

Agl16, a Thermophilic Glycosyltransferase Mediating the Last Step of *N*-Glycan Biosynthesis in the Thermoacidophilic Crenarchaeon *Sulfolobus acidocaldarius*

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Recently, the S-layer protein of *Sulfolobus acidocaldarius* was shown to be N-linked with a tribranched hexasaccharide, composed of Man₂Glc₁GlcNAc₂ and a sulfated sugar called sulfoquinovose. To identify genes involved in the biosynthesis and attachment of this glycan, markerless in-frame deletions of genes coding for predicted glycosyltransferases were created. The successful deletion of *agl16*, coding for a glycosyltransferase, resulted in the S-layer protein and archaellins having reduced molecular weights, as visualized by Coomassie staining or immunoblotting. This analysis indicated a change in the *N*-glycan composition. Nano-liquid chromatography-tandem mass spectrometry (LC-MS/MS) analyses confirmed that the glycan of the S-layer protein from the *agl16* deletion mutant was a pentasaccharide, which was missing a terminal hexose residue. High-performance liquid chromatography (HPLC) analyses of the hydrolyzed *N*-glycan indicated that the missing hexose is a glucose residue. A physio-logical characterization of the *agl16* deletion mutant revealed a significant effect on the growth at elevated salt concentrations. At 300 mM NaCl, the doubling time of the $\Delta agl16$ deletion strain affected the assembly and function of the archaellum, as exemplified by semisolid Gelrite plate analysis, in which the motility is decreased according to the *N*-glycan size.

The majority of the *Archaea* possess a proteinaceous surface layer (S-layer) as the sole cell wall structure (1), which contributes to cell stability and maintenance of the cell morphology. Archaeal S-layer proteins (2–10), as well as other surface-exposed proteins, like pilins (11), sugar-binding proteins (12), and the archaeal flagellins (13, 14), now termed archaellins (15), are posttranslationally modified via glycosylation. This posttranslational modification affects protein stability (16) and influences the assembly of surface-exposed proteins (17).

Protein glycosylation, in which sugar residues or oligosaccharides are covalently linked mainly to either selected asparagine (*N*-glycosylation) or serine and threonine residues (*O*-glycosylation) within a peptide sequence, is probably the most predominant posttranslational modification. In contrast to *Bacteria*, where *N*-glycosylation is a rare process, found only in a few species (18), almost all sequenced *Archaea*, with just a few exceptions, possess the gene encoding the key enzyme of the *N*-glycosylation process, the oligosaccharyltransferase AglB (19, 20). In the past few years, research on archaeal *N*-glycosylation performed in three members of the *Euryarchaea* revealed that *Archaea* exhibit features of both eukaryal and bacterial glycosylation pathways (21). Genes involved in this *N*-glycosylation pathway are designated *agl* (archaeal glycosylation) (21, 22).

Surface-exposed proteins of the crenarchaeon *Sulfolobus acidocaldarius*, like cytochrome $b_{558}/566$ (23), sugar-binding proteins (24), the large S-layer protein SlaA (7), and the archaellin (14), are modified by the attachment of glycans. The cytochrome $b_{558}/566$, which possesses *O*- and *N*-linked glycans, was the first crenarchaeal glycoprotein in which the composition and structure of the *N*-glycan were determined (25). Analysis of *O*-linked sugars detected only mannose residues (23), whereas the detailed analyses of the *N*-linked glycans revealed a tribranched hexasaccharide

composed of Glc1Man2GlcNAc2 and a sulfated sugar called sulfoquinovose. Analysis of the S-layer glycoprotein SlaA confirmed the same N-glycan and revealed an astonishingly high glycosylation density (7). Beside the high glycosylation density, which is more similar to the highly glycosylated eukaryal cell wall protein ECM33 of Saccharomyces cerevisiae, the presence of the chitobiose core is a further eukaryal characteristic of the N-glycans which has exclusively been found in Sulfolobus so far. Furthermore, incorporation of the rare sulfated sugar sulfoquinovose (QuiS) into the N-glycan has not been detected in any other Archaea. Sulfoquinose has so far been found only as the head group of the nonphosphorous lipid sulfoquinovosyldiacylglycerol, which is incorporated into the photosynthetic membrane of all higher plants, mosses, ferns, and algae, as well as in most photosynthetic bacteria (26, 27). Deletion of the UDP-sulfoquinovose synthesis gene in S. acidocaldarius resulted in the loss of incorporation of sulfoquinovose in the N-glycan as well as one mannose and the glucose residues, implying the highly ordered assembly process of the N-glycans (14). However, unlike some Archaea, in which glycosyltransferases (GTases) are clustered directly next to the oligosaccharyltransferase AglB (19), no aglB-based gene clusters of GTases are present in Sulfolobus, which makes the identification of GTases participating in the N-glycosylation more difficult. Identification of GTases on the basis of their homologies to known N-glycosyla-

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TABLE 1 Plasmids used in this study

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Plasmid	Insert	Reference
pSAV407	Gene targeting plasmid, pGEM-T Easy backbone, pyrEF cassette of S. solfataricus ^a	28
pSAV1233	In-frame deletion of <i>agl16</i> cloned into pSVA407 with ApaI and BamHI	This study
pMZ1	Gene targeting plasmid with a C-terminal Strep tag and $10 \times$ His tag	30
pSAV1450	pRN1-based shuttle vector with <i>lacS</i> reporter gene of <i>S. solfataricus^b</i>	28
pSVA1273	Full <i>agl16</i> gene insert cloned into pMZ1 with NcoI and BamHI	This study
pSAV1274	agl16 insert from pSVA1273 cloned into pSVA1450 with NcoI and EagI	This study

^{*a*} *pyrEF* encode the enzymes orotate phosphoribosyltransferase and orotidine-5-monophosphate decarboxylase, respectively.

^b lacS encodes β-D-galactosidase of S. solfataricus.

tion components in *Eukarya, Bacteria*, or *Archaea* led to no specific hits for *S. acidocaldarius*, and deletion studies of several genes coding for predicted GTases in the genome of *S. acidocaldarius* revealed no alteration of the *N*-glycan linked to the S-layer protein SlaA except for *saci0807*.

In this study, we identified the first thermophilic GTase, Agl16 (Saci0807), that participates in the crenarchaeal *N*-glycosylation process. Recently developed genetic tools for *S. acidocaldarius* as well as biochemical and mass-spectrometric methods were used to verify the predicted function of the Agl16 GTase in the biosynthesis of the S-layer and archaellin *N*-glycan.

MATERIALS AND METHODS

Strains and growth conditions. The strain *S. acidocaldarius* MW001 ($\Delta pyrE$) (28) and the $\Delta agl16$ mutant were grown in Brock medium at 75°C and pH 3, adjusted using sulfuric acid. The medium was supplemented with 0.1% (wt/vol) NZ-amine and 0.1% (wt/vol) dextrin as carbon and energy sources (29). Gelrite (0.6%) selection plates were supplemented with the same nutrients (as listed above), with the addition of 10 mM MgCl₂ and 3 mM CaCl₂. For second selection, plates containing 10 μ g ml⁻¹ uracil and 100 μ g ml⁻¹ 5-fluoroorotic acid (5-FOA) were added. For the growth of the uracil auxotrophic mutant MW001 and $\Delta agl16$ strains, 10 μ g ml⁻¹ uracil was added to the medium. The cell growth was monitored by measuring the optical density at 600 nm. For propagation of plasmids, *Escherichia coli* DH5 α cells were used.

Construction of a deletion plasmid. Plasmids used in this study are listed in Table 1. To verify whether the gene saci0807, encoding the GTase Agl16, is indeed important for the N-glycosylation pathway, a markerless deletion mutant of this gene was created as previously described (28, 31). Briefly, the uracil-auxotrophic strain MW001, in which the pyrE gene is disrupted, was transformed with plasmid pSVA1233. To construct the plasmid, 800 to 1,000 bp of the up- and downstream fragments of agl16 were PCR amplified. At the 5' end of the upstream and at the 3' end of the downstream fragment, the restriction sites for ApaI and BamHI were introduced by the forward and reverse primers, respectively. The upstream reverse primer and the downstream forward primer were designed to each incorporate 15 bp of the reverse complement strand of the other primer, resulting in a 30-bp overlapping stretch. The up- and downstream fragments were fused by an overlapping PCR, using the 3' ends of the upand downstream fragments as primers. The amplified overlapping PCR fragment was purified by electrophoresis in 0.8% agarose gel using a Nucleospin extract kit (Macherey-Nagel, Düren, Germany) and digested with ApaI and BamHI. After repurification, the overlap fragment was ligated into an ApaI- and BamHI-predigested plasmid, pSVA407, containing a pyrEF cassette (28). The constructed plasmid, pSVA1233, was

transformed into *E. coli* DH5 α and selected on LB plates containing 50 µg ml⁻¹ ampicillin. The accuracy of the plasmid was ascertained by sequencing. In order to avoid restriction in *S. acidocaldarius*, the plasmid was methylated by transformation in *E. coli* ER1821 cells containing pM.EsaBC4I (available from NEB), which expresses a methylase.

Transformation and selection of the deletion mutant in S. acidocaldarius. Preparation of competent cells was performed according to the protocol of Kurosawa and Grogan (32). Briefly, S. acidocaldarius strain MW001 was grown in Brock medium supplemented with 0.1% (wt/vol) NZ-amine and 0.1% dextrin until an optical density at 600 nm (OD_{600}) between 0.1 and 0.3 was reached. Cooled cells were harvested by centrifugation $(2,000 \times gat 4^{\circ}C \text{ for } 20 \text{ min})$. The cell pellet was washed once each in 50 ml, 10 ml, and 1 ml of ice-cold 20 mM sucrose (dissolved in demineralized water) after mild centrifugation (2,000 \times g at 4°C for 20 min). The final cell pellet was resuspended in 20 mM sucrose to attain an OD₆₀₀ of 10 and stored in 50-µl aliquots at -80°C. Methylated pSVA1233 plasmid (400 to 600 ng) was added to the 50-µl aliquot of competent MW001 cells and incubated for 5 min on ice before transformation in a 1-mm-gap electroporation cuvette at 1,250 V, 1,000 Ω , and 25 mF using a Bio-Rad gene pulser II. Directly after transformation, 50 µl of a 2×-concentrated recovery solution (1% sucrose, 20 mM beta-alanine, 20 mM malate buffer [pH 4.5], 10 mM MgSO₄) was added to the sample and incubated at 75°C for 30 min under mild shaking conditions (150 rpm). Before plating, the sample was mixed with 100 µl of heated 2×-concentrated recovery solution, and two 100-µl portions were spread on Gelrite plates containing Brock medium supplemented with 0.1% NZ-amine and 0.1% dextrin. After incubation for 5 to 7 days at 78°C, large brownish colonies were used to inoculate 50 µl of Brock medium containing 0.1% NZ-amine and 0.1% dextrin, each culture was incubated for 3 days of 78°C. Each culture, which was confirmed by PCR to contain genomic incorporated plasmid pSVA1233, was grown in Brock medium supplemented with 0.1% NZamine and 0.1% dextrin until an OD of 0.4. Aliquots of 40 µl were spread on second-selection plates supplemented with 0.1% NZ-amine, 0.1% dextrin, and 10 mg ml⁻¹ uracil and incubated for 5 to 7 days at 78°C. Newly formed colonies were streaked onto new second-selection plates to ensure that they were formed from single colonies, before each colony was screened for the genomic absence or presence of the agl16 gene by PCR. The deletion of agl16 was further confirmed by sequencing and reverse transcription-PCR (RT-PCR).

RT-PCR. Primers used in this study are listed in Table 2.The RNA isolation was performed using an RNeasy Plus mini-extraction kit (Qiagen, Hilden, Germany), according to the manufacturer's instructions. The concentration of RNA was analyzed by spectrophotometrically. To eliminate the genomic DNA, two repetitive 2 h of incubation with DNase I (Fermentas, St. Leon-Rot, Germany) followed by repurification steps with the RNeasy Plus mini-extraction kit were performed. Purified total RNA was checked by PCR for the lack of contaminating chromosomal DNA, before cDNA synthesis with a First Strand cDNA synthesis kit (Fermentas) was used, according to the manufacturer's instruction. To verify the deletion of *agl16* by RT-PCR, a specific forward primer (4101) and reverse primer (4106) amplifying the internal region of *agl16* and, as a control, primers specific for the internal region of *aglB* (1727 and 1730) were designed and used in a PCR with single-stranded cDNA as a template.

S-layer isolation. Cell pellets of frozen or fresh cells from a 50-ml culture were incubated under shaking condition for 45 min at 37°C in 40 ml of buffer A (10 mM NaCl, 1 mM phenylmethylsulfonyl fluoride [PMSF], 0.5% sodium lauroylsarcosine) with the addition of a small amount of DNase. Samples were centrifuged for 40 min in an Optima Max-XU ultracentrifuge (Beckman Coulter) at 21,000 rpm, yielding a brownish tan pellet. The pellet was resuspended and incubated for 30 min at 37°C in 1.5 ml of buffer A. Repetitive steps of washes in buffer B (10 mM NaCl, 0.5 mM MgSO₄, 0.5% SDS), incubation for 20 min at 37°C, and subsequent centrifugation (tabletop centrifuge at 14,000 rpm), yielded in

TABLE 2 Oligonucleotide primers used in this study

Primer	Sequence $(5'-3')^a$	Restriction site	Description and target gene
1848	CTCTAC <u>GGGCCC</u> ACTTCTGTCTCCCCACTGG	ApaI	Up-for, $\Delta agl16$
1849	AAGGTCTGGATGTTGTAAAAATTATGTCCATGACATCACAGGACA		Up-rev, $\Delta agl16$
1850	TGGACATAATTTTTACAACATCCAGACCTTATACATGAATCATCC		Down-for, $\Delta agl16$
1851	CGCCGA <u>GGATCC</u> AACCATGGAATTGTAGCC	BamHI	Down-rev, $\Delta agl16$
4101	GTCCTCAAGACCCAATTCACTAAC		RT-PCR-for, agl16
4106	GCCGACTGCAGCATTAGTAATGACTCCACACCAC		RT-PCR-rev, agl16
2799	CATGC <u>GGATCC</u> TTAACTTAAACTTTTATAGAGCTCAAGC	BamHI	Forward primer complementing agl16
4100	CATGC <u>CCATGG</u> ATGTATAAGGTCTGGATGTTGACCCCAC	NcoI	Reverse primer complementing agl16
1729	CTGCTGCAATTACAGCGTTC		RT-PCR-for <i>aglB</i>
1730	AACCGTGAGCTACTTCAGAC		RT-PCR-rev aglB

^a Underlining indicates restriction sites.

a translucent tan pellet of S-layer proteins. Purified S-layer proteins were washed once with water and then stored in water at 4°C.

Induction and immunostaining of the archaellin FlaB. Pellets of fresh 50-ml cultures of MW001 and the $\Delta agl16$ strain were resuspended and grown for 4 h at 79°C in 40 ml of Brock medium lacking NZ-amine as a carbon and energy source to induce the production of archaella (33). Centrifuged cells were resolved in 1 ml of 20 mM Tris HCl and 500 mM NaCl buffer. Aliquots of 30 µl were loaded on a 11% SDS-PAGE gel, electrophoresed (100 V), and blotted onto a polyvinylidene difluoride (PVDF) membrane. For detection, the primary antibody raised in rabbits against a FlaB peptide (Eurogentec) and an alkaline phosphatase-coupled anti-rabbit IgG antibody (Sigma-Aldrich, St. Louis, MO) were used. The chemifluorescence signal was measured in a Fujifilm LAS-4000 luminescent image analyzer (Fujifilm, Düsseldorf, Germany).

Motility assay. Swimming motility was analyzed on semisolid plates consisting of 0.15% Gelrite supplemented with Brock medium with a reduced amount of NZ-amine (0.0015%) and either 0 mM or 300 mM NaCl. To inoculate the plates, aliquots of 5 to 7 μ l, corresponding to the OD₆₀₀ of MW001 or $\Delta agl16$ cultures (grown to an OD₆₀₀ of 0.5), were dropped on soft Gelrite plates. Plates were incubated 4 to 9 days in a humid chamber at 75°C. Swimming behavior of the different *Sulfolobus* strains was analyzed by measuring the swimming radius.

Complementation of agl16 by cloning of agl16 into the S. acidocaldarius expression vector. To complement the defect in the biosynthesis of the N-glycan in the $\Delta agl16$ strain, the gene saci0807 (agl16) was amplified from genomic DNA of S. acidocaldarius, introducing BamHI and NcoI restriction site at the 5' ends of primers 2799 and 4100, respectively. The PCR fragment was cloned into pMZ1 (30), yielding vector pSVA1273, the plasmid was digested with NcoI and EagI and the insert containing the agl16 gene was ligated into pSVA1450, an improved expression vector (M. Wagner and S. V. Albers, unpublished data) based on pCmalLacS (34), yielding pSVA1274. The pCmalLacS plasmid contains an inducible maltose promoter that is used for the expression of agl16 in S. acidocaldarius. Before transformation into the $\Delta agl16$ deletion strain, the plasmid was methylated (see "Transformation and selection of the deletion mutant in S. acidocaldarius" above). Transformed cells were plated on first-selection Gelrite plates, and single colonies were picked and induced in Brock medium with 0.4% maltose. Pellets of a culture grown to an OD of 0.8 were used for the S-layer isolation.

Tryptic digestion for the mutant S-layer samples. Purified S-layer mutants were run on a 2 to 8% precast gel (Invitrogen, Paisley, United Kingdom) and stained with Novotex colloidal blue stain (Invitrogen). The stained band was excised from the gel and cut into relatively small pieces and destained, using 400 ml of 50% (vol/vol) acetonitrile in 0.1 M ammonium bicarbonate (pH 8.4) and dried in a SpeedVac. Dried gel pieces were typically reswollen in 20 µl of 50 mM ammonium bicarbonate (pH 8.4) containing 0.5 ng ml⁻¹ trypsin (Promega catalog no. V5111) and incubated at 37°C, overnight. The supernatant was transferred to a clean Eppendorf tube. In order to stop the tryptic reaction, the gel pieces were incubated (37°C for 10 min) in 0.1% trifluoroacetic acid (TFA) (50 ml)

followed by incubation (37°C for 10 min) in acetonitrile (100 ml) to shrink the gel pieces. The supernatant was then added to the previously collected glycopeptides and the final volume reduced to \sim 20 to 40 µl for nano-LC-ES-MS analysis.

Nano-LC-ES-MS/MS analysis. Tryptic digests were analyzed by nano-LC-ES-MS/MS using a reverse-phase nano-HPLC system (Dionex, Sunnyvale, CA) connected to a quadrupole time-of-flight (TOF) mass spectrometer (Q-STAR Pulsar I; MDS Sciex). The digests were separated by a binary nano-high-performance liquid chromatography (nano-HPLC) gradient generated by an Ultimate pump fitted with a Famos autosampler and a Switchos microcolumn switching module (LC Packings, Amsterdam, The Netherlands). An analytical C_{18} nanocapillary (75 μ m [inside diameter] by 15 cm; PepMap) and a micro-precolumn C₁₈ cartridge were employed for on-line peptide separation. The digest was first loaded onto the precolumn and eluted with 0.1% formic acid (Sigma) in water (HPLC grade; Purite) for 4 min. The eluant was then transferred onto an analytical C18 nanocapillary HPLC column and eluted at a flow rate of 150 nl/min using the following gradient of solvent A [0.05% (vol/ vol) formic acid in a 95:5 (vol/vol) water/acetonitrile mixture] and solvent B [0.04% formic acid in a 95:5 (vol/vol) acetonitrile/water mixture]: 99% A from 0 to 5 min, 99 to 90% A from 5 to 10 min, 90 to 60% A from 10 to 70 min, 60 to 50% A from 70 to 71 min, 50 to 5% A from 71 to 75 min, 5% A from 75 to 85 min, 5 to 95% A from 85 to 86 min, and 95% A from 86 to 90 min. Data acquisition was performed using Analyst QS software with an automatic information-dependent-acquisition (IDA) function.

HPLC analyses. The *N*-glycans of purified S-layer derived from a 25-ml culture were hydrolyzed in 2 N TFA for 20 h at 80°C. After the treatment, the S-layer was removed from the sample by centrifugation in a tabletop centrifuge at 14,000 rpm for 20 min. An 80-µl portion of the clarified supernatant was acidified with H_2SO_4 (4 µl, 1 M). Subsequently this solution was analyzed by high-performance liquid chromatography (HPLC) using an 8-µm H⁺ IEX resin column (250 by 8 mm [inside diameter]; Grom, Rottenburg, Germany) at 80°C and a mobile phase of 5 mM H_2SO_4 (0.35 ml min⁻¹).

Transmission electron microscopy (TEM). For TEM, cells were fixed with 2.5% glutaraldehyde for 15 min at room temperature. Cell suspensions were placed directly on glow-discharged Formvar-coated 200-mesh copper grids (Plano, Wetzlar, Germany). The samples were washed once with water, negatively stained for 15 s with 2% uranyl acetate, and air dried. Grids were investigated on a JEOL 3010 300-kV transmission electron microscope (JEOL, Eching, Germany).

RESULTS

Identification of *saci0807*, its genomic localization, and protein characterization. To elucidate the *N*-glycosylation pathway in *S. acidocaldarius*, the genome was screened for the presence of genes coding for predicted GTases. However, none of these identified GTases (Table 3) possesses a high sequence similarity to known eukaryal, bacterial, or archaeal GTases involved in the *N*-glycosy-

TABLE 3 Overview	of genes	encoding	predicted	GTase	in S.
acidocaldarius			-		

GTase family	3D fold, mechanism of sugar transfer	<i>S. acidocaldarius</i> gene(s) encoding GTase	No. of GTase genes
4	GT-A, inverting GT-B, retaining	saci0124, saci0275, saci0956, saci1011, saci1051, saci1499, saci1909, saci1911, saci1915, saci1927 saci0807, saci0869, saci1249,	10
		saci1827, saci1904, saci1907, saci1914, saci1916, saci1921, saci1922, saci1923	
5	GT-B, retaining	saci1201	1
35	GT-B, retaining	saci0294	1
39	GT-C, inverting	saci0421	1
66	GT-C, inverting	saci1274 (aglB)	2
Not classified		saci0201, saci0870, saci1865, saci2027	4

lation pathway. Deletion studies of a variety of putative GTases in the genome of *S. acidocaldarius* were fruitless, except that deletion of *saci0807*, encoding a predicted GTase, resulted in an alteration in the biosynthesis of the *N*-linked hexasaccharide. The gene *agl16* (*saci0807*) encodes a soluble 40.6-kDa protein with a glycosyltransferases 1 domain (PF00534) spanning residues 167 to 330, whereas outside this region the protein contains no other known domain. However, the broad range of functions related to the glycosyltransferase 1 domain did not allow a precise prediction of the function of this GTase. The CAZy database grouped Agl16, based on similarity of short sequence motifs, in GTase family 4, with a GT-B fold and a retaining enzyme according to the stereochemistry of the substrates and reaction products. Besides the vast number of different GTase sequence families, only a limited number of structural folds have been determined, in which the majority of GTases are found to be either GT-A or GT-B folded. In GT-A-folded proteins, only a single catalytic domain is found, whereas GT-B-folded enzymes possess two catalytically distinct domains, separated by a stretch which binds the acceptor. All domains possess structural elements which are similar to the Rossmann fold. The C-terminal domain of the GT-B-folded GTases mediates the binding of the nucleotide-activated sugar. In contrast to the GT-A GTases, the catalysis of GT-B GTases is independent of the metal ion cofactor, and these enzymes do not possess the metal ion-binding DxD motif. However, in Agl16 we could identify a DxD motif; whether this motif is indeed important for the catalysis by binding a cofactor is still unclear.

BLAST analysis against known protein structures of GT-Bfolded GTases revealed a crystal structure (residues 172 to 338) of an archaeal GTase from *Archaeoglobus fulgidus* (W. Zhou, F. Forouhar, K. Conover, R. Xiao, T. B. Acton, G. T. Montelione, L. Tong, J. F. Hunt, PDB no. O30192, 2005) which showed 26% sequence identity to Agl16. This GTase is homologous to the bacterial mannosyltransferase WbaZ, which catalyzes an α -Man $p(1 \rightarrow 3)$ - β -D-Galp linkage in the O-antigens from *Salmonella enterica* (35) and from *E. coli* K30 (36). Modeling of the Agl16 to the archaeal WbaZ homolog (PDB no. 2f9f) showed the two-domain structure, including the Rossmann-like fold.

The analysis of the genetic localization of *alg16* revealed two genes which might also participate in the *N*-glycosylation pathway by activating the sugar precursor. Directly upstream, the gene *saci0806* encodes a predicted phosphoglucomutase/phosphomannomutase and the gene *saci0802* encodes a predicted 3-hexulose-6-phosphate synthase (Fig. 1). In our proposed model, Saci0802 would activate the sugar by transferring a phosphate from nucleotide triphosphate onto the C₆ atom of the sugar. The phosphoglucomutase/phosphomannomutase is predicted to transfer the phosphate to create sugar-3-phosphate. This phosphorylated sugar would then be nucleotide activated by a yet-unknown enzyme. The nucleotide-activated sugar acts as a donor



FIG 1 Proposed activation and transfer pathway of the last hexose residue of the *N*-glycan in *S. acidocaldarius*. (A) Proposed biosynthesis and transferase pathway of the last hexose. The structure of the unidentified hexose is shown as glucose. *saci0802* encodes a proposed 3-hexulose-6-phosphate synthase; *saci0806* is annotated as a phosphoglucomutase, and *agl16* codes for a soluble glycosyltransferase. Glycan symbols: blue sphere, glucose; green sphere, mannose; orange sphere, sulfoquinovose; blue square, GlcNAc. (B) Physical map of the gene region of *S. acidocaldarius*, where the gene coding for the glycosyltransferase Agl16 is located. The illustrated region encompasses *saci0802* to *saci0808*. The outlined gene encodes the glycosyltransferase Agl16. The genes *saci0802* and *saci0806* are most likely also involved in the activation of the hexose.



FIG 2 Confirmation of the in-frame $\Delta agl16$ mutant. (A) Gene deletion was confirmed by PCR using the outside primers against the flanking regions of *agl16* and template DNA isolated from the background strain MW001, the mutant lacking *agl16*, or the plasmid pSVA1233, used for the homologous recombination incorporating the up- and downstream region of deleted *agl16*. (B) RT-PCR confirmed the deletion of the gene *agl16*, encoding a glycosyltransferase. The cDNA (C) served as a template in PCR amplification using primers against the internal region of either *aglB* or *agl16*. In each case, PCR amplifications were also performed using genomic DNA (G) as a positive control and total RNA (R) as a negative control.

for the GTase Agl16, which either directly transfers the sugar to the premature *N*-glycan or to the dolichyl phosphate. However, the direct transfer to the *N*-glycan is more likely, because all eukaryal GTase transferring sugars onto dolichyl phosphate, like Dpm1 or Alg5, are membrane bound.

Markerless in-frame deletion of agl16. To delete agl16, 800 to 1,000 bp stretches flanking the gene were amplified and fused by PCR. The resulting overlapping fragment was inserted into the plasmid pSAV0407, including the pyrEF genes for selection. The obtained vector was transformed in the background strain S. acidocaldarius MW001 and selected first for the integration and second for the segregation of the plasmid, yielding either the background strain or the in-frame deletion mutant. The deletion of agl16 was first confirmed by PCR amplification of the gene area with upstream forward and downstream reverse primers, resulting in either a 3,125-bp or a 2,090-bp fragment for the background strain MW001 or the $\Delta agl16$ mutant, respectively (Fig. 2). To verify the $\Delta agl16$ mutant, a reverse transcription-PCR (RT-PCR) was performed. RNA of the background strain MW001 and the $\Delta agl16$ strain, grown to mid-exponential growth phase, was isolated and used as a template for the synthesis of single-stranded cDNA. The control PCR used the genomic DNA, cDNA, and RNA as templates, as well as primers against the internal region of either agl16 or aglB, encoding the oligosaccharyl transferase (Fig. 2). The PCR confirmed the lack of genomic DNA in the RNA sample as well as the functional synthesis of the cDNA for the background strain MW001 and for the mutant. PCR with primers against the internal region of agl16 confirmed the absence of agl16 in the deletion mutant strain.

Deletion of *agl16* reduced the molecular weight of the S-layer protein SlaA. As described above, the GTase Agl16 was first identified by screening a variety of GTase deletion mutants for defects in the *N*-glycan biosynthesis. To detect differences in the *N*-glycan size, the S-layer glycoprotein SlaA was used as a reporter protein. Isolated S-layer proteins from the background as well as from the *agl16* deletion strain were separated on an 11% SDS-PAGE gel (Fig. 3). Staining with Coomassie brilliant blue revealed that the samples contained similar amounts of the S-layer glycoprotein. Furthermore, the SlaA from the $\Delta agl16$ deletion mutant migrated faster than the wild-type protein (Fig. 3). This was the first indication that the Agl16 is involved in the *N*-glycosylation pathway. To determine whether the defect in the *N*-glycosylation was indeed caused by the deletion of *agl16* and was not a result of polar effects, the deletion strain was complemented with an empty plasmid (control) or with a plasmid containing the full-length *agl16* gene. The S-layer of the complemented strain ($\Delta agl16 + pSVA1274$) showed no alteration in the protein size, implying the successful restoration of the full *N*-glycan biosynthesis pathway by the expression of the *agl16* gene, excluding any polar effects in the deletion mutant (Fig. 3). Deletion of the *agl16* resulted in a reduced FlaB protein size.

As it had been previously shown that deletion of *agl3*, encoding a UDP-sulfoquinovose synthase, resulted in an altered N-glycan on SlaA and the archaellin FlaB (14), we reasoned that the deletion of agl16 might also lead to a different migration behavior of the FlaB protein in SDS-PAGE. Therefore, cell pellets from the background strain S. acidocaldarius MW001, the $\Delta agl3$ strain (14), the $\Delta agl3$ strain complemented with pSVA1266 (14), the $\Delta agl16$ strain, and the $\Delta agl16$ strain complemented with pSVA1274 were separated by 11% SDS-PAGE and immunoblotted with antibodies raised against FlaB. Both FlaB proteins from the $\Delta agl16$ and from the $\Delta agl3$ deletion strain migrated faster than the wild-type protein (Fig. 4), whereas in the complementation strains, the original FlaB wild-type size was restored. As was shown for the S-layer glycoprotein, FlaB in the $\Delta agl16$ strain was small compared to FlaB in the wild-type and $\Delta agl3$ strains. This difference most likely is due to the fact that only one sugar residue is lost within the N-glycan in the $\Delta agl16$ strain, whereas in the $\Delta agl3$ strain, the N-glycan is reduced to three instead of six sugar residues (14). Although



FIG 3 Effects of the *agl16* deletion on *S. acidocaldarius* S-layer glycoprotein. Equivalent amounts of the S-layer protein SlaA from *S. acidocaldarius* MW001, the $\Delta agl16$ strain, the $\Delta agl16$ strain with control plasmid pSVA1450, and the $\Delta agl16$ strain complemented with pSVA1274 containing *agl16* were separated by 11% SDS-PAGE and Coomassie blue stained.



FIG 4 Glycosylation defects on the archaellin FlaB from *S. acidocaldarius*. Equivalent amounts of cells from *S. acidocaldarius* MW001 (lane 1), the $\Delta agl16$ strain (lane 2), the $\Delta agl16$ strain complemented with pSVA1274 (lane 3), the $\Delta agl3$ strain (lane 4), and the $\Delta agl3$ strain complemented with pSVA1266 (lane 5) were separated by 11% SDS-PAGE and immunoblotted with antibodies raised against FlaB.

equivalent amounts of cells were used, as confirmed by Coomassie staining (data not shown), the immunoblot signal was stronger in the wild-type and complemented strains, implying a higher FlaB protein concentration present in these cells.

The SlaA N-glycan composition is changed in the $\Delta agl16$ deletion mutant. In order to verify that the observed changes in the size of the S-layer glycoprotein and the archaellin FlaB (Fig. 3 and 4) are indeed caused by the lack of one sugar residue of the N-glycan within the $\Delta agl16$ mutant, the N-glycan composition was analyzed. Thus, glycopeptides of the trypsin-digested S-layer proteins from wild-type S. acidocaldarius MW001 and the $\Delta agl16$ mutant were analyzed by nano-LC-ES-MS/MS. The glycopeptides of the wild-type S. acidocaldarius were previously shown to carry a hexasaccharide with the composition Glc1Man2GlcNAc2QuiS, with some glycoforms being truncated at presumed intermediate steps of biosynthesis and with the trisaccharide Man₁GlcNAc₂ being the most dominant truncated structure (7). As shown in Fig. 5, for the tryptic peptides T26 and T28 (for designations of the tryptic peptides, see reference 7), the intact hexasaccharide is no longer present in the $\Delta agl16$ mutant, as the corresponding glycopeptides

of m/z [1561.5]³⁺ for the T24 and m/z [1151.6]⁴⁺ for the T26 were missing (7). Instead, the largest molecular ions were 1507.37³⁺ for T24 glycopeptides and 1111.59⁴⁺ for T26 glycopeptides, which correspond to pentasaccharide-linked peptides. These data confirmed that indeed a single hexose is missing in the *N*-glycan composition in the $\Delta agl16$ mutant. The identity of the hexose is either mannose or glucose, as both are present as terminal sugars. Since mannose and glucose have the same mass, the identity of the missing hexose cannot be unequivocally assigned from the MS data. Therefore, the *N*-glycan of the $\Delta agl16$ mutant has a composition of either Glc₁Man₁GlcNAc₂QuiS or Man₂GlcNAc₂QuiS, as shown in Fig. 5.

HPLC analyses lead to identification of the missing hexose as glucose. To identify the nature of the missing hexose in the N-glycan of the $\Delta agl16$ deletion mutant, the N-glycans of isolated SlaA from the background strain MW001 and the $\Delta agl16$ mutant strain were hydrolyzed with TFA, and the supernatant containing the hydrolyzed monosaccharides were analyzed by HPLC. Based on the composition of the fully assembled N-glycan tree, the Glc/Man ratio of the N-glycan from the MW001 background strain should be 1:2 (7, 25). However, HPLC analyses revealed a higher Glc ratio in the background strain than expected. The increase of the Glc peak within the HPLC analyses is due to the presence of GlcN derived from GlcNAc hydrolysis, which runs at the same retention time in the HPLC analysis. Furthermore, in analyses of the N-glycan composition of the $\Delta agl3$ strain, which includes only Man₁GlcNAc₂ and lacks the terminal glucose residue, a peak with the same retention time as Glc was apparent. Nevertheless, as the GlcNAc and QuiS concentrations are equal in MW001 and the $\Delta agl16$ strain, the changes in the Glc/Man concentration in the mutant and background strain made identifying the missing hexose possible. HPLC analyses of three independent isolated S-layers that were TFA hydrolyzed revealed average Glc/ Man ratios of 9.2:1 in the MW001 background strain and 4.4:1 in the $\Delta agl16$ deletion strain, showing that the Glc concentration in the $\Delta agl16$ deletion strain is 2-fold reduced (Table 4). Based on the changes in the Glc concentration, it is proposed that Agl16 is a glycosyltransferases transferring the terminal Glc residue to the N-glycan.



FIG 5 Detailed analysis of the *S. acidocaldarius* $\Delta agl16$ mutant background S-layer glycopeptide. The total ion chromatograms of the glycan profiles for the tryptic peptides T24 (A) and T26 (B) are shown. The annotated peaks show that the truncated Hex₂QuiS₁HexNAc₂ glycoform is present on the observed glycopeptides, whereas the full glycan is missing. For reference, a cartoon depicting the full glycan is provided for each spectrum. Glycan symbols: blue sphere, glucose; green sphere, mannose; orange sphere, sulfoquinovose; blue square, GlcNAc.

	Retention time (min)		Area (mV*s)		Concn (µM)		Slope for calibration	Ratio
Sample	Glucose	Mannose	Glucose	Mannose	Glucose	Mannose	$(mV \cdot s \cdot \mu M^{-1})$	(glucose:mannose)
Standards								
Glucose	14.31		4,775		10,000		0.48	
Mannose		15.17		4,234		10,000	0.42	
Background strains								
MW001 S1	14.31	15.19	645	68.6	1,351	162.0		8.3:1
MW001 S2	14.33	15.18	646	69.3	1,353	163.7		8.3:1
MW001 S3	14.32	15.20	537	43	1,125	101.6		11.1:1
Mutants								
$\Delta agl16$ S1	14.34	15.18	97.6	21.4	204	50.5		4.0:1
$\Delta agl16$ S2	14.35	15.16	51.16	9.52	107	22.5		4.8:1
$\Delta agl16$ S3	14.33	15.17	112.31	23.51	235	55.5		4.2:1

TABLE 4 HPLC analyses of TFA treatzed isolated S-layer from the background strain MW001 and the $\Delta agl16$ deletion mutant

Deletion of the agl16 showed a drastic reduction of motility. We reasoned that the reduced FlaB protein concentration in the deletion mutants might be caused by a diminished stability or impaired assembly of the archaellum by the altered N-glycan composition. To test the functionality of the archaellum in the deletion strains, a motility assay was performed as previous described (33). Aliquots of the nonmotile S. acidocaldarius $\Delta aapF \Delta flaH$ deletion strain (33) and the hypermotile S. acidocaldarius $\Delta aapF$ strain (37) were used as control strains. The S. acidocaldarius background strain MW001, the $\Delta agl3$ strain, the S. acidocaldarius $\Delta agl16$ strain, and the complemented strain were dropped (or dripped) onto semisolid Gelrite plates containing either 0 mM NaCl or 300 mM NaCl and incubated for 4 and 9 days. In contrast to the hypermotile S. acidocaldarius $\Delta aapF$ strain, for which swimming could be detected after 4 days of incubation, the nonmotile S. acidocaldarius $\Delta aapF \Delta flaH$ deletion strain showed no motility even after 9 days of incubation (Fig. 6). The background strain MW001 showed strong motility after 9 days of incubation, whereas the $\Delta agl3$ and $\Delta agl16$ deletion strains exhibited no and drastically reduced swimming abilities, respectively. The swimming area appeared fuzzy at the colony border for the $\Delta agl16$ strain, showing a reduced but not totally impaired swimming ability, whereas the ability to swim in the $\Delta agl3$ strain seemed totally abolished. The complementation of $\Delta agl16$ restored motility, as the colony showed a swimming radius similar to that of the background strain MW001 (Fig. 6). The incubation under elevated salt conditions (300 mM NaCl) abolished motility, as even the hypermotile S. acidocaldarius $\Delta aapF$ strain did not show any swimming behavior. However, the deletion strains showed a drastic reduction of growth at high salinities, compared to the background strain or the hyper- or nonmotile strains (Fig. 6). The ability to grow at high salinity is dependent on the glycan size. The $\Delta agl3$ deletion strain showed no obvious growth after 4 days of incubation, whereas after 9 days of incubation, only a slight increase in cell density could be detected. After 4 days of incubation, the growth of the $\Delta agl16$ deletion strain was reduced compared to the background strain MW001; however, the defect was not so striking as in the $\Delta agl3$ deletion strain. Complementation of the $\Delta agl16$ recovered the growth defect, as the complementation $\Delta agl16$ + pSVA1274 strain showed the same colony density and size as the background strain MW001.



FIG 6 Effects on the motility from *S. acidocaldarius* by interfering with the *N*-glycan assembly. Equivalent amounts of cells (5 to 6 μ l) from *S. acidocaldarius* $\Delta aapF \Delta flaH$ (lane 1), *S. acidocaldarius* $\Delta aapF$ (lane 2), the background strain *S. acidocaldarius* MW001(lane 3), *S. acidocaldarius* $\Delta agl3$ (lane 4), *S. acidocaldarius* $\Delta agl16$, and *S. acidocaldarius* $\Delta agl16$ complemented with pSVA1274 (lane 6), were dripped onto a 0.15% Gelrite plate and incubated for 4 and 9 days at 75°C. The panels on the right show the enlarged colony border and the motility zone (taken from the left panel: 0 mM NaCl, 9-day incubation), as well as the *N*-glycan composition of MW001 and the $\Delta agl16$ and $\Delta agl3$ mutants. The ability to swim corresponds to the *N*-glycan size.



FIG 7 TEM analysis of negatively stained cells of MW001 (A), the $\Delta agl3$ mutant (B), and the $\Delta agl16$ mutant (C) showed the lack of archaella on each of the *agl* deletion mutants. The MW001 background strain exhibits Aap pili (8 to 10 nm; white arrow) as well as the archaella (10 to 14 nm, black arrow); in the *agl* deletion mutants, only the Aap pili could be detected. Bars, 200 nm (A) and 100 nm (B and C).

The cells of the agl3 and agl16 deletion mutants appeared nonarchaeallated. To confirm that the impaired or reduced motility of the agl mutants was caused by unstable archaella due to the N-glycan defects, the presence of cell appendages in each mutant was analyzed by transmission electron microscopy (TEM). In contrast to the MW001 background strain, which exhibits Aap pili with a diameter of 8 to 10 nm and archaella with a diameter of 10 to 14 nm (Fig. 7A), no archaella could be detected in either agl deletion mutant (Fig. 7B and C). The analyses were repeated with cell cultures that had undergone tryptone starvation to induce archaellum production, as previously described (33). Even under inducing conditions, archaella could not be detected in either mutant by TEM. This confirmed that the archaellum is unstable or cannot be assembled on the cell surface of the agl deletion mutants. Although the agl16 deletion mutant showed little motility on semisolid Gelrite plates (Fig. 6), this analysis underlines the importance of the terminal hexose for stable archaellum assembly. However, in contrast to the archaella, the pilins were present in each of the mutants. Also, the Aap pili are predicted to be Nglycosylated but with a lower ratio than the archaellins; moreover, the pili seem to be much more stable even under harsh detergent treatment (A. Henche and S.-V. Albers, personal communication). The EM images gave further indication that the S-layer stability or assembly is affected in these agl mutants. In contrast to that of the MW001 background strain, the S-layer of the mutants showed defects causing the cytoplasmic membrane to bulge out (Fig. 7). This instable S-layer might not affect the growth under normal growth condition but is more obvious at elevated salt concentrations, in which the stability of the Slayer is more affected.

The alteration of the *N*-glycan resulted in growth retardation upon salt stress. As shown in the motility assay defects in the N-glycan biosynthesis have strong effects on the growth under elevated salinities. Previously we reported the significant reduction of the growth rate for the $\Delta agl3$ strain under salt stress, in which the biosynthesis of the sulfoquinovose incorporated in the N-glycan was interrupted (14). The S-layer as the sole surface envelope in S. acidocaldarius has to stabilize the cell surface and protect it from external conditions. As defects in the biogenesis of these glycoproteins presumably have an effect on the interaction and the stability of the protective S-layer, we measured the growth rate for the background and the $\Delta agl16$ strains. Under standard growth conditions (Brock medium, 0.1% NZ-amine and 0.1% dextrin, 0 mM NaCl, 79°C and pH 3), no significant changes in the doubling times of the mutant ($T_d = 8.59 \pm 0.23$ h) compared to the background strain MW001 ($T_d = 8.72 \pm 0.26$ h) were observed (Fig. 8). The effect of the reduced glycan structure became evident when the strains were grown at higher salinities. At 200 mM NaCl, the doubling time of the $\Delta agl16$ strain was increased $(T_d = 14.82 \pm 1.22 \text{ h})$, whereas the doubling time of the background strain ($T_d = 8.62 \pm 0.55$ h) was not altered compared to that under standard growth conditions. At 300 mM NaCl, the growth of the background strain was only slightly affected ($T_d =$ 13.01 \pm 0.37 h), while the deletion mutant was profoundly affected. The doubling time of the mutant increased to twice $(T_d =$ 24.57 \pm 1.75 h) the doubling time of the background strain. In comparison to the $\Delta agl3$ strain, for which the doubling time was $63.9 \pm 2 h$ (14) at this salt concentration, the effect of one missing sugar in the S-layer N-glycan had a much lower impact than the deletion of three sugars from the glycan tree. However, at 400 mM NaCl, only the background strain, MW001, was capable of sustaining growth after a long lag phase ($T_d = 34.84 \text{ h} \pm 6.48 \text{ h}$), whereas the mutant showed no obvious growth.



FIG 8 Response to salt stress in the $\Delta agl16$ strain. S. acidocaldarius MW001 (black line) or $\Delta agl16$ (gray line) were grown in Brock medium at different salt concentrations (0 to 400 mM NaCl). Growth was measured as the optical density at 600 nm (OD₆₀₀). Shown are values obtained from an experiment done in triplicate; repeats generated a similar result.

DISCUSSION

Various surface-exposed proteins from S. acidocaldarius have been shown to be modified by the attachment of glycan, e.g., Slayer protein (7), cytochrome $b_{558}/566$ (23, 25), archaellins (14), or sugar-binding proteins (24). Most of them are shown to be modified via the N-glycosylation process. Recently, we identified the first enzyme participating in the N-glycosylation process in S. acidocaldarius. Agl3, a UDP-sulfoquinovose synthase, creates the nucleotide-activated sugar precursor UDP-sulfoquinovose, which is assembled to form a branch of the N-glycan (14). However, additional enzymes, in particular glycosyltransferases (GTases), necessary for the assembly of the full-length N-linked glycan are still unidentified in S. acidocaldarius (38). Unlike the agl gene cluster within Haloferax volcanii, in which the oligosaccharyltransferase AglB and other enzymes involved in the N-glycosylation process are found (39), genes coding for predicted GTases are scattered all over the S. acidocaldarius genome and not located near the aglB gene (Table 3). This scattered distribution of GTase seems common within thermophilic organisms (19), making the identification of N-GTases in thermophilic Archaea more difficult. Various deletion studies of genes coding for predicted GTases within the genome of S. acidocaldarius so far have identified only one GTase involved in the biosynthesis of the N-glycan of the S-layer glycoprotein. In the present study, we report this first identified glycosyltransferase involved in the N-glycosylation in a crenarchaeon. The markerless in-frame deletion of agl16 in S. acidocaldarius resulted in a reduced molecular size of the large S-layer protein SlaA and the archaellin FlaB. The slight reduction of the molecular size of these proteins implied that one or more sugar residues are absent in the N-glycan structure. Nano-LC-MS/MS analysis confirmed the presence of a pentasaccharide lacking one terminal hexose residue. HPLC analyses of the sugar composition of isolated S-layer proteins from the $\Delta agl16$ and the background strains revealed a reduced Glc concentration in the agl16 mutant, which indicate that Agl16 is important for the transfers of the terminal glucose residue onto the premature N-glycan. The recently developed expression system in S. acidocaldarius (28) enables us to express and purify Agl16 homologously. The characterization of Agl16 will give us new insight in the glycosyltransferase reaction, e.g., which donor and acceptor are used. For instance Agl16 might use soluble nucleotide-activated (UPD or GDP) glucose precursors or dolichyl phosphate-linked glucose residues as donors. Additionally, an activity assay of Agl16 would also reveal the preferred acceptor, whether it is dolichyl phosphate or directly the premature N-glycan tree linked to dolichyl phosphate. In Haloferax volcanii, the last assembly step of the N-glycan is mediated by AglS, which transfers the terminal mannose of the pentasaccharide (40). In contrast to the soluble Agl16, AglS is a membrane-bound GTase, which transfers the last mannose residue from a dolichyl phosphate-activated mannose onto the N-glycan tree. This assembly step is similar to the last eukaryal N-glycan biosynthesis step, in which the membrane-bound GTase uses dolichyl phosphate-linked mannose or glucose residues as the sugar donor (41, 42). All other soluble eukaryal GTases use nucleotide-activated sugar residues in the first N-glycan biosynthesis steps. Therefore, it is most likely that Agl16 also uses a nucleotideactivated sugar precursor as the sugar donor.

The deletion of *agl16* in *S. acidocaldarius* causes significant effects on the phenotype, as the presence of only the partial *N*-glycan

structure on the S-layer protein and the archaellins resulted in a reduced growth rate at elevated salt concentration as well as impaired function of the archaellum. A similar effect of a truncated N-glycan was also observed in other Archaea. The observation that underglycosylation affects archaellum filament assembly was first made in Methanococcus maripaludis, where deletion of MMP0350, encoding a putative acetyltransferase, was proposed to be responsible for the acetylation of the first two N-glycan sugar residues (43). Also, in Methanococcus voltae, deletion of aglA, involved in the final biosynthetic step of the trisaccharide, had a similar effect on the stability of the archaella, as the $\Delta aglA$ strain exhibited only few and shortened archaella (22). The finding that archaellins need to be modified with at least a subset of the N-glycan sugar to assemble and function has recently also been shown in H. volcanii (17). Defects in the N-glycosylation introduced by agl gene deletions as well as the replacement of the asparagines within the Nglycosylation sites in the archaellins resulted in the loss of motility and archaellum assembly in H. volcanii (17). However, in contrast to H. volcanii and M. voltae in which a trisaccharide is the minimal N-glycan size to allow motility (17, 22, 44), S. acidocaldarius displays a strong dependency on the fully assembled tribranched hexasaccharide, as even one missing sugar residue in the N-glycan almost completely abolished motility. A correlation of salinity and growth was also shown in H. volcanii, in which the deletion of aglD resulted in a reduction of the growth rate in media with high salinities (3.7 M or 4.8 M NaCl), whereas under standard growth conditions (1.7 M NaCl), no change in growth was observed (10).

Beside *agl16*, the genes *saci0802* and *saci0806*, lying upstream of *agl16*, are predicted to be involved in the *N*-glycan biosynthesis step. Mutants with deletions of these genes will help determine whether these gene products, as it is proposed in Fig. 1A, activate the last hexose residue before it is transferred by Agl16. Additionally, analyses of gene expression might give insight into a coordinated regulation of these first genes involved in the *N*-glycosylation process.

In conclusion, this study identified the first crenarchaeal GTase, Agl16, catalyzing the last biosynthesis step of the *N*-glycan in *S. acidocaldarius*. The deletion of *agl16* demonstrated the importance of the fully assembled *N*-linked hexasaccharide for the growth and proper assembly and function of the archaellum of *S. acidocaldarius*. The comparison to the $\Delta agl16$ mutant and the previous reported $\Delta agl3$ mutant demonstrated that motility is strongly dependent on *N*-glycan size.

This study has now extended the field of *N*-glycosylation research to crenarchaeotes and contributes to the increasing data available on archaeal *N*-glycosylation and enzymes responsible for biosynthesis and assembly of *N*-glycans. Further studies will be undertaken to uncover the complete *N*-glycosylation pathway in *S. acidocaldarius.*

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