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The bindosome is a structural component of the *Sulfolobus solfataricus* cell envelope

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**Abstract**  Sugar binding proteins of the thermoacidophile *Sulfolobus solfataricus* function together with ABC transporters in the uptake of sugars. They are synthesized as precursors with a class III signal peptide that are normally found in archaeal flagellins and bacterial type IV pilins. The functional expression of sugar binding proteins at the cell surface is dependent on the bindosome assembly system (Bas) that is homologous to bacterial type IV pilin assembly systems. The Bas system consists of an assembly ATPase, BasE; a membrane anchoring protein, BasF; and three small class III signal peptide containing proteins BasABC. Expression of BasEF in a *S. solfataricus* ΔbasEF strain restored the uptake of glucose, while an ATPase mutant of BasE was unable to complement. BasEF was detergent-extracted from *S. solfataricus* membranes as a stable protein complex. Solute binding proteins can be extracted from the cell surface as two high molecular mass complexes of 600 and 400 kDa, wherein the largest complex also contains the main S-layer protein SlaA. Electron microscopic analysis of the cell surface of the wild-type and ΔbasEF strain indicates that the absence of the BasEF complex causes an alteration in cell morphology and the corrugation of the S-layer pattern that is reversed by complementation with the BasEF complex. These results suggest an interaction between the S-layer and the sugar binding proteins that contribute to cell shape.

**Keywords**  Cell envelope · Archaea · Sulfolobus · S-layer

**Introduction**

*Sulfolobus solfataricus* is a thermophilic crenarchaeon belonging to the family of Sulfolobales and it optimally grows at 80°C and at a pH value of 3–4. It was originally isolated from sulfur-rich volcanic areas in the USA, Italy and Iceland and its metabolism and physiology is adapted to these environments (Zillig et al. 1980). From the different strains from Sulfolobales that have been tested for their ability to grow on various minimal media of sugars such as glucose, arabinose and disaccharides such as cellobiose, maltose and lactose, only *S. solfataricus* was found to grow on minimal sugar media (Grogan 1989). The natural environment of *S. solfataricus* is relatively poor in substrates, which necessitates efficient mechanisms of substrate transport across the membrane. In recent years, the ABC transporters in *S. solfataricus* have been studied extensively and several sugar binding proteins were identified that bind sugars at subnanomolar concentrations (Albers et al. 1999a; Elferink et al. 2001).

GlcS is a binding protein that binds glucose and galactose, while AraS binds arabinose, fructose and xylose. In bacteria, binding proteins are directed to the extracellular compartment by means of an N-terminal signal peptide that
upon translocation across the cytoplasmic membrane is removed by a signal peptidase. However, sequence analysis of GlcS and AraS showed that these proteins bear a type IV pilin-like signal peptide or class III signal peptide at their N-terminus (Albers et al. 1999b; Albers and Driessen 2002). Class III signal sequence bearing proteins are normally assembled into an extracellular macromolecular complex such as the pilus or flagellar structures. Class III signal peptides are cleaved by a dedicated type IV signal peptidase which upon processing removes only the positively charged N-terminus leaving a hydrophobic domain of about 20 amino acids attached at the N-terminus of the mature protein (Albers and Driessen 2002; Craig et al. 2004). By the removal of the positive charges, translocation of the protein across the cytoplasmic membrane is possible whereupon the hydrophobic N-terminus acts as scaffold for the assembly of the protein into a supramolecular structure at the extracellular surface of the membrane. In vitro cleavage assays showed that the class III signal peptides of GlcS and AraS are indeed processed by the type IV signal peptidase of S. solfataricus Pbd (Albers et al. 2003). Importantly, the presence of the class III signal peptides predicts that an assembly system is required for the functional expression of the sugar binding proteins at the cell surface. S. solfataricus contains five operons that encode subunits that are homologous to components of bacterial type IV pili assembly systems. Recently, we have shown that the Bas system of S. solfataricus is responsible for the functional expression of sugar binding proteins in the cell envelope (Zolghadr et al. 2007). The Bas system consists of the assembly ATPase, BasE; a membrane protein, BasF; and three small proteins BasABC that are synthesized with a class III signal peptide and that possibly form a pseudopilus-like structure. Deletion of the basEF and basABC genes results in a growth deficiency on various sugars that for uptake depend on a class III signal peptide bearing sugar binding protein. Although the uptake of glucose was impaired in the ΔbasEF and ΔbasABC mutants, the translocation of the binding proteins across the cytoplasmic membrane seemed unaffected suggesting that the Bas system may be involved in correct localization of the binding proteins.

BasE belongs to the family of type IV pil assembly/type II secretion ATPases (Planet et al. 2001; Peabody et al. 2003; Albers and Driessen 2005) that share several conserved motifs in the Walker A and B sites, and the Asp and His Box boxes. In the bacterial ATPases, these conserved motifs are located at the C-terminal domain (CTD) that constitutes the ATP binding and hydrolysis site (Possot and Pugsley 1994; Sandkvist 2001). BasF is a membrane protein that due to its similarity with the EspN protein of the type II secretion system of Vibrio cholerae (Abendroth et al. 2005) is likely to interact with BasE to constitute the functional assembly system.

Currently, the precise role of the Bas system in the localization of the sugar binding proteins remains unclear. We now show that the uptake of glucose is directly dependent on the ATPase activity of BasE, and that BasE and BasF are interacting proteins. Localization studies indicate that the sugar binding proteins are contained in high molecular mass complexes that are associated with the S-layer. The deletion of the basEF genes does not interfere with complex formation but results in cells with an altered morphology and a distorted S-layer. These data indicate that sugar binding proteins are functionally associated with the S-layer.

**Experimental methods**

*S. solfataricus* growth conditions

*S. solfataricus* PBL2025 (Schelert et al. 2004) and the ΔbasEF strain (Zolghadr et al. 2007) were grown in Brock medium at pH 3 and 80°C (Brock et al. 1972) and the medium was supplemented with 0.1% (w/v) of tryptone as sole carbon and energy source. Growth of cells was monitored by measuring the optical density at 600 nm.

Construction of the *basE* mutant

The lysine residue 237 in the Walker A site of BasE was replaced by an alanine by site-directed mutagenesis. The pMZ11 plasmid harboring the basEF genes (pSVA69) was used as a template for the generation of the mutation using the primers 5'-ACGACATTTAATATAAGAGCATTCCC TGAA-3' and 5'-TTTTCAGGGAATGCTCTTATATTAA ATGTG-3'. The mutated basEF was transferred to the SSV1 virus vector with BlnI and EagI restriction sites as described, yielding the plasmid pSVA107 (Zolghadr et al. 2007).

Glucose uptake assays

The uptake of d-[14C]-glucose (250 Ci/mmol) by *S. solfataricus* cells was measured as described (Albers et al. 1999a).

Isolation of cell envelope

Cells were harvested at OD600 of ~1 and resuspended in a buffer containing 50 mM Hepes/NaOH at pH 8 and supplemented with 1 mM PMSF and a small amount of DNase I. Cells were broken by 10 cycles of 10 s sonication and 50 s rest (Soniprep 150, LA Abcoude). Unbroken cells were removed by low spin centrifugation at 4,300 rpm and 4°C for 15 min, and cell envelopes were harvested by
ultracentrifugation at 100,000×g for 1 h at 4°C. Cell envelope pellets were resuspended in 50 mM Hepes/NaOH buffer at pH 8 and frozen in liquid nitrogen and stored at −80°C. Cell envelopes included the cytoplasmic membrane as well as the S-layer and all other extracellular cell attached components.

Purification of BasE/F from *Sulfolobus* cell envelopes

BasE and BasF were co-expressed in the *S. solfataricus ΔbasEF* strain using pSVA75 (Zolghadr et al. 2007) with a C-terminal tandem tag (Strep-tag and 10× His-tag) on BasF. Cell envelopes from the expression strain were isolated as described above and solubilized in buffer A (50 mM Hepes buffer at pH 8) supplemented with 0.05% DDM for 1 h at room temperature. Solubilized membrane proteins were separated from non-solubilized membranes by ultracentrifugation (100,000 ×g for 1 h at 4°C), and the supernatant fraction was applied to a nickel NTA affinity column. Unbound proteins were eluted with 50 mM imidazole and specifically bound proteins were eluted with 200 mM imidazole in buffer A.

For size exclusion chromatography analysis of the BasEF complex, a Superdex 200 column was equilibrated with 50 mM Hepes/NaOH, pH 8 supplemented with 0.05% (w/v) DDM. The chromatography was run with same buffer at 1 ml/min on a AKTA purifier (GE Healthcare) and the fractions were collected at 1 ml per fraction. BasE and BasF antibodies against purified BasE and anti-BasF antibodies. BasF antibodies were generated against a synthetic peptide (QKYKSRAREIRDDIC) of BasF in rabbits and BasE antibodies against purified BasE (Genescript).

Purification of binding proteins by ConA chromatography

Isolated cell envelopes from *S. solfataricus* PBL2025 and *ΔbasEF* cells were solubilized in buffer A (20 mM Hepes buffer at pH 7.5, 100 mM NaCl) supplemented with 2% DDM for 1 h at room temperature. The solubilized fraction was separated from non-solubilized membranes by ultracentrifugation at 100,000×g for 1 h at 4°C. Solubilized membrane proteins were passed onto a 1 ml ConA Sepharose column equilibrated with 10 ml buffer A. The column was washed with 30 ml buffer A, and bound membrane proteins were eluted with buffer A supplemented with 0.05% DDM and 150 mM methyl-α-D-mannopyranoside. The flow through, wash steps and elution fractions were analyzed on 12% SDS-PAGE gel. The uptake of ν-[¹⁴C]-glucose (250 Ci/mmol) by *S. solfataricus* cells was measured as discussed previously (Albers et al. 1999a).

Fixation, freeze substitution and embedding of *S. solfataricus* cells

High-pressure freezing of samples was carried out with high-pressure freezer (Leica EM-PACT 2) as described previously (Rachel et al. 2002). The samples were freeze substituted in a Leica EM AFS (automatic freeze substitution system, Leica, Vienna, Austria) with pure acetone containing 2% (w/v) osmium tetroxide and 0.1% (w/v) uranyl acetate, and 5% H2O at −90°C for 72 h, −60°C for 8 h and −30°C for 6 h. Samples were washed for 1 h in acetone at room temperature and infiltrated in Epon 812 (Ted Pella, Inc., USA) for 30 h and the Epon was polymerized at 60°C for 24 h. Ultrathin sections were cut with a Leica UCT ultramicrotome and mounted on carbon-coated copper grids. Transmission electron micrographs were recorded using a slow-scan CCD camera (TVIPS, Gauting, Germany) on a Philips CM 12 (FEI, Eindhoven, The Netherlands) operated at 120 kV, which is also equipped with a Gatan TV 673 wide-angle camera (Gatan, München) for searching and focusing.

Freeze-etching of *S. solfataricus* cells

Cells were harvested by centrifugation at 5,000 rpm for 15 min and a 1.5-µl aliquot of the concentrated cell suspension was loaded into a gold-carrier and frozen in liquid nitrogen. The sample was cut with a cold knife (T ≤ −185°C) and etched for 4 min (T = −97°C; p ≤ 1.3 × 10⁻⁵ Pa) in a CFE-50 freeze-etch unit (Cressington, Watford, UK). The freeze-etched sample was shadowed (1 nm Pt/C 45°, 10 nm C 90°) and cleaned on H2SO4 (70%). Replicas were picked up with 700 mesh hexagonal copper grids. Transmission electron micrographs were taken as above.

Results

Characterization of BasEF by site-directed mutagenesis

To determine the role of the ATPase activity of BasE, the conserved lysine residue (K237) of the Walker A motif was replaced by an alanine residue by site-directed mutagenesis. In general, the introduction of this mutation in homologous assembly ATPases results in a loss of the ATPase activity and a functional defect (Herdendorf et al. 2002; Sexton et al. 2004). The mutated basE gene was subcloned with basF into the virus expression vector pSVA9 of *S. solfataricus* to yield pSVA107. The pSVA107 plasmid was introduced into the ΔbasEF strain and glucose uptake was measured and compared to a strain transformed with pSVA75 containing the native basEF genes (Zolghadr...
et al. 2007; Albers et al. 1999a). The cells complemented with BasE(K237A)F showed a glucose transport activity similar to the negative control, whereas cells transformed with the wild-type BasEF showed high levels of glucose uptake (Fig. 1). The results demonstrate that the ATPase activity of BasE is essential for active glucose transport in *S. solfataricus*.

**BasE and BasF co-purified as a complex**

Since the expression of BasEF in the ΔbasEF strain restored the glucose uptake, we performed a detergent extraction and purification of the Bas complex from these cells in order to determine the complex subunit composition. Herein, BasE and BasF were co-expressed in *S. solfataricus* cell envelopes with a C-terminal tandem tag (Strep-tag/10× His-tag) on BasF. After solubilization BasF was purified using His-tag affinity chromatography and eluted with the expected molecular size of 75 kDa (Fig. 2a) and various other proteins. The identity of BasF was confirmed by immunoblotting using an antibody directed against BasF (Fig. 2b). The elution fraction contained another protein with a size of ~60 kDa (Fig. 2a) which was identified as BasE by immunoblotting using a polyclonal antibody raised against the purified BasE. BasF was also identified by mass spectrometry (data not shown). However, this protein band also contained tf55α, a subunit of the thermosome of *S. solfataricus* which has a similar mass as BasE which explains the greater band intensity of BasE as compared to BasF. These data indicate that BasF and BasE are interacting proteins forming a stable membrane protein complex.

To determine if BasEF is present in a high molecular mass complex, the partial purified BasEF complex was subjected to size exclusion chromatography using a Superdex 200 column and 0.1% dodecylmaltoside (Fig. 3a). Fractions were analyzed on SDS-PAGE (Fig. 3b), and by immunoblotting with BasF and BasE antibodies (Fig. 3c, d, respectively). The first elution peak with an approximate mass of 1 MDa contained the tf55α protein which is forming octa- and nonamers hence the high molecular weight peak (Fig. 3b, lane 1; Schoehn et al. 2000). The second peak, which corresponded to a size of 100–200 kDa, contained the BasE and BasF proteins (Fig. 3b, lane 3). The third peak contained a protein with a mass of approximate 40 kDa (Fig. 3b, lane 4). Mass spectroscopy analysis failed to reveal the identity of 40 kDa protein. The estimated size of the BasEF elution fraction was smaller than expected. A stoichiometric complex of BasEF is predicted to have a molecular mass of ~135 kDa. However, ATPases homologous to BasE form a hexameric ring (Herdendorf et al. 2002; Yamagata and Tainer 2007), and thus a putative BasE hexamer alone would correspond to a mass of ~360 kDa. The amount of BasF molecules in the complex is unknown, but considering that the BasF is 75 kDa while taking the

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**Fig. 1** Uptake of glucose by the ΔbasEF strain without (open circles) and with a plasmid containing BasEF (pSVA75) (closed squares) or BasE(K237A)F (pSVA107) (open squares)

**Fig. 2** Co-purification of the BasE and BasF proteins expressed in *S. solfataricus* ΔbasEF cells. Isolated membranes were solubilized in 2% DDM, and solubilized proteins were passed over a Ni-NTA affinity column and eluted with 100 mM imidazole. Fractions were analyzed on SDS-PAGE (a), and by Western blotting using antibodies against BasF (b) and BasE (c). Start material, flow through and the 200 mM imidazole elution fraction were loaded in lanes 1, 2 and 3, respectively.
detergent micelle into account, a minimum size of 500 kDa is expected for the BasEF complex. The intense appearance of the BasE band in Fig. 3b (lane 3) is most likely due to the presence of trimers/dimers of tf55 as identified by mass spectrometry. Therefore, the eluted fraction most likely corresponds to a stoichiometric BasEF complex.

Sugar binding proteins on S. solfataricus are assembled into high molecular complexes

To determine if the sugar binding proteins are contained in large complexes at the cell surface, binding proteins were extracted from cell envelopes derived from S. solfataricus wild-type and ΔbasEF cells using a low Triton X-100 concentration, and partially purified by ConA lectin affinity chromatography. The identity of the proteins was confirmed by mass spectrometry (Fig. 4). When analyzed by Blue-native PAGE, the ConA-enriched binding proteins were found in two main complexes with apparent molecular masses of about 600 and 400 kDa, respectively. These two complexes were excised, eluted and subsequently run on an SDS-PAGE (b) to show their subunit composition. For comparison the initially isolated Con A fraction is displayed in the first lane. The identity of the proteins was determined by mass spectrometry.

The complexes observed in the BN-PAGE showed a distinct protein composition on SDS-PAGE. Protein complex 1 contained the main S-layer protein (SlaA) and a protein at 120 kDa that by mass spectrometry was
identified as a putative dipeptide binding protein, SSO1273 (Fig. 4b). SlaA has previously been detected to represent a major component of the ConA purification fraction (Ellen et al. 2009) and found to be glycosylated (data not shown). Protein complex 2 consists of the glucose binding protein (GlcS) and another putative dipeptide binding protein SSO3043 as validated by mass spectrometry (Fig. 4). Both SSO3043 and SSO1273 are synthesized with a type I signal peptide. It should be emphasized that the annotation of SSO3043 and SSO1273 as putative dipeptide binding proteins may not necessarily mean that these proteins bind dipeptides as previous analysis of such proteins revealed that most represent sugar binding proteins (Elferink et al. 2001; Nanavati et al. 2006). Especially, SSO3043 is surrounded by enzymes annotated to be involved in sugar degradation. Interestingly, when the growth medium was supplemented with 0.4% arabinose, the arabinose binding protein (AraS) was induced and found to be a part of protein complex 2 on BN-PAGE (Fig. 4a, b, Band 2). These results demonstrated that the (sugar) binding proteins are contained in high molecular mass complexes with a relatively uniform size. The presence of SlaA in one of these complexes suggests that the binding proteins are associated with the S-layer.

Analysis of S. solfataricus surface layer

In an attempt to visualize the bindosome structure in the cell envelope of S. solfataricus, we examined the cell surface by transmission electron microscopy (TEM). Herein, cells were subjected to high-pressure freezing, freeze substitution and TEM to visualize highly defined structures of the cell envelope of the PBL2025 control strain and the ΔbasEF mutant strain. TEM images showed a well-preserved cell surface from both strains in ultrathin sections (Fig. 6). Interestingly, the morphology of wild-type cells appeared as highly lobed and irregular shaped cocci, while the ΔbasEF cells were less lobed, almost regular cocci (Fig. 6a, b). Possibly, these structural changes in the cell envelope resulted from the basEF deletion. The membrane and S-layer were clearly distinguished and enclose a quasi-periplasmic space with 25 nm in width (Fig. 6c, d). The periplasmic space appeared well contrasted, due to the exposure of the cells to OsO4 and uranyl acetate during the freeze substitution. This indicates the presence of an abundance of biologically relevant components. Immuno-gold labeling was performed to study the
localization of binding protein in the cell envelope. However, the antibodies used to localize GlcS, the glucose binding protein, gave very poor labeling, while another antibody against the A1A0 ATPase (Küper et al. 2010) resulted in significant and dense labeling of the cytoplasmic membrane (data not shown). Therefore, no final conclusion about the localization of GlcS could be obtained.

In another approach, the surface layer of the PBL2025 and \( \Delta \)basEF cells was visualized by freeze-etching freshly grown cells (Fig. 7). In these freeze-etch preparations, the S-layer of strain PBL2025 showed a crystalline lattice of pseudo-hexagonal symmetry, reflecting S-layer protein complexes with internal threefold symmetry, arranged on hexagonal lattice, in a similar manner as described (Fig. 7a, c). The crystalline lattice of the \( \Delta \)basEF S-layer appeared to be less corrugated (Fig. 7b, d).

As additional evidence that the Bas system is indeed involved in determining the cell shape of \( \textit{S. solfataricus} \), we complemented the \( \Delta \)basEF strain with the plasmid expressing the BasEF complex. From these cells, TEM images were obtained as described before for the wild-type and \( \Delta \)basEF cells as well as freeze-etch preparations to visualize the S-layer. Expression of BasEF restored the lobed cell shape (Fig. 8b/d) as compared to the round \( \Delta \)basEF (Fig. 6c/d). Also, the S-layer appeared like in the wild type, although disturbances were visible in the regular arrangement. These data show that the deletion of basEF results in a disturbance of the surface layer lattice, which likely causes a different cell morphology and an altered localization of the sugar binding protein complexes in the cell envelope.

Discussion

Here, we have explored several approaches to study the function of the Bas system and its role in the correct localization of the sugar binding proteins in the cell envelope of \( \textit{S. solfataricus} \). The Bas system plays an important role in the functional assembly of sugar transport in \( \textit{S. solfataricus} \). This is evident from the sugar transport defect of the basEF deletion mutant that can be complemented by the expression of BasEF. However, expression of BasE(K237A)F with a mutation of the catalytic lysine residue of the ATP binding site of BasE did not result in complementation. This implies that the ATPase activity of BasE is required for the correct localization and/or assembly of the sugar binding proteins. In a previous study, we could demonstrate that BasEF are not
needed for the translocation of sugar binding proteins across
the cytoplasmic membrane (Zolghadr et al. 2007). Moreover,
in the absence of BasEF, the extracellular binding proteins
still bind sugars with high affinity. However, these
cells are unable to grow on sugars that for uptake depend on a
binding protein that is synthesized with a class III signal
peptide, and also sugar transport is impaired. BasE and BasF
are homologous to subunits of bacterial type II secretion and
type IV pili assembly systems. BasE is a cytosolic ATPase
and BasF is a membrane protein that likely functions to
anchor BasE to the membrane. Indeed, BasE and BasF form a
complex, but this complex seems labile under the detergent
extraction conditions tested. Size exclusion chromatography
hints at the presence of a stoichiometric complex rather than
a hexameric arrangement of the BasE subunit. Stabilization
of the BasEF complex for future structural analysis remains
challenging as the complex is readily degraded by endoge-
nous proteases (unpublished results).

An analysis of the ConA purified sugar binding proteins
by BN-PAGE and size exclusion chromatography revealed
that these proteins are present in a large complex. These
complexes could be separated as symmetric peaks on size
exclusion chromatography. However, the binding proteins
purified from the ΔbasEF strains were present in a similar
complex indicating that BasEF is not required for complex
formation per se. Possibly, the role of BasEF is to correctly
localize the binding protein complexes at the surface layer
envelope since some of the binding proteins were found to
be associated with the major S-layer protein SlaA. Analysis
of the surface envelope by high-pressure freezing and freeze
substitution yielded high-resolution images of cytoplasmic
membrane, quasi-periplasmic space and S-layer. The TEM
images of the S-layer suggested that the Bas system does
not just generate a “traditional” filament comparable to
flagella or pili, but that it rather is involved in or contributes
to shaping the cell envelope. It appears that the deletion of
basEF has a disordering effect on the self-assembly of the
membrane-integral SlaB (Veith et al. 2009). In this respect,
the sugar binding complexes isolated from the cell surface
may represent partially dissociated complexes and part of
a large native structure in the cell envelope that is a struc-
turally associated with the S-layer. The quasi-periplasmic
space of S. solfataricus is composed of dense material, most
probably glycan chains from lipids, glycosylated proteins
and extracellular proteins (see, e.g., the scheme of the
S-layer architecture of Halobacterium spp., Kessel et al.
1988), which may be essential for the overall stability
of cell morphology and cell wall. Such high density may

Fig. 7 Electron micrographs of
freeze-etched S-layers of the
PBL2025 (a, c) and ΔbasEF
(b, d) strains. Sizes of the bars
are in a and b are 500 nm and in
c and d 200 nm
limit the diffusion of extracellular solutes, and assuming that this space amongst others is filled with sugar binding proteins, a specific organization of these binding proteins may facilitate the diffusion of solutes from the outer surface of the S-layer to the cytoplasmic membrane as it is known from Gram-negative bacteria (Brass et al. 1986).

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