $I_{in}$ and is no longer anionic lipid-regulated) and the activation barrier for transport is lowered. Although both phenomena are most likely interrelated, the ionic activation of transport must involve more than a release of the CBS module from the membrane. Whatever the precise structural rearrangements may be, the CBS module in conjunction with anionic lipids play a central role in osmotic, i.e. ionic strength, sensing.

How Universal Is “Simple” Electrostatic Control of Membrane Transport?—Within the potassium channels, human TREK-1 and the archaeal MJK$_4$ (22, 23), protein-lipid interaction serves to regulate the activity in response to membrane stretch and cell swelling (hypo-osmotic stress). The sensing domains in these channels consist of a positively charged C-terminal cluster, which interacts with the membrane surface. This interaction can be modified by introducing either polylysine (a cationic molecule used to mask the charge of anionic lipids) or changing the fraction of phosphatidylinositol bisphosphate (PIP$_2$) in the membrane. PIP$_2$ has a valence of $-4$ at pH 7.0 and changing the fraction of PIP$_2$ will have a large effect on the surface potential of the membrane (24).

Electrostatic interactions between protein and membrane surfaces are also of major importance for the regulation of the natively unstructured protein myristoylated alanine-rich C-kinase substrate (MARCKS) (25). In this protein, a basic and hydrophobic region has been identified to be responsible for concentrating the signaling lipid PIP$_2$, which constitutes about

FIGURE 7. The effect of pH on the activity and osmotic activation of parental OpuA, OpuAΔ12, and OpuAΔ12insHis. Proteoliposomes were loaded with 9 mM MgSO$_4$, 9 mM Na$_2$ATP plus 50 mM potassium P, at the corresponding pH, which is equiosmolar with 70 mM potassium P, of the outside medium. Uptake of $[^{14}$C]glycine betaine by OpuA (●), OpuAΔ12 (○), and OpuAΔ12insHis (▲) in 70 mM potassium P, and supplemented with 0 mM KCl (A), 100 mM KCl (B), and 200 mM KCl (C) is shown. Proteoliposomes were composed of 50 mol % DOPE, 12 mol % DOPC, and 38 mol % DOPG.

FIGURE 8. Schematic representation of the activity of OpuA at low (12 mol %) and high (38 mol %) of DOPG. The transporter (ligand binding receptor, translocator including ATP binding cassette) is depicted by the main cylinder, the CBS module in orange, the surface-exposed residues in blue (cationic) or yellow (neutral substitutions), and the anionic C terminus in red; the curled tail indicates the deletion of the tandem CBS domain. Anionic lipids are depicted by red circles and neutral lipids are in gray. Top, wild type OpuA; middle, OpuAΔ119 (= ΔCBS); bottom, OpuA[K3R2].
1% of the total lipids in the plasma membrane. The hydrophobic residues penetrate into the lipid bilayer and the basic residues attract three PIP$_2$ molecules by means of nonspecific electrostatic interactions. Calmodulin, complexed with Ca$^{2+}$, has the ability to wrap around the basic cluster of MARCKS, reversing its charge from positive to negative and repelling the complex from the membrane. The interaction of MARCKS with the membrane regulates PIP$_2$ sequestration and connects the local intracellular Ca$^{2+}$ concentration to PIP$_2$, allowing interaction of these signaling molecules with other biologically important molecules.

Next to PIP$_2$, the bioactive lipids phosphatidic acid (PA) and lysophosphatidic acid (LPA) are important signaling lipids in the membrane. Although their abundance in the membrane is low, they are involved in many intracellular processes and are important intermediates in lipid biosynthesis (26). Recently, it has been shown that the phosphomonoester headgroups of LPA can form inter- and intramolecular hydrogen bonds, thereby increasing the protonation state of LPA (27). These results suggest that hydrogen bonds between LPA and basic residues in proteins play an important role in protein–lipid interactions. An example is the CTP:phosphocholine cytidyltransferase, the key regulatory enzyme in the synthesis of phosphatidylcholine. The activity of cytidyltransferase is dependent on the content of anionic lipids in the membrane, including PA and phosphatidylglycerol (28). Cytidyltransferase contains an amphipathic helix that has been identified as responsible for the physical properties of the membrane (29). A similar amphipathic effect has been observed on the myristoylated alanine–rich C-kinase MARCKS (25). Cytidyltransferase and MARCKS are both amphipathic proteins that bind to membranes in a two-step process. One step involves electrostatic interactions and serves to localize the protein to the surface, the other step constitutes the hydrophobic interaction. For the CBS module of OpuA, it is not clear whether hydrophobic interactions occur between the protein domains and the membrane. Knowing that CBS tandem domains have a distinct hydrophobic core and a more hydrophilic exterior, as indicated by the various crystal structures of CBS homologues (8, 11, 30, 31), hydrophobic interactions with the membrane seem less important than electrostatic interactions.

In conclusion, the surface-exposed cationic residues of the CBS module play an important role in osmotic (i.e. ionic) regulation of OpuA. We propose that the CBS module may interact (directly or indirectly) with the anionic membrane surface and lock the transporter in an inactive state at low ionic strength. Above threshold values of ionic strength, the electrostatic interactions are screened and OpuA is activated. The strength of the electrostatic interaction is tuned by the C-terminal tail, which does not require a specific structure but simply depends on the number of co- and counterions in the protein and on the membrane. By introducing histidines in the C-terminal, the transporter became responsive to the internal pH.

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