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Structure of the sucrose-specific porin ScrY from *Salmonella typhimurium* and its complex with sucrose

Doris Forst¹, Wolfram Welte², Thomas Wacker¹ and Kay Diederichs²

The X-ray structure of a sucrose-specific porin (ScrY) from *Salmonella typhimurium* has been determined by multiple isomorphous replacement at 2.4 Å resolution both in its uncomplexed form and with bound sucrose. ScrY is a noncrystallographic trimer of identical subunits, each with 413 structurally well-defined amino acids. A monomer is built up of 18 anti-parallel β-strands surrounding a hydrophilic pore, with a topology closely similar to that of maltoporin. Two non-overlapping sucrose-binding sites were identified in difference Fourier maps. The higher permeability for sucrose of ScrY as compared to maltoporin is mainly accounted for by differences in their pore-lining residues.

Gram-negative bacteria protect their vulnerable cytoplasmic membrane by a peptidoglycan layer and an outer membrane (OM)1.2. It has been estimated that ~10% of all genes in E. coli are involved in transport of small nutrient molecules across the OM, the periplasmic space with the peptidoglycan and the cytoplasmic membrane (CM)3. Sugar-specific, active transport systems in the CM with K_m values in the micromolar range maintain a low concentration of monosaccharides, disaccharides and unbranched oligosaccharides in the periplasm^{4,5}. In contrast, sugar transport across the OM is purely passive, driven by diffusion. According to Fick's equation the flux of a species of molecules accross the OM is proportional to the concentration difference between the external space and the periplasmic space. The high value of the proportionality constant, the permeability, is due to the porins, the major protein component of the OM. So-called 'general diffusion porins'6,7 or 'nonspecific porins' form water-filled channels through which ions and polar molecules, smaller than the exclusion limit of the channel, may pass. Nonspecific porins are between 30,000-50,000 M_r and form trimers. The ionic permeability of general diffusion porins can be measured as current flowing through porins reconstituted into black lipid membranes6 or through small OM fragments in patch-clamp experiments8. Plots of current against ion concentration are linear, showing that ions pass through the OM in a diffusion-like process. The linear relation of current and ion concentration and the differences in the permeability of anions and cations can be rationalized qualitatively by theoretical models9.

The structures of five general diffusion porins have been determined by X-ray crystallography¹⁰⁻¹³. These structures are variations of a 16-stranded antiparallel β -barrel with nearest-neighbor connections between strands. The connections form loops (L) of varying length on the external side and mostly hair-pin turns (T) on the periplasmic side of the OM. The third external loop (L3) folds along the inner barrel wall and constricts its lumen in the center of the membrane, giving the channel an hourglass-like cross section that allows for a high, but charge-dependent permeability¹⁴.

The permeation of solute molecules through the channel can be treated with reaction-rate theories, either transition state theory¹⁵ or Kramers theory¹⁶. The latter appears to be more appropriate to aqueous protein–substrate systems. The rate coefficient for permeation contains a Boltzmann factor exp[- Δ G'/RT] with the activation free-energy Δ G^{*} = Δ H^{*}-T Δ S^{*} of the entry process (formula (14) in ref. 16). The entropy term -T Δ S^{*} will represent a considerable fraction of the free-energy barrier, as the permeating molecules are confined to the narrow cross section of the channel.

Under growth-limitation conditions the cells must strive to increase the permeability further to yield a sufficient flux of nutrient molecules at shallow concentration gradients. Increasing the density of nonspecific porins in the OM and increasing the cross section of the porin channel are inappropriate means because the OM is already densely packed with porins¹⁷ and because an increased exclusion limit would weaken the protective effect of the OM. Another means to increase the permeablity is to decrease the free energy barrier for entrance into the channel so that the rate of entrance is increased. This can be effected by introducing binding sites specific for certain nutrient molecules, such as sugars or nucleosides, since the negative contribution of the enthalpy of one or several binding sites will decrease ΔG^* . Binding sites arranged along the inner channel wall thus will increase the permeability compared to the case of a nonspecific channel, as long as the concentrations of the substrate molecules are well below their dissociation constants (K_d) .

Indeed, porins specific for different small nutrient molecules are expressed by Gram-negative bacteria under conditions of growth limitation. One example is maltoporin (also called LamB due to its role as a receptor for phage λ) which is specific for maltose and maltooligosaccharides. It is coded on the *mal* regulon together with a maltose transport complex of the ATP binding cassette (ABC) type and a periplasmic binding protein¹⁸. Maltoporin is expressed at disaccharide concentrations lower than ~100 µM¹⁹, while the K_ds of the binding sites for disaccharides of both glycoporins are in the 10 mM range. The K_ds drop for oligosaccharides with 3, 4 and 5 glucosyl units but remain constant at a value of ~300 µM²⁰ for longer ones.

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Institut für Biophysik und Strahlenbiologie, Albert-Ludwigs-Universität, Albertstr. 23, 79104 Freiburg im Breisgau, Germany. ²Fakultät für Biologie, Universität Konstanz, Box M656, 78457 Konstanz, Germany.

Correspondence should be addressed to W.W. email: Wolfram. Welte@uni-konstanz.de



The structure of maltoporin with and without bound maltooligosaccharides and also with sucrose has been determined by X-ray crystallography^{21–24}. Maltoporin possesses the same overall structure as the general diffusion porins, except that its barrel contains 18 strands. As expected from binding experiments²⁰, maltoporin possesses a chain of 3–5 low affinity binding sites for glucosyl groups extending along the wall of the channel.

Schmidt *et al.*²⁵ have discovered a sucrose uptake system on plasmid pUR400 in *Salmonella typhimurium* and *Escherichia coli* that confers to the cells, under growth limitation conditions, the ability to grow on sucrose as a sole carbon source. The *scr* regulon codes for a sucrose-specific phosphoenoltransferase-dependent transport (PTS) complex in the cytoplasmic

Fig. 1 Ribbon superposition of ScrY (yellow) and maltoporin (blue) with the periplasmic side and the external side at the bottom and top, respectively. The barrel consists of 18 antiparallel strands and has a kidneyshaped crossection of between 30 Å and 50 Å diameter. The height varies between 20 Å near the trimer axis and 50 Å at the trimer periphery. The tilt angles of the β -strands relative to the trimer axis is between 35° and 55°, and the barrel shear number⁷³ amounts to +22. This figure was prepared with RIBBONS⁷⁴.

membrane. It was later found that one gene, *scrY*, codes for a porin that is specific for sucrose and maltooligosaccharides and that the *scr* regulon is also found in the genome of *Klebsiella pneumoniae*^{26,27}.

N-terminal sequencing²⁸ showed that mature ScrY and maltoporin possess 483 and 421 residues respectively. Upon comparing sequences it was found that a C-terminal ScrY domain of 411 residues can be aligned with the Maltoporin sequence with an overall identity of around 20%^{26,27}. The N-terminal 72 residues of ScrY have no equivalent in maltoporin.

We have crystallized $ScrY^{29}$ and solved by multiple isomorphous replacement (m.i.r.) the structure of the pore domain (residues 71–483) at 2.4 Å resolution both with and without bound sucrose.

Crystallographic problems and solutions

Three presumably interrelated problems rendered the structure determination of ScrY difficult. First, the space group reported for the low-resolution data²⁹ of the preliminary crystallographic investigation proved irreproducible. At resolution of 4 Å and beyond, additional reflections $(-h + k + 1 \neq 3n)$ were found between the reflections of the obverse-indexed hexagonal-rhombohedric lattice (-h + k + l = 3n), turning the lattice into a hexagonal-primitive one. These reflections were of negligible intensity at low resolution (6 Å), but were of the same average intensity as those of the R3 lattice at resolution better than 3 Å, overall increasing the proportion of weak reflections. This finding can be explained by a slight distortion of the R3 space group, leading to the P3(1,2) subgroup with a trimer in the asymmetric unit, whose non-crystallographic symmetry axis is close to, but does not coincide with, the rotation axis of the R3 system. The three-fold rotational symmetry of the



Fig. 2 View of the membrane spanning surfaces of the glycoporin trimers. a, ScrY; b, maltoporin. Aromatic residues are added to the ribbon representation of both porins. The two girdles of aromatic residues that bound the hydrophobic zone of 30 Å width are discernible. In ScrY they are less regular and some aromatic residues are within the hydrophobic zone. Moreover, maltoporin differs from ScrY by an additional clustering of aromatic residues in the external loops. This figure was prepared with Molscript⁷⁵ and Raster3D⁷⁶.



hexagonal primitive lattice was well fulfilled, and the absence of intensites of the (0,0,l) reflections with $l \neq 3n$ confirmed the space group P3_(1,2).

Second, we found the ScrY crystals to be highly anisomorphous, with crystals even from the same crystallization setup exhibiting variations of unit cell axes in the order of 2-3%, leading to difficulties in interpretation of difference Patterson and difference Fourier syntheses. *R*-factors between native crystals ranged mostly between 20-30%. Therefore, complete data sets had to be collected on single crystals.

Third, ScrY crystals were twinned, with twinning fraction α evenly distributed in the range 0–0.5. Procedures as described in the Methods section were used to determine the twinning fraction according to equation 1 and for detwinning the data according to equation 2.

Despite these difficulties initial single isomorphous replacement phases were obtained by solving the difference Patterson map of the cocrystallized KAu(CN)₂-derivative. Interpretation of other derivatives followed (Table 1). It was found that detwinning provided cleaner difference Patterson functions and was necessary for their interpretation. However, for heavyatom refinement and phasing the observed (and not the detwinned) amplitudes were used, as detwinning produces datasets of lower completeness, and we were able to identify derivative datasets with low twinning fractions. M.i.r phasing allowed us to obtain an interpretable density map at 2.9 Å resolution, into which residues 71–483 of the mature sequence could be built in several macrocycles of X-PLOR refinement and manual improvements of the model, using O.

After completing the final X-PLOR refinements, SHELXL refinements of the native and sucrose-complexed models confirmed the twinning fraction as determined by equation 1 and showed that inclusion of α as a refinable parameter (as opposed to no twinning treatment) decreased both R_{free} and R_{cryst} by only 0.12% and 0.20% respectively. However, results of these refinements were discarded since — in addition to requiring almost prohibitive computing resources — it was not possible to restrain the positions of NCS-related water molecules in a simple way. Therefore, final statistics (Table 2) except for twinning fractions are based on X-PLOR refinement.

Fig. 3 Structural alignment between ScrY and E. coli maltoporin. Capital letters denote structurally equivalent residues based on a 3.8 Å cutoff, whereas residues without a structural counterpart are shown in lower case letters. In addition to the 18 strands of the main β -barrel, short pieces of regular secondary structure (E β strand; G, 3₁₀-helix) as classified by DSSP⁷⁷ are also given.

Description of the structure

The model of the C-terminal 413 residues of ScrY represents the complete transmembrane domain. As in all porins whose structures have been analyzed so far, ScrY is basically a β -barrel with nearest neighbor connections between strands and with large loops (L) of irregular length exposed to the external surface. L3 is attached by polar and apolar interactions to the inner barrel wall. It constricts the channel cross section near the center to 8 × 11 Å so that the channel lumen possesses an hour-glass ross section with a large external and a par-

shaped longitudinal cross section with a large external and a narrower periplasmic vestibule.

Like other porins, ScrY forms stable trimers in the outer membrane. In contrast to the structures of the nonspecific porins, which all consist of 16 β -strands, the two glycoporins ScrY and maltoporin²¹ possess 18 strands. The observed barrel structure has a kidney-shaped cross section with dimensions of roughly 50 Å × 28 Å and a height of between 25 Å near the trimer center and 50 Å at the trimer periphery.

When the MIR structure analysis of ScrY was begun, the low sequence similarity with maltoporin led us to expect considerable structural differences between the two porins. However, the two independently solved structures show a surprisingly large degree of structural conservation (Fig. 1). The gross differences between both glycoporins are in the external loops which tend to tilt over the pore vestibule in both porins like petals and thereby form a constriction at the entry to the common vestibule of all three monomers. The residues of the phage- λ binding site of maltoporin on L9, L4, L6 are not conserved in ScrY. L9 contains a short 310-helix and thereafter a Ca2+-binding loop segment. In the latter, the ion is bound in a distorted tetragonal-bipyramidal geometry. The longest loop L1, which consists of 29 amino acids and contains a short 310-helix, folds down towards the constriction site and is involved in hydrogen bonds with L3. These hydrogen bonds stabilize the structure of both loops. Generally, ScrY is less 'closed' than maltoporin as L4, L6, L8 and L9 are shorter. The similarity of the backbones of both glycoporins is quite high in view of the modest sequence homology: using a 3.8 A cutoff, 315 of the 413 Ca atoms of ScrY can be superimposed with those of maltoporin with a root-mean-square (r.m.s.) deviation of only 1.3 Å.

Among the periplasmic turns (T), T5 is larger in ScrY and protrudes in a radial direction from the barrel, whereas T4 is markedly shorter in ScrY. T5 is involved in the crystal-contacts and T8 folds into the channel mouth.

The monomer-monomer interface

Trimer formation is stabilized through many polar interactions between monomers at the top and bottom of the β -barrel. In between, there exists an apolar region formed by small



hydrophobic amino acids. The low average B-factor of around 18 Å² indicates the rigidity of this region. On the periplasmic side of the trimer center, near the N and C termini, a cluster of six phenylalanines, formed by residues 73 and 75 of each monomer, is found. Here, the aromatic rings from adjacent monomer interfaces are mutually perpendicular to each other, optimizing quadrupole interactions³⁰ and connecting adjacent monomers.

In ScrY and all *E. coli* porins, loop L2 fills a gap above L3 of a neighboring monomer^{11,21}. This loop contributes considerably to the stability of the trimer because it connects adjacent porins in a manner similar to an outward-bent hook.

The membrane exposed surface

The peripheral transmembrane surface of the ScrY porin trimer can be divided into three zones that encircle the trimer: a hydrophobic zone with a width of up to 33 Å composed of aliphatic residues (such as Ala, Leu, Ile and Val) and two hydrophilic zones at the periplasmic and external surfaces. Two aromatic girdles form borderlines between the hydrophobic zone and the two hydrophilic zones (Fig. 2a). A neutron diffraction analysis³¹ showed that the hydrocarbon core of detergent micelles just covers the hydrophobic zone of OmpF porin, which is thus identified as the contact interface with the lipid chains in the outer membrane. The accumulation of aromatic amino acids at the polar/apolar border is a common feature of all known structures of porins and is also seen, but is less obvious in other membrane proteins³². These girdles are thought to seal the porins into the outer membrane lipid bilayer as the 'amphiphilic' aromatic residues are well suited for partitioning into the interphase between the hydrocarbon and polar membrane layers^{10,12}. The comparison with maltoporin (Fig. 2b) and with nonspecific porins shows that the girdles in ScrY are much less regular, with more aromatic residues in the hydrophobic zone. We speculate that the irregularity of the girdles in ScrY may contribute to the specific toxicity of ScrY to E. coli cells at high expression levels26 since this would likely impair the sealing of the porins with the membrane lipid bilayer, possibly resulting in membrane instability or porin accumulation somewhere before the outer membrane.

The 'greasy slide' and its 'ionizable track'

Schirmer *et al.*²¹ identified six aromatic residues on the inner surface wall of the maltoporin from the external vestibule across the constriction site toward the periplasmic vestibule. As sugar binding by proteins often involves stacking of pyranosyl rings on aromatic rings³³ these authors proposed that these aromatics are a

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Fig. 4 Images⁷⁸ comparing the hourglassshaped inner channel surfaces of *a*, ScrY complexed with two sucrose molecules and *b*, maltoporin. The cutting plane is spanned by the trimer axis and a radial beam connecting the axis and the constriction site center. The molecular surface near Asp, Glu, Arg, Lys is colored red whereas near Phe, Tyr, Trp it is colored green.

chain of binding sites or a binding slide for maltose and maltoligosaccharides. In structures of maltoporin complexed with maltooligosaccharides, most of the glucosyl groups are in van der Waals contact with some of the aromatic residues.

In ScrY, the four external of the six aromatic residues found in maltoporin are

absolutely conserved (Trp 151, Tyr 118, Tyr 78, Trp 482; Fig. 3), even up to the level of a similar side chain conformation. The fifth, Trp 358, is conservatively replaced by Phe 435 whereas the sixth near the periplasmic mouth is missing. Following Wang *et* $al.^{23}$, we use the terms S1–S5 for the glucosyl binding sites near Trp 151, Tyr 118, Tyr 78, Trp 482 and Phe 435.

The structure of the residues lining the constriction site (78-80, 110, 114, 118, 120-121, 140, 142, 161, 187-188, 194-201, 204, 207, 322-323) is rather similar in both glycoporins. Of these residues, all except three are conserved or conservatively replaced and most adopt a similar side chain conformation in ScrY and maltoporin. In spite of the very similar architecture of both glycoporins around the constriction site, the cross section of the channel in ScrY is larger (8.5 Å \times 11 Å) than in maltoporin (7 Å \times 10 Å). This is due to three major exchanges of L3-residues protruding into the lumen: Asn 192 in ScrY becomes Arg 109 in maltoporin, but with a reversed orientation so that the side chain is oriented into the lumen in maltoporin; Asp 201 and Phe 204 in ScrY are replaced by Tyr 118 and Asp 121 of maltoporin, the latter two residues protruding more into the constriction site lumen than the former. The larger cross section of the constriction site and the wider external channel mouth qualitatively explain the much higher single channel conductance of ScrY in 1 M KCl (1,4 nS) as compared to maltoporin (0,15 nS)28.

In the neighborhood of the greasy slide, the degree of conservation of ionizable residues — the 'ionic track' of Schirmer²³ — is high and above the average conservation found between ScrY and maltoporin. Many of these are conserved even in their side chain conformation between ScrY and maltoporin. Some of them were found to be involved in hydrogen bonding of two maltose molecules, a maltotriose molecule, a maltohexaose molecule and a sucrose molecule bound to maltoporin^{22,23}.

Sucrose binding sites in ScrY

The structure of ScrY cocrystallized with sucrose reveals two sucrose molecules, 'Suc1' and 'Suc2', bound to the channel surface (Fig. 4*a*) with occupancies of 0.7 and 0.8 respectively. Their glucosyl groups are in van der Waals contact with Tyr 78 (S3), located in the constriction site, and with Trp 482 (S4), adjacent to the periplasmic end of the constriction site (Figs 5, 6*b*), the same sites as for the glucosyl groups (g3 and g4) of a maltose found in maltoporin²². The sucrose molecules form hydrogen bonds with the ionizable track in the neighborhood of the greasy slide and of L3, as shown schematically in Fig. 7.

Suc1 and Suc2, as well as a single sucrose molecule in malto-

Fig. 5 Constriction site viewed (stereo) from the periplasmic space with superimposed electron density map. Suc2 is seen in the front bound near Trp 482 (S4 binding site), Suc1 is seen in the background near Tyr 78 (S3 binding site). Green: $2F_{obs} - F_{calc}$ density at 2.0 σ . Cyan: $F_{obs} - F_{calc}$ density at 2.0 σ . Cyan: $F_{obs} - F_{calc}$ density at 1.5 σ of the ScrY-sucrose complex with phases calculated after refinement of the model without sucrose.

porin²³, are all similarly positioned with the glycosyl group nestled against the greasy slide, the fructosyl group bending away from it and the nonreducing glucosyl-C4 pointing towards the periplasm. The greasy slide seems to bind the sugars in a defined direction, that is with the nonreducing end towards the periplasm.

This orientation has been found for all bound sugars in glycoporins²²⁻²⁴.

Sucrose binding to S3 is sterically blocked in maltoporin due to the presence of Tyr 118 and Arg 109, both of which protrude into the lumen of the constriction site (Fig. 6a). These maltoporin residues as well as Asp 121 are represented in Fig. 6a by their transparent van der Waals surface. In addition to rendering the binding of sucrose to S3 sterically impossible, these residues of maltoporin represent a 'bolt' in the constriction site blocking the path between S3 and S4. The remainder of this region is very similar in both glycoporins. Indeed, the glucosyl group of the single sucrose molecule found by Wang *et al.*²³ in maltoporin binds to a position between S2 and S3, just before the constriction site.

From the very low permeation rate of sucrose through maltoporin, one can infer that an extended conformation of sucrose, which should permeate through maltoporin efficiently, is very unfavorable. Our results demonstrate how ScrY, in contrast to maltoporin, is constructed to allow the bulky sucrose to slide easily through the channel. The observation that, in the absence of ScrY, *E. coli* cannot grow on low sucrose concentrations as sole carbon source²⁶ shows that ScrY achieves a high permeability for this disaccharide.

Other protein-sucrose complexes

In contrast to maltose and maltoligosaccharides, sucrose in crystalline form and bound to proteins has a globular conformation with the glucosyl and the fructosyl moieties bent towards each other. The torsion angles, Φ , Ψ , of sucrose³⁴ are found to be similar in crystallographical and molecular modelling studies to the values of crystalline sucrose (108°/-45°), indicating a similar overall conformation in solution and in the crystal^{34–36}. In ScrY, the torsion angles about the two glycosidic bonds of sucrose, Φ , Ψ , are found to be 75°/-31° for Suc1 and 90°/-61° for Suc2.

In a complex with lentil-lectin³⁷, sucrose exhibits torsion angles of 107° and -58°, with a water intercalated between glucosyl-O2 and fructosyl-O3. All of the glucosyl hydroxyls are hydrogen bonded to the protein but only two fructose hydroxyls form indirect hydrogen bonds to the protein through waters. Moreover, the glucosyl group is stacked on a phenylalanine ring. The features of the bound sucrose in lentil-lectin are also found in ScrY and maltoporin (Fig. 7). The two glucosyl groups are more intimately attached to the channel wall than the fructosyl groups, that is they are in van der Waals contact with aromatic residues and form a larger number of hydrogen bonds to the protein. As the fructosyl group contributes less to the free energy of

binding and since maltose can be considered as two glucosyl groups the binding of maltose to ScrY should be stronger than that of sucrose to ScrY. Indeed, experimental determination of the respective binding constants²⁸ yield K_D values of 6.25 mM and 50 mM for maltose and sucrose.

Oligosaccharide binding sites involving chains of aromatic residues flanked by potential hydrogen bond forming residues have also been found in other proteins. Cellulose-binding domains of cellulolytic enzyme complexes from fungi and bacteria indicated binding sites for three adjacent crystalline cellulose chains with aromatic residues interacting with roughly every second glucose unit of one chain³⁸. *E. coli* maltodextrin phosphorylase shows stacking of a glucosyl group on a Tyr³⁹. The authors discuss two factors contributing to the favorable stacking of glucosyl groups on aromatic residues: hydrophobic interactions and electrostatic interactions between partial positive charges on the hydrogens bound to carbons of the sugar ring and the π electrons of the aromatic ring.

On the other hand, chains of aromatic residues are not a general feature of oligosaccharide binding, as seen from the structure of a bacterial endoglucanase⁴⁰, and lysozyme⁴¹. C-type (Ca²⁺-dependent) lectins show how the surroundings of a Ca²⁺ atom, bound to the surface of a protein, can create a low-affinity binding site for monosaccharides (K_D between 6 and 22 mM) which can bind two neighboring equatorial sugar hydroxyls to the protein^{42,43}. Ca²⁺ has also been found to be involved in indirect binding of glucose through water molecules⁴⁴ or by apparently optimizing the conformation of residues for hydrogen bonding to sugar hydroxyls⁴⁵.

Glycoporins possess a series of low-affinity binding sites for glucosyl groups. Binding occurs through hydrophobic contacts with aromatic residues and hydrogen bonding to ionizable residues. The binding sites are arranged as an extended slide from the extracellular mouth to the periplasmic mouth of the channel. The observed increase of binding constants for maltooligosaccharides with increasing number of glucosyl residues in maltoporin²⁰ and ScrY²⁸ suggests a series of four or five binding sites with the correct spacing and geometry to bind such helical maltooligosaccharide chains^{21,24}. As binding sites increase the sugar permeability of the outer membrane, it is expected that glycoporins confer a growth advantage to bacteria whose growth is limited by low concentration in the culture medium of disaccharides like maltose or sucrose or maltotriose and may be indispensable for the uptake of maltooligosaccharides with more than three glucosyl residues. Indeed in the presence of maltooligosaccharides longer than three glucose units as the only available car-



bon source, *E. coli* cells need maltoporin or ScrY for growth²⁶. It is comprehensible to ascribe this requirement to the greasy slide, as permeation of these longer chain molecules with their high entropy of threading through the channel will depend upon a high-affinity binding site. As Schirmer *et al.* have pointed out, the complementarity of the greasy slide geometry to the natural left-handed superhelical conformation of maltooligosaccharides is one feature underlying the high affinity²¹.

The binding sites with the highest affinities are found close to the constriction site. This is in accord with the high entropy barrier associated with threading the sugar molecule through this zone. Binding at any position on the greasy slide may be 'flexible'; that is, sugars might find alternative hydrogen bonding partners at similar free energies. Mutational studies are in accord with this view, as point mutations of aromatic residues of the 'greasy slide' in maltoporin do not inhibit uptake of maltooligosaccharides^{46,47}. The resulting free-energy profile of a glucosyl unit sliding along the chain of binding sites seems to be smooth, with only shallow barriers in between sites⁴⁸.

Interestingly, differences in side chain conformations between residues of the native and sucrose-complexed form are only slightly greater than those of residues not involved in sucrose binding. The side chain conformations of sucrose-binding residues in the native **Fig. 6 a**, Constriction site viewed (stereo) from the periplasmic space with Suc1 bound to S3. The residues forming the constriction site are shown. All of these, except for Asn 192, Asp 201 and Phe 204, are conserved in maltoporin up to a similar side chain conformation. The space occupied by the three altered residues (Arg 109, Tyr 118, Asp 121 in maltoporin) is indicated by its transparent van der Waals surface. **b**, Greasy slide, stereo side view with the external side oriented to the top, as represented by aromatic residues Trp 51, Tyr 118, Tyr 78, Trp 482, Phe 435 (S1–S5) with Suc1 bound near Tyr 78 (S3) and Suc2 bound near Trp 482 (S4). This figure was prepared with GRASP⁷⁸.

and complexed structures can be overlayed with an r.m.s.d. of 0.24 Å, which is the same value as found when comparing all side chains. This rigidity of the residues lining the sucrose path should minimize frictional forces on the sliding sugar molecule by the protein. According to the reaction rate theory of Kramers, a low frictional coefficient in an overdamped protein system ensures a high reaction rate¹⁶ that results in a high channel permeability.

Division of permeation into four reactions

A full description of permeation should include the following consecutive reactions :

(i) Diffusion of sugars from the external medium to the vestibule.

(ii) Collisions with and diffusion on the vestibule surface resulting in binding to the external end of the greasy slide.

(iii) One-dimensional diffusion along the greasy slide.

(iv) Escape of sugar from the periplasmic end of the greasy slide.

At equilibrium the flux of sugar through each partial reaction is equal and expected to result from evolutionary optimization for high permeability. The apparently low frictional coefficient of the sliding reaction (iii) could be a consequence of such an optimization.

The larger the concentration gradient driving the first reaction, the better is the process optimized. It seems not to be perfect yet as Jordy et al.49 have concluded from the order of magnitude of the on-rate constants for maltooligosaccharide binding that the sugar transport is not limited by the diffusion to the channel entrance; in other words, that the sugar hits the channel many times before it is bound to the greasy slide in the constriction site. Nevertheless the external vestibule should be engineered to minimize the number of hits of sugar molecules needed for a successful capturing. If maltoporin and ScrY have been optimized for capturing different sugars, their respective vestibule surfaces should differ in their respective coating with residues. Maltooligosaccharides that represent an extended ribbon-like structure with one face being less polar might for instance stick to patches of aromatic residues. Sucrose molecules with their globular shape and their numerous hydrogen bonding groups might preferentially stick to ionizable residues. A weak association of the sugar with the vestibular surface could



Fig. 7 LIGPLOT⁷⁹-style scheme of hydrogen bonding network of Suc1 bound to S3 and Suc2 bound to S4.

improve the diffusional flow of sugars through reaction (ii)⁵⁰. In Fig. 4 both glycoporins are shown with aromatic residues at the surface coded in green and ionizable residues (Asp, Glu, Arg, Lys) coded in red. Indeed, maltoporin has a noticeable density of aromatic residues on its vestibule surface (see also Fig. 2) while in ScrY ionizable residues predominate.

The periplasmic side of ScrY

Schülein *et al.*²⁸ constructed a deletion mutant ScrY Δ 3–73 and studied its properties. The deletion increased the cation *versus* anion permeability P_c/P_a and increased the association with the peptidoglycan.

Due to absence of electron density, the N-terminal 70 residues of the ScrY crystal structure could not be traced and modeled. Inspection of the sequence shows a segment with a clear heptad 'a'-'g' pattern (A. Lupas, personal communication) indicating a 99% probability for a coiled-coil tertiary structure⁵¹ for 42 residues starting at Ile 4. The heptads are indicated in the diagram in Fig. 8. Hydrophobic residues Leu, Ile, Ala, Val are on positions 'a' and 'd', while Glu residues are on position 'e' and Arg and one Lys on position 'g'. No structural prediction exists for the segment between Gln 46 and Lys 70.



It appears likely that three α -helices, one from each ScrY monomer, associate to form a left-handed coiled-coil. The latter would consist of 12 α -helical turns and have a length of roughly 65 Å. Assuming an interhelical distance of 11 Å (ref. 52) and a diameter of 12 Å, the cross section of the coiled-coil would be roughly 23 Å. The packing of the protein in the unit cell allows accomodation of such a cylindrical volume without steric problems.

The length of the N-terminal extension is much shorter than the 400 Å predicted for the coiled coil of the periplasmic lipoprotein Omp α from *Thermotoga maritima*⁵³, shorter than the estimated distance of 200 Å (ref. 53) between the outer and inner membrane of *E. coli*, shorter than the coiled-coil predicted within the II-domain of TolA which was reported to bind porins⁵⁴ and of roughly the same length as the coiled-coil predicted for the murein lipoprotein (Braun's lipoprotein) of *E. coli*^{55,56}.

ScrY is coexpressed with a sucrose specific PTS system for which no periplasmic binding protein is known. Free diffusion of the sugars in the periplasmic space would reduce the efficiency of sugar uptake as some sugar molecules would diffuse back to the external phase. We speculate that the N-terminal coiledcoil functions as a low affinity binding site for sugars, effectively replacing a sucrose-specific periplasmic binding protein. The oppositely ionizable residues on the 'e' and 'g' heptad positions of the coiled-coil are potential hydrogen bonding partners for sugar hydroxyls. Oppositely ionizable residues on 'e' and 'g' positions have been shown to contribute to the stability of other coiled-coils57 in accord with crystal structures where these charged pairs form salt bridges58. However, theoretical considerations⁵⁹ and pK_a shift values of acidic residues in coiled-coils⁶⁰ suggest that these ionizable residues remain solvated and thus are potential hydrogen bonding partners for sugar hydroxyls in vivo. The question if they really form low-affinity sugar binding sites must be answered experimentally.

Methods

Crystallization and data collection. The isolation and purification of ScrY is as previously described^{29,61}. ScrY was crystallized by vapor diffusion using the sitting drop method at 17 °C. The drop contained 5–7 mg ml⁻¹ protein, 20 mM Tris/Cl at pH 7.7, 100 mM LiCl, 20 mM MgSO₄, 1.2% β-D-octylglucopyranoside (β-OG), at least 1% Hexyldimethylamineoxide (C₆DAO), 1% β-D-heptylglucopyranoside (β-HG) and 6–9% PEG-2000. The concentration of PEG in the reservoir was 12–15 %. For cocrystallization with sucrose, we added 100 mM sucrose to the drop. Heavy atom derivatives were produced by soaking crystals or by cocrystallizing the protein in the presence of the heavy metal compound. Crystals grew within three to six weeks to a size up to 400 µm in each direction. The space group is P3₂ ($a = b = 112.1 \pm 1.5$ Å, $c = 147.0 \pm 2$ Å) with one trimer per asymmetric unit. The Matthews-parameter of the crystal has a value of 3.8 Å³/Da, indicating a solvent (plus detergent) content of 68%.

The crystals diffracted to better than 2.0 Å resolution using synchrotron radiation (EMBL beamline X11 at DESY, Hamburg, Germany). The native data for the initial structure solution reported here as well as the heavy atom derivatives and sucrose data were collected on an image-plate-detector system (STOE, Darmstadt, Germany) using CuKα-radiation from a rotating anode generator (40 kV, 100 mA). Crystals were aligned with the c axis along the spindle,

and frames of 0.5° oscillation were collected over 65° at 14 °C. The resulting images were processed with XDS⁶².

Crystallographic procedures. Heavy atom binding sites were located in difference Patterson maps and refined using DAREF⁶³. The multiple isomorphous replacement (MIR) phases were improved by solvent flattening

Fig. 8 Sequence of the N-terminal 70 residues of ScrY showing heptad pattern of residues 4-45.

| Table 1 Crystallographic data | | | | | | | | | |
|---|--------------------------|----------------|---------------------|-----------------------|--------------------|--------------------------------------|--------------------------------------|---|-----|
| Dataset | Twinning fraction (%) | Resolution (Å) | Completeness (%) | Number of reflections | Number of sites | R _{sym} ¹ (%) | R _{iso} ² (%) | Phasing power ³ in shells 99–4 Å 4–2.9 Å | |
| Native 0 | 0 | 2.89 | 83.5 | 38,033 | L R | 10.4 | - | - | - |
| Native 1 | 3 | 2.89 | 93.5 | 42,612 | - | 9.7 | 29.9 | - | - |
| Native 2 | 3 | 2.29 | 92.6 | 82,849 | - | 8.3 | 24.9 | - | - |
| Sucrose soak | 3 | 2.29 | 78.7 | 72,892 | - | 16.1 | 30.1 | - | |
| KAu(CN) ₂ | 3 | 2.89 | 83.7 | 38,145 | 9 | 9.7 | 28.0 | 2.2 | 1.3 |
| cis-Pt(NH ₃) ₂ C | l ₂ 3 | 2.89 | 95.7 | 43,612 | 9 | 11.3 | 29.1 | 1.2 | 0.7 |
| K ₂ ReCl ₄ | 12 | 2.89 | 85.4 | 38,912 | 9 | 10.6 | 23.4 | 1.1 | 0.6 |

 ${}^{1}R_{sym} = \sum_{hkl} \sum_{i} |I_{hkl} \langle I_{hkl} \rangle | / \sum_{hkl} \sum_{i} I_{hkl}; \langle I_{hkl} \rangle = \text{average of } I_{hkl}.$

 ${}^{2}R_{iso} = \sum |I - I_{p}| / \sum |I_{p}|$; $|I_{p}|$ is taken from dataset Native 0.

Phasing power is mean value of heavy atom structure factor amplitude divided by lack of closure.

R

P

T

R

N

C

N

R

h

s

⁴Reference data to determine the twinning fraction α and R_{iso} was data set Native 0.

and non-crystallographical symmetry (NCS) averaging, using RAVE64 or DM65,66. Skeletonizing and model building were done with the program O⁶⁷. The model was refined using X-PLOR⁶⁸ using published protocols⁶⁹. Water molecules were identified in difference density maps and only included into the model if they were present in all three monomers. 2Fobs - Fcalc electron density maps were averaged over the three monomers of the asymmetric unit. The final model of uncomplexed ScrY was used for finding sucrose binding sites in datasets of sucrose-complexed ScrY. Anisomorphism between uncomplexed and sucrose-complexed crystals was dealt with by performing 20 cycles of rigid-body minimization prior to electron density map calculation. 70 N-terminal residues in each monomer were not included into the model, as there was no interpretable electron density for this segment. Solvent molecules and one Ca2+ atom per monomer were added in the final stages of refinement. All atoms were under strong NCS-restraints during the refinement.

Analysis of twinning. In P3(1,2), four non-equivalent ways to index a

given reflection exist, described by (h k l) (k h - l) (-h-k l) or (-k -h -l). For each dataset, only one of these possibilities is suitable for merging with other datasets or difference coefficient calculation, and can be identified on the basis of a low *R*-factor with respect to a reference dataset. For some of these calculations we found that not just one of these possibilities yielded a low *R*-factor but a second also gave an *R*-factor better than that expected for unrelated datasets. The second possibility always corresponded to a rotation of the unit cell by 180° about the axis bisecting the unit cell vectors *a* and *b*: (h-k, k-h, l-). Thus, a hemihedral twinning law of the

$$I_{obs} (h k l) = (1 - \alpha)I_{true}(h k l) + \alpha I_{true}(k h - l)$$
(1)

with a twinning fraction α has to be used to describe the intensities of ScrY datasets. Detwinning can be done using

$$I_{true}(h k l) = ((1 - \alpha)I_{obs}(h k l) + \alpha I_{obs}(k h - l)) / 1 - 2\alpha (2)$$

as long as the twinning fraction a is markedly lower than 0.5. Determination of α by intensity statistics⁷⁰ was not possible as the non-crystallographic symmetry axis nearly coincides with the twinning operation. Therefore α was determined with a method similar to that of Gomis-Rüth et al.²¹ as follows:

(i) For all native datasets, the correlation coefficient between the (h k l) and the (k h -l) reflections (or their symmetry mates) was evaluated; and the dataset with the lowest correlation was defined as the reference. Intensities I_{ref} from this dataset

were preliminarily assumed to be unaffected by twinning. (ii) For each measured dataset with intensities I', the *R*-factor

$$R(\alpha) = (\sum |I' - I(\alpha)|) / \sum I'$$

with $I(\alpha) = (1 - \alpha)I_{ref}(h \ k \ l) + \alpha I_{ref}(k \ h - l)$ with respect to the reference dataset with intensities I_{ref} was calculated, and the value of α that gave the lowest $R(\alpha)$ was taken as the twinning fraction of this dataset. The accuracy of α determined by this procedure was checked with model calculations, and found to be correct within a few percent over the range $\alpha = 0 - 0.5$. Furthermore, the procedure gave good results even in the case of anisomorphism or heavy atom substitution. As an alternative to $R(\alpha)$, we also used the correlation coefficient between I' and $I(\alpha)$, with similar results.

(iii) When a model for ScrY porin became available, the F_{calc} derived from the model were used as the $\alpha = 0$ reference. In this way, the twinning fraction of the previously used reference dataset was found to be $\alpha = 1\%$.

Furthermore, the final model of ScrY was subjected to SHELXL-

| Table 2 Crystallographic | refinement statisti | cs | | |
|--|---------------------|-------------------|--|--|
| | native | sucrose-bound | | |
| esolution (Å) | 15-2.4 | 15-2.4 | | |
| 3₂ cell axes (Å) | 112.1,112.1,147.0 | 111.8,111.8,147.8 | | |
| winning fraction ¹ α | 0.011 | 0.018 | | |
| -factors R _{free} / R _{crvst} (%) | 22.8/21.4 | 24.6/20.7 | | |
| lumber of reflections ($F > \sigma F$) for R_{free} , R_{cryst} | 1,036 / 62,256 | 891 / 54,444 | | |
| ompleteness overall ($F > \sigma F$) (%) | 78.3 | 67.4 | | |
| lumber of: | | | | |
| Non-hydrogen protein atoms | 9,606 | 9,606 | | |
| Sucrose atoms | 0 | 138 | | |
| Solvent molecules | 330 | 330 | | |
| Calcium ions | 3 | 3 | | |
| .m.s. deviation from ideality: | | | | |
| Bond length (Å) | 0.010 | 0.011 | | |
| Bond angles (°) | 1.68 | 1.73 | | |
| Dihedral angles (°) | 28.3 | 28.4 | | |
| t.m.s. deviation of <i>B</i> -factors of covalently bound atoms (Å ²) | 1.8 | 1.9 | | |
| R.m.s. deviation from non-crystallographic ymmetry, all atoms (Å) | 0.10 | 0.10 | | |

'Resulting from SHELXL-96 refinement of α.

 ${}^{2}R_{cryst} = \sum_{i} |\mathbf{F}_{c}| - |\mathbf{F}_{c}| / \sum_{i} |\mathbf{F}_{c}|$ where $|\mathbf{F}_{c}|$ and $|\mathbf{F}_{c}|$ are the observed and scaled calculated structure factor amplitude respectively.

96⁷² refinement, with the twinning fraction α used as a refineable parameter in the program, thus treating twinning in a mathematically adequate way.

Coordinates. The coordinates of the complexed and uncomplexed forms have been deposited in the Brookhaven Protein Data Bank (accession codes 1A0T and 1A0S respectively.)

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