



# Lipid Anchoring of Archaeosortase Substrates and Midcell Growth in Haloarchaea

® Mohd Farid Abdul-Halim, a Stefan Schulze, a Anthony DiLucido, a Friedhelm Pfeiffer, b Alexandre Wilson Bisson Filho, c & Mechthild Pohlschroder Bernard Wilson Bisson Filho, c

ABSTRACT The archaeal cytoplasmic membrane provides an anchor for many surface proteins. Recently, a novel membrane anchoring mechanism involving a peptidase, archaeosortase A (ArtA), and C-terminal lipid attachment of surface proteins was identified in the model archaeon Haloferax volcanii. ArtA is required for optimal cell growth and morphogenesis, and the S-layer glycoprotein (SLG), the sole component of the H. volcanii cell wall, is one of the targets for this anchoring mechanism. However, how exactly ArtA function and regulation control cell growth and morphogenesis is still elusive. Here, we report that archaeal homologs to the bacterial phosphatidylserine synthase (PssA) and phosphatidylserine decarboxylase (PssD) are involved in ArtA-dependent protein maturation. Haloferax volcanii strains lacking either HvPssA or HvPssD exhibited motility, growth, and morphological phenotypes similar to those of an  $\Delta artA$  mutant. Moreover, we showed a loss of covalent lipid attachment to SLG in the ΔhvpssA mutant and that proteolytic cleavage of the ArtA substrate HVO\_0405 was blocked in the ΔhvpssA and ΔhvpssD mutant strains. Strikingly, ArtA, HvPssA, and HvPssD green fluorescent protein (GFP) fusions colocalized to the midcell position of H. volcanii cells, strongly supporting that they are involved in the same pathway. Finally, we have shown that the SLG is also recruited to the midcell before being secreted and lipid anchored at the cell outer surface. Collectively, our data suggest that haloarchaea use the midcell as the main surface processing hot spot for cell elongation, division, and shape deter-

**IMPORTANCE** The subcellular organization of biochemical processes in space and time is still one of the most mysterious topics in archaeal cell biology. Despite the fact that haloarchaea largely rely on covalent lipid anchoring to coat the cell envelope, little is known about how cells coordinate *de novo* synthesis and about the insertion of this proteinaceous layer throughout the cell cycle. Here, we report the identification of two novel contributors to ArtA-dependent lipid-mediated protein anchoring to the cell surface, HvPssA and HvPssD. ArtA, HvPssA, and HvPssD, as well as SLG, showed midcell localization during growth and cytokinesis, indicating that haloarchaeal cells confine phospholipid processing in order to promote midcell elongation. Our findings have important implications for the biogenesis of the cell surface.

**KEYWORDS** archaea, *Haloferax volcanii*, S-layer, archaeosortase, cell division, cell elongation, cell shape, cell surface, haloarchaea, lipid anchoring

Microbial cell surface proteins play critical roles in many important biological processes, including bioenergetics, mediation of intercellular communication, nutrient uptake, surface adhesion, and motility. Cell surface proteins also play impor-

Citation Abdul-Halim MF, Schulze S, DiLucido A, Pfeiffer F, Bisson Filho AW, Pohlschroder M. 2020. Lipid anchoring of archaeosortase substrates and midcell growth in haloarchaea. mBio 11:e00349-20. https://doi.org/10.1128/mBio.00349-20.

**Invited Editor** Sonja-Verena Albers, University of Freiburg

Editor M. Stephen Trent, University of Georgia
Copyright © 2020 Abdul-Halim et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0
International license.

Address correspondence to Alexandre Wilson Bisson Filho, bisson@brandeis.edu, or Mechthild Pohlschroder, pohlschr@sas.upenn.edu.

This article is a direct contribution from Mechthild Pohlschroder, a Fellow of the American Academy of Microbiology, who arranged for and secured reviews by Rachel Whitaker, University of Illinois at Urbana-Champaign, and Parkson Chong, Temple Libitority.

Received 13 February 2020 Accepted 18 February 2020 Published 24 March 2020

<sup>&</sup>lt;sup>a</sup>Department of Biology, University of Pennsylvania, Philadelphia, Pennsylvania, USA

<sup>&</sup>lt;sup>b</sup>Computational Biology Group, Max Planck Institute of Biochemistry, Martinsried, Germany

Department of Biology, Rosenstiel Basic Medical Science Research Center, Brandeis University, Waltham, Massachusetts, USA

mBio<sup>®</sup>

tant roles in cell elongation and shape maintenance, but how this is achieved in archaea is not well understood (1).

The structural organization of cellular surfaces is one important readout of how cells coordinate growth, morphogenesis, and division. In both bacteria and eukaryotes, a multitude of growth modes have been characterized, with cells inserting new envelope material almost all along the cell surface (2), bipolarly (3), and unipolarly (4), and in some cases, different modes can be interchangeable (5, 6). In the case of archaea, which lack a peptidoglycan cell wall, glycosylated S-layer and other proteins are commonly the sole components of the cell envelope (7, 8), where they typically show a twodimensional (2D) crystal-like arrangement. This poses an interesting problem for archaeal surface protein organization, and currently, there are no data about the mechanisms of archaeal cell elongation control (9).

While many proteins are anchored to the cell surface via transmembrane (TM) domain insertion into the membrane, some are anchored through covalent N-terminal attachment of a lipid moiety (8). Recently, a novel mechanism was discovered whereby proteins are anchored to the membrane through a lipid moiety covalently attached to a processed C terminus. In archaeal cells, processing and lipid modification of these C-terminal anchored proteins are mediated by enzymes known as archaeosortases, with archaeosortase A (ArtA) of the model archaeon Haloferax volcanii being the most studied example (10-13). Proteins recognized and processed by H. volcanii ArtA contain a distinct C-terminal tripartite structure consisting of a conserved PGF motif, followed by a hydrophobic domain and then a stretch of positively charged residues. Molecular biological and biochemical analyses determined that ArtA does indeed process a diverse set of proteins, including both Tat and Sec substrates, that have been shown to play roles in motility and mating (10, 12). Most notably, this includes the S-layer glycoprotein (SLG), which is the sole component of the H. volcanii cell wall.

A previous in silico study by Haft and coworkers noted that in Methanosarcina acetivorans C2A, Methanosarcina mazei Gö1, as well as several other methanogens, the artA gene is located next to the gene that encodes an archaeal homolog of bacterial phosphatidylserine synthase (PssA) (14). Based on the degree of sequence similarity and its substrate specificity, the archaeal PssA homolog belongs to the PssA subclass II, similar to PssA found in Gram-positive bacteria such as Bacillus subtilis, as opposed to the PssA of Gram-negative bacteria, such as Escherichia coli, which belongs to subclass I (15). Work in B. subtilis has elucidated most of the biochemistry involved in the reaction catalyzed by PssA, which involves the transfer of a diacylglycerol moiety from a CDP-phosphatidyl lipid to L-serine to make phosphatidylserine (16, 17). Phosphatidylserine can subsequently be decarboxylated to phosphatidylethanolamine by the enzyme phosphatidylserine decarboxylase (PssD), which has been characterized from Sinorhizobium meliloti and B. subtilis (17, 18). However, unlike bacterial PssA, in vitro study of the archaeal PssA homolog from Methanothermobacter thermautotrophicus (MTH\_1027) revealed that this protein catalyzes the transfer of the archaetidic acid moiety of CDP-archaeol onto the hydroxyl group of L-serine to form the polar lipid archaetidylserine (CDP-2,3-di-O-geranylgeranyl-sn-glycerol: L-serine O-archaetidyltransferase) (15). Mirroring the phosphatidylethanolamine biosynthesis reaction in bacteria, it was postulated that archaetidylserine could also undergo decarboxylation to archaetidylethanolamine by an archaeal PssD homolog, a putative archaetidylserine decarboxylase.

Distant homologs to PssA and PssD are encoded in the H. volcanii genome, which we refer to as HvPssA (HVO\_1143) and HvPssD (HVO\_0146), respectively. In this study, we show that HvPssA and HvPssD are involved in ArtA-dependent C-terminal protein maturation, which involves proteolytic cleavage and lipid anchoring. An interplay between ArtA, HvPssA, and HvPssD is further supported by their colocalization at midcell. These analyses reveal, to the best of our knowledge for the first time, that cell elongation happens from the midcell in archaea.

## Methanosarcina acetivorans C2A

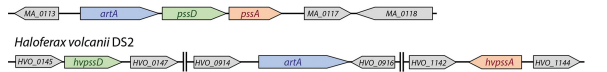


FIG 1 Schematic representation of artA, pssA, and pssD distribution across Euryarchaeota. Shown are M. acetivorans (top) and H. volcanii (bottom) genomic organization of artA and genes encoding homologs to PssA (HvPssA) and PssD (HvPssD). The M. acetivorans artA, pssA, and pssD gene lengths are 834 bp, 627 bp, and 744 bp, respectively. The H. volcanii artA, hvpssA, and hvpssD gene lengths are 912 bp, 672 bp, and 606 bp, respectively.

#### **RESULTS**

Synteny of artA, pssA, and pssD genes in Methanosarcina strains. Based on the juxtaposition of the genes encoding the archaeosortase (artA) and the putative membrane lipid biosynthesis archaetidylserine synthase (pssA) in Methanosarcina spp. (14), we hypothesized that homologs of the H. volcanii PssA (HvPssA) might be involved in the lipidation of ArtA substrates.

Upon further in silico analyses, we were intrigued that in several Methanosarcina species, the artA and pssA genomic region flanked a homolog of the pssD gene (Fig. 1; see also Fig. S1 in the supplemental material), probably encoding archaetidylserine decarboxylase, suggesting that they could be involved in consecutive steps of a proposed archaetidylethanolamine biosynthetic pathway. We decided to extend our studies of ArtA-dependent protein processing in H. volcanii toward the orthologs HvPssA (HVO\_1143) and HvPssD (HVO\_0146), even though the genes are not clustered with ArtA (HVO\_0915) in this species (Fig. 1). The three genes are unclustered, not only in Haloferax spp. but also in more than 70 haloarchaeal genomes.

It should be noted that while the Methanosarcina spp. and H. volcanii PssA and PssD homologs share a significant similarity, these proteins have only 30 to 35% sequence identity to the in vitro-characterized enzymes from Methanobacterium thermautotrophicus, S. meliloti, and B. subtilis (15, 17, 18). Thus, it is possible that the H. volcanii proteins may act on variants of the substrates processed by these experimentally characterized homologs.

H. volcanii  $\Delta hvpssA$  and  $\Delta hvpssD$  mutant cells exhibit growth, morphology, and motility phenotypes similar to those of the AartA mutant strain. In order to determine whether HvPssA and/or HvPssD are involved in the archaeosortasedependent processing pathway, we generated H. volcanii hvpssA and hvpssD deletion mutants (Fig. S2) using the pop-in/pop-out method (19).

We had previously shown that the H. volcanii \( \Delta artA \) mutant strain exhibits various severe phenotypic defects (e.g., poor growth, atypical morphology, and impaired motility), perhaps due, at least in part, to defective processing of the SLG, an ArtA substrate (10). S-layer disruption may affect cell stability and may interfere with morphology. Also, as archaea do not have a cell wall, the S-layer may take over the function as a stator for the archaellar motor. The latter is supported by the fact that ArlF and ArlG (formerly referred to as FlaF and FlaG, respectively) were shown to interact with the S-layer glycoprotein (20, 21). We thus subjected the  $\Delta hvpssA$  and  $\Delta hvpssD$ mutant strains to analysis for these physiological responses. Both deletion strains exhibit a growth defect similar to that of the ΔartA mutant strain compared to the H53 parent strain used in these studies (Fig. 2A). For all genes, normal growth is rescued by complementation, expressing the deleted gene in trans from a plasmid. Moreover, the  $\Delta hvpssA$  and  $\Delta hvpssD$  mutant strains are partially impaired in motility; the defect, however, is less severe than that in the  $\Delta artA$  mutant. While no halo is observed after 5 days of  $\Delta artA$  mutant incubation at 45°C, a small halo is formed by the  $\Delta hvpssA$  and ΔhvpssD mutant strains (Fig. 2B).

Light microscopic examination of the parental strain at mid-exponential-growth stage shows predominantly disk-shaped cells, while the  $\Delta artA$  mutant cells exhibit a

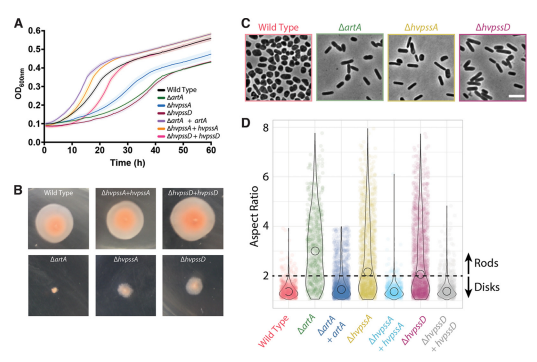


FIG 2 Absence of HvPssA or HvPssD leads to defects in growth, cell morphology, and motility. (A) Wild-type (strain H53) and  $\Delta artA$ ,  $\Delta hvpssA$ , and  $\Delta hvpssD$  mutant cells were grown with shaking in 96-well plates with a total volume of 200  $\mu$ l of liquid semidefined CA medium, and the growth of six biological replicates was monitored at the  $OD_{600}$ , with recordings taken every 30 min. For complementation analysis, artA, hvpssA, or hvpssD was expressed from pTA963 under the tryptophan-inducible p.tna promoter. The wild-type and deletion strains were transformed with an empty pTA963 plasmid as a control. (B) The wild-type (strain H53) and artA, hvpssA, or hvpssD deletion and complementation strains from individual colony on solid agar plates were individually stab inoculated with a toothpick into semisolid 0.35% agar in CA medium supplemented with tryptophan, followed by incubation at 45°C. (C) Phase-contrast images were taken from wild-type and mutant cells during mid-exponential-growth phase (OD<sub>600</sub>, 0.3) and immobilized under 0.5% agarose pads. (D) Violin distributions of aspect ratio measurements from single cells. Rodlike (aspect ratio, >2) cells were prevalent to disklike (aspect ratio, <2) cells in the mutant strains compared to the wild type. Biological replicates were collected on three different days, and data were analyzed from >1,000 cells under each condition by automated image segmentation. Scale bars =  $5 \mu m$ .

predominantly rod-shaped phenotype (Fig. 2C). Again, the  $\Delta hvpssA$  and  $\Delta hvpssD$ mutant strains show a similar but less severe phenotype. The vast majority of cells from cultures are rods, but we have observed a few disk-shaped cells in liquid cultures for each of these strains. This phenotype is fully complemented, and disk-shaped cells are observed when HvPssA and HvPssD are expressed in trans in the  $\Delta hvpssA$  and  $\Delta hvpssD$ mutant strains, respectively (Fig. S3A).

To properly quantify the differences between the wild-type and mutant strains, we segmented cell images and measured the aspect ratio from each population, which is the ratio between the longer and shorter axes of the cell. Subsequently, we observed that H. volcanii rods have an aspect ratio above 2, while disks predominantly fall under this cutoff value (Fig. S3B). Using this arbitrary threshold, we automatedly determined the proportions of rods from wild-type and mutant cells (Fig. 2D and S3C), confirming our initial observation that the  $\Delta artA$ ,  $\Delta pssA$ , and  $\Delta pssD$  mutant populations are significantly enriched in rod-shaped cells compared to the wild type.

Thus,  $\Delta hvpssA$  and  $\Delta hvpssD$  mutant strains exhibit phenotypes similar to that of the ΔartA mutant strain for three independent physiological effects, supporting the hypothesis that the encoded proteins are involved in the ArtA-dependent processing pathway.

HvPssA and HvPssD are required for SLG lipid modification. To determine whether the drastic cell morphology transitions and the phenotypic similarities between the  $\Delta hvpssA$ ,  $\Delta hvpssD$ , and  $\Delta artA$  mutants are due to the inhibition of the covalent lipid modification of ArtA substrates, we investigated the lipidation of the SLG

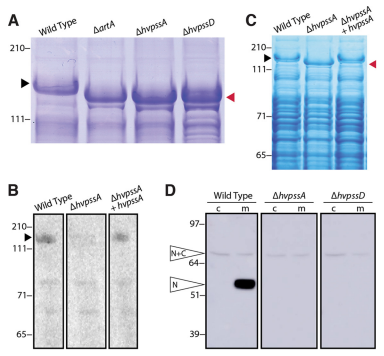


FIG 3 HvPssA and HvPssD are critical for HVO\_0405 C-terminal processing and SLG lipidation. (A) Coomassie-stained LDS-PAGE gel of cell extracts from H. volcanii H53 (wild-type [WT]) and \( \Delta artA, \) ΔhvpssA, and ΔhvpssD mutant strains. The ΔartA, ΔhvpssA, and ΔhvpssD mutant SLG (red arrowhead) exhibited a mobility shift compared to the WT SLG (black arrowhead). (B) Fluorography of protein extracts isolated from H53 (WT), ΔhvpssA mutant, and hvpssA complementation (ΔhvpssA + hvpssA) cells grown in the presence of 1 µCi/ml [14C]mevalonic acid. Significant labeling of SLG (black arrowhead) is only detected in the WT and hvpssA complementation (ΔhvpssA + hvpssA) extracts. (C) Coomassie staining of the gel used for fluorography. The SLG mobility shift in the  $\Delta hvpssA$  mutant (red arrowhead) is reverted upon hvpssA expression in trans. (D) Western blot analysis of cytoplasmic (c) and membrane (m) fractions of H53 (WT), ΔhypssA mutant, and ΔhypssD mutant strains expressing, in trans, HVO 0405-6×His. The N-terminal domain of HVO\_0405 was detected using anti-HVO\_0405-N-term antibodies. Hvo\_0405 not processed by ArtA and the N-terminal HVO\_0405 processed by ArtA are labeled "N+C" and "N," respectively. The C-terminal domain, which carries a His tag, has not been analyzed in this experiment. Numbers indicate molecular mass in kilodaltons.

mediated by ArtA (11). Initially, as an indirect analysis, we examined the effect of hvpssA and hvpssD deletions on SLG electrophoretic mobility in a lithium dodecyl sulfate (LDS)-PAGE gel, as a mobility shift is observed in an artA deletion strain (11). While the similarities of Coomassie-stained band intensities for SLG isolated from the ΔartA, ΔhvpssA, and ΔhvpssD mutant strains and their parent strains indicate a similar SLG abundance, electrophoretic mobility demonstrates similar migration shifts of the SLG isolated from all three deletion strains compared to the SLG from the parent strain (Fig. 3A). These shifts are reverted by complementation with the respective gene in trans (Fig. S4A). To corroborate this observation, we set out to directly measure the lipid modification of SLG in the  $\Delta hvpssA$  mutant strain. These experiments proved that lipid labeling of the SLG with radiolabeled mevalonic acid, an archaeal lipid precursor, is severely impaired in the ΔhvpssA mutant strain compared to the parent strain (Fig. 3B) and that this phenotype can be complemented by expressing HvPssA in trans. This is conclusive evidence for HvPssA being closely or even directly coupled with ArtAdependent protein lipidation. Because the phenotypes of the  $\Delta hvpssD$  mutant strain closely match those of the  $\Delta hvpssA$  mutant strain in all other assays, including the indirect gel-based assay for SLG modification (Fig. 3A to C), it is highly likely that HvPssD is also involved in ArtA-dependent SLG lipidation.

HvPssA and HvPssD are required for proteolytic ArtA-substrate processing. To determine whether HvPssA/HvPssD-dependent lipidation is required for ArtAdependent C-terminal processing, we investigated the C-terminal proteolytic cleavage

of a second ArtA substrate. As a reporter, we used the strain-specific domain fusion protein HVO\_0405, with its centrally located cleavage site, because the cleaved and uncleaved versions of HVO\_0405 can be easily distinguished by LDS-PAGE separation and subsequent immunostaining (12).

Cleavage of C-terminally His-tagged HVO\_0405, expressed in trans, does not occur in the  $\Delta hvpssA$  and  $\Delta hvpssD$  mutant strains. In contrast, this ArtA substrate is cleaved in the corresponding parent strain, as is evident from our Western blot analysis of membrane fractions using antibodies against the N-terminal part of HVO 0405 (Fig. 3D and S4B). Blocking proteolytic cleavage does not lead to an increased amount of the full-length form. The same has been observed in an  $\Delta artA$  mutant (12) and has been attributed to the instability of the domain-fused version of this protein.

Midcell localization of ArtA, HvPssA, and HvPssD promotes cell site-specific lipidation in vivo. Given the dependence of ArtA activity on HvPssA and HvPssD and their importance to cell growth and morphogenesis, we decided to investigate whether ArtA, HvPssA and HvPssD were recruited to perform at specific regions of H. volcanii cells. Strikingly, ArtA, HvPssA, and HvPssD tagged with monomeric superfolder green fluorescent protein (msfGFP) localize at midcell (Fig. 4A). As controls, we also imaged a tagged version of FtsZ1, that has previously been shown to localize to midcell and was speculated to participate in cell division (22), as well as msfGFP not fused to any protein. FtsZ1-msfGFP shows a localization pattern almost identical to those of ArtA, HvPssA, and HvPssD, while the msfGFP protein by itself is not recruited to midcell (Fig. 4A). Furthermore, we confirmed that cells carrying msfGFP-tagged versions of ArtA, HvPssA, and HvPssD present morphological profiles similar to that of the wild type and thus must be functional (Fig. S3D). Finally, time-lapse videos of cells cultivated within microfluidics suggest that ArtA, HvPssA, and HvPssD proteins are recruited to the midcell right after daughter cells are born and persist for most of the cell cycle, including during cytokinesis (Fig. 4B, blue arrowhead, and Movie S1).

Considering that deletions of artA, hvpssA, and hvpssD each drastically perturbed growth (Fig. 2A), induced cells to stay in a rodlike shape (Fig. 2C), and do not seem to play an essential role in cell division, we hypothesize that the lipid anchoring of ArtA substrates specifically at midcell might be important for cell elongation and morphogenesis. Interestingly, we also noticed a correlation between the presence of ArtA, HvPssA, and HvPssD at the midcell and actual cell elongation in the cell population (Fig. 4B, red arrowhead). To investigate this further, we expressed a second copy of the SLG which was tagged with msfGFP, a GFP variant shown to be fluorescent upon Sec-dependent transport in bacteria (23). Interestingly, the SLG-msfGFP fusion accumulated at the midcell site instead of localizing around the cell envelope (Fig. 4C and Movie S2). Thus, even though both versions of SLG, tagged and untagged, are synthesized, they do not seem to coassemble across the S-layer. Additionally, overexpression of the SLG-msfGFP fusion caused severe growth and morphological defects (Fig. 4D). Interestingly, cells rapidly recovered in less than 1 day (approximately 6 generations) following depletion of the SLG-msfGFP, returning to their normal cell shape (Fig. S5). It might be possible that secreted SLG-msfGFP cannot be incorporated into the 2D proteinaceous crystal array. This scenario could explain the observed SLG-msfGFP foci at the poles (Fig. 4C), as an excess of secreted but unassembled SLG protein could aggregate at the pole, where the S-layer array must show discontinuities. However, independent of whether this SLG construct was secreted or not, our results strongly suggest that nascent SLG is targeted specifically to the midcell.

Last, we investigated the morphological transitions in H. volcanii protoplast cells during de novo S-layer synthesis. If the midcell-confined ArtA, HvPssA, and HvPssD are in fact promoting the lipidation of recently secreted SLG molecules at the cell surface, one would be able to observe midcell-localized reshaping during protoplast recovery. As expected, protoplasts generated by the addition of proteinase K either within microfluidics or in bulk cultures adopt a round-like shape (Fig. 4E, left), suggesting that the S-layer might be the structure that ultimately determines the archaeal cell shape. As the protease is washed out with fresh medium, cells rapidly reshape exclusively from

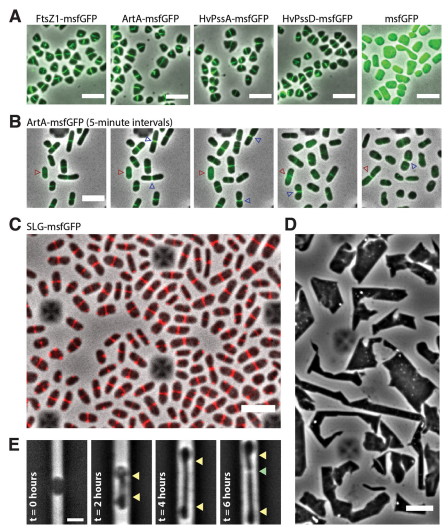


FIG 4 Midcell localization of the lipid-anchoring and processing machinery in H. volcanii. (A) Snapshots of merged phase-contrast (gray) and fluorescein isothiocyanate (FITC; green) channels of cells expressing FtsZ1-msfGFP, ArtA-msfGFP, HvPssA-msfGFP, HvPssD-msfGFP, and soluble msfGFP. Cells were immobilized under 0.5% agarose pads prepared with CA medium. (B) Time-lapse images of cells growing inside a CellASIC microfluidic device. Images of merged phase-contrast (gray) and FITC (green) channels were taken every 5 min for 12 h. Blue arrowheads indicate cell division events, while red arrowheads label one example of a cell elongating only after the arrival of ArtA-msfGFP to the midcell. (C) Snapshot of SLG-msfGFP (red) midcell localization. (D) Phase-contrast images of H. volcanii cells under prolonged overexpression (24 h) of the SLG-msfGFP fusion. (E) H. volcanii cells reshape and elongate preferentially at the midcell during protoplast recovery. Cells were loaded into the microfluidic chamber, and the S-layer was chemically removed by the addition of 1 mg/ml proteinase K and recovered with fresh medium (t = 0 h). Yellow arrowheads indicate the cell area extended until cell division (green arrowhead). Scale bars =  $5 \mu m$ .

the midcell position (Fig. 4E). Altogether, our observations suggest that the midcell area in H. volcanii cells is not only dedicated to cell division but is also a central hub for outbound cell extension and other cellular processes.

## **DISCUSSION**

Our data confirmed the hypothesized involvement of the lipid biosynthesis enzyme homologs HvPssA and HvPssD in the C-terminal posttranslational modifications of ArtA substrates. With respect to physiological effects, the  $\Delta hvpssA$  and  $\Delta hvpssD$  mutants showed similar but slightly less severe phenotypes than that resulting from the deletion of artA (Fig. 2). We furthermore experimentally determined the effects of hvpssA and hvpssD deletions on proteolytic cleavage and lipid labeling (Fig. 3).

Since the gene encoding HVO\_0405 resulted from the fusion of two previously independent genes, this protein provided us with an excellent tool for the analysis of ArtA-related proteolysis in H. volcanii, as the size difference between precursor- and ArtA-processed mature proteins is large enough to be detected using immunostaining (12). This allowed us to clearly demonstrate that ArtA-dependent proteolytic cleavage is blocked when either hvpssA or hvpssD is deleted in H. volcanii. This block could be bypassed by plasmid-based gene complementation. These results strongly suggest that lipidation and proteolysis are intricately connected with proteolytic cleavage only occurring if the modifying lipid and/or HvPssA or HvPssD is present. By sequence homology, H. volcanii HvPssA is predicted to be involved in lipid biosynthesis, specifically, the generation of the polar lipid archaetidylserine from CDP-archaeol. While the functionally characterized homolog (from M. thermautotrophicus) is only distantly related (30% to 35% sequence identity), the confirmed involvement of HvPssD strongly supports the hypothesis that HvPssA and HvPssD generate archaetidylethanolamine.

Interestingly, a recent characterization of a bacterial rhombosortase, a nonhomologous analog of archaeosortase (24), showed direct involvement of a glycerophosphoethanolamine-containing moiety in the process. Analogously, lipid-attached ethanolamine may be directly involved in the membrane anchoring of ArtA substrates. In this scenario, instead of contributing directly to the ArtA-mediated substrate cleavage and/or lipid anchoring, HvPssA and HvPssD catalyze the final steps in a pathway that generates archaetidylethanolamine, a substrate required by this process. This opens the way for another hypothesis regarding the ArtA reaction mechanism. In this scenario, ArtA acts similarly to sortase A in bacteria, wherein ArtA cleaves the substrate through thioesterification, forming a thioester acyl-enzyme intermediate, which is consistent with the identification of Cys-173 as an active-site residue (13). The nucleophilic attack of an amine resolves this intermediate, but instead of a pentaglycine branched lipid II, the reactive amine nucleophile is archaetidylethanolamine (Fig. 5A). Such a mechanism would directly result in a covalently modified protein C terminus. While an archaetidylethanolamine lipid was reported to be absent from Halobacterium salinarum or Haloarcula marismortui (25), it is present in H. volcanii and several other haloarchaea, albeit at various abundances (26).

As lipid analysis does not cope with protein-bound lipids, the low concentration of archaetidylethanolamine is not surprising, even though the SLG is highly abundant and archaetidylethanolamine may be used as its membrane anchor. Nevertheless, a detectable amount of archaetidylethanolamine in the H. volcanii membrane suggests the functional roles of HvPssA and HvPssD, catalyzing the synthesis of archaetidylserine and its decarboxylation to archaetidylethanolamine, respectively. These enzymes perhaps associate with or even form a complex with ArtA, resulting in a majority of the synthesized archaetidylethanolamines to be immediately used to modify the SLG and other ArtA substrates for their membrane anchoring upon C-terminal processing. Thus, only a small amount may be left free in the membrane. The lipid attached to an EDTA-soluble fraction of the SLG has been analyzed by mass spectrometry and was identified as archaetidic acid (27). However, as the lipid has been released from the protein by alkaline hydrolysis, this procedure may have hydrolyzed and thus removed the ethanolamine headgroup.

While investigating the interdependence between ArtA and HvPssA or HvPssD, we observed the recruitment of these proteins to the midcell in *H. volcanii* (Fig. 4A). Considering these data and the observed midcell localization of newly inserted SLG molecules (Fig. 4C), we propose a model for S-layer assembly, lipidation, and growth in haloarchaea (Fig. 5B). First, SLG is recruited to the midcell, where it is transported across the cytoplasmic membrane in a Sec-dependent manner (28). Following secretion, SLG is processed and linked to archaetidylethanolamine by ArtA, requiring HvPssA or HvPssD for archaetidylethanolamine synthesis and/or interaction with ArtA.

There are still key aspects of haloarchaeal growth and shape control that are not addressed by our model. For example, it is still not clear how the deletion of either artA, hvpssA, or hvpssD generates a rod-shaped cell population (Fig. 2C). Furthermore,

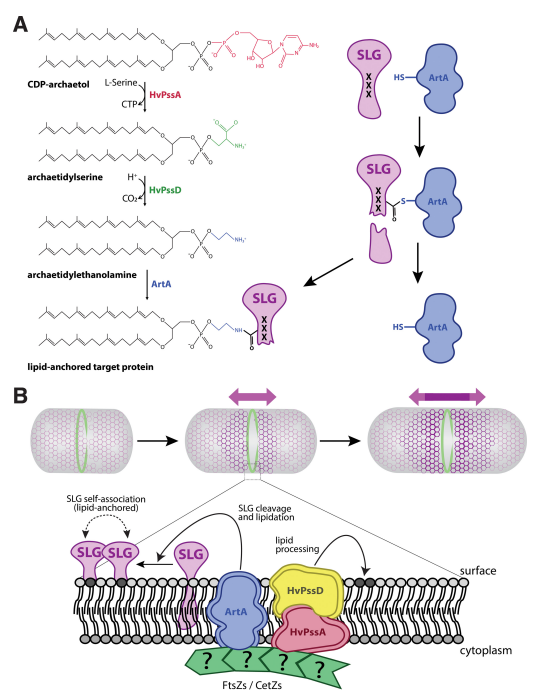


FIG 5 A model for lipid attachment and cell growth involving HvPssA, HvPssD, and ArtA. (A) In our speculative model, CDP-archaeol is converted to archaetidylethanolamine in two steps involving HvPssA and HvPssD. ArtA acts as a peptidase and covalently links its active-site cysteine to a newly generated C terminus of its target protein, simultaneously releasing the C-terminal peptide. Then, the free amino group of ethanolamine attacks the thiocarboxylate, which marks the covalent attachment of the target protein to the ArtA active-site cysteine. This results in covalent attachment of the lipid to the C terminus of the target protein as a carboxamide, simultaneously releasing ArtA. The process of cleavage and lipidation is dependent on HvPssA and HvPssD, either by binding of archaetidylethanolamine to ArtA or by protein-protein interaction between ArtA and HvPssA or HvPssD. (B) Recruitment of ArtA, HvPssA, or HvPssD to the midcell promotes anchoring of surface proteins and insertion of new SLG into the S-layer at midcell, contributing to cell elongation and division.

although it has been shown that the S-layer is not essential in other archaea (29, 30), ectopic overexpression of our SLG-msfGFP fusion drastically impacted the morphology of H. volcanii cells (Fig. 4D) beyond the lack of SLG processing and lipidation (11). Therefore, it is possible that our SLG-msfGFP fusion is actually blocking the transport of

or interaction with other yet-unknown surface proteins essential for shape maintenance. The concept of having different classes of surface-modifying proteins counteracting each other has been demonstrated in bacteria, where different classes of penicillin binding proteins (PBPs) act on the peptidoglycan cell wall to control cell width homeostasis in rod-shaped cells (31).

Interestingly, just like ovococcoid bacteria that are capable of midcell elongation and lack a clear dedicated elongation machinery, haloarchaea may be also employing cytoskeletal polymers to direct different subcomplexes for cell elongation and cell division (32). This evidence is even more striking for coccoid bacteria, for which a single point mutation in FtsZ is able to induce cell elongation in Staphylococcus aureus (33). However, haloarchaea might have conserved at least two distinct elongation modes in addition to cell division, generating disklike and rodlike populations (34). This scenario would also corroborate the morphological malleability of haloarchaea, being capable of assuming unusual shapes like triangles and squares (35, 36).

In spite of the bacteria that use the localization of specialized proteins to promote midcell elongation, it is important to point out that these mechanisms are likely not evolutionarily related to the proposed haloarchaeal S-layer lipidation and cell elongation. First, even though there are examples of bacterial species with an S-layer that carries out peptidoglycan cell wall synthesis at the midcell (6), their new S-layer material is inserted as patches distributed all around the cell (37, 38). Second, the lack of conservation in the protein architecture between archaeal and bacterial S-layers argues that they may have emerged independently of each other (8, 39).

In conclusion, by applying a set of different experimental approaches, we have confirmed that two putative lipid biosynthesis enzymes, HvPssA and HvPssD, are involved in the proteolytic cleavage and lipid labeling of ArtA substrates specifically at midcell. We have also proposed, to the best of our knowledge, the first molecular model for archaeal cell elongation.

#### **MATERIALS AND METHODS**

Strains and growth conditions. The plasmids and strains used in this study are listed in Table 1. H. volcanii strain H53 and its derivatives were grown at 45°C in semidefined Casamino Acids (CA) medium supplemented with tryptophan (50  $\mu$ g ml<sup>-1</sup> final concentration) (40). Cells were cultivated either in liquid medium (orbital shaker at 250 rpm) or on solid 1.5% agar. Difco agar and Bacto yeast extract were purchased from Becton, Dickinson, and Company. Peptone was purchased from Oxoid. To ensure equal agar concentrations in all plates, agar was completely dissolved in the media prior to autoclaving, and the autoclaved media were stirred before plates were poured. Escherichia coli strains were grown at 37°C in NZCYM medium (Fisher Scientific) supplemented with ampicillin (100 µg/ml).

Plasmid preparation and H. volcanii transformation. DNA polymerase, DNA ligase, and restriction enzymes were purchased from New England BioLabs. Plasmids were initially transformed into E. coli DH5 $\alpha$  cells. Plasmid preparations were performed using the QIAprep Spin miniprep (Qiagen) kit. Prior to H. volcanii transformation, plasmids were transformed into the Dam- E. coli strain DL739. H. volcanii transformations were performed using the polyethylene glycol (PEG) method (40). All oligonucleotides used to construct the recombinant plasmids are listed in Table S1.

Generation of chromosomal hvpssA and hvpssD deletions in H53. Chromosomal deletions were generated by homologous recombination (pop-in/pop-out), as previously described (19). Plasmid constructs for use in the pop-in/pop-out knockout process were generated by using overlap PCR, as described previously (41), as follows: approximately 700 nucleotides flanking the hypssA gene were PCR amplified and cloned into the haloarchaeal suicide vector pTA131. The hvpssA upstream flanking region was amplified with oligonucleotides FW\_pssA\_KO\_Xbal and RV\_pssA\_up, while the hvpssA downstream flanking region was amplified using FW\_pssA\_dw and RV\_pssA\_KO\_XhoI (oligonucleotides are listed in Table S1). The hvpssA upstream and downstream flanking DNA fragments were fused by PCR using oligonucleotides FW\_pssA\_KO\_Xbal and RV\_pssA\_KO\_Xhol, followed by cloning into pTA131 digested with Xbal and Xhol. The insertion of the correct DNA fragment into the cloning site of the recombinant plasmid was verified by sequencing using the same oligonucleotides. The final plasmid construct, pFH38, contained upstream and downstream hvpssA flanking regions and was transformed into the parental H53 H. volcanii strain. To confirm the chromosomal replacement event at the proper location on the chromosome, colonies derived from these techniques were screened by PCR using the FW\_pssA\_KO\_Xbal and RV\_pssA\_KO\_Xhol oligonucleotides. The hvpssA deletion mutant generated in strain H53 was designated FH38 (Table 1). For the generation of the plasmid construct for chromosomal hvpssD deletion, approximately 700 nucleotides flanking the hvpssD gene were PCR amplified and cloned into the haloarchaeal suicide vector pTA131. The upstream flanking region was amplified with oligonucleotides FW\_pssD\_KO\_Xbal and RV\_pssD\_up, while the downstream flanking region was amplified using FW\_pss-D\_dw and RV\_pssD\_KO\_Xhol. The flanking DNA fragments were fused by PCR using oligonucleotides

TABLE 1 Strains and plasmids used in this work

Plasmid or strain	Genotype and/or description <sup>a</sup>	Reference or source
Plasmids		
pTA131	pBluescript II with BamHI-Xbal fragments from pGB70 harboring p.fdx-pyrE2, Amp <sup>r</sup>	19
pTA963	pyrE2 and hdrB markers, Trp-inducible (p.tna) promoter, Amp <sup>r</sup>	43
pFH25	pTA963 carrying hvo_0405 with C-terminal 6×His tag	12
pFH38	pTA131 carrying 700 bp upstream and 700 bp downstream hvpssA flanking region	This work
pFH43	pTA131 carrying 700 bp upstream and 700 bp downstream hvpssD flanking region	This work
pFH39	pTA963 carrying hvpssA with C-terminal 6×His tag	This work
pFH44	pTA963 carrying hvpssD with C-terminal 6×His tag	This work
pFH55	pTA963 carrying $hvo\_0405$ N-terminal LVIVD domain with C-terminal 6×His tag	This work
Strains		
DH5lpha	E. coli F $^ \phi$ 80 $\Delta$ lacZ $\Delta$ M15 (lacZYA-argF)U169 recA1 endA1 hsdR17( $r_K^ m_K^-$ ) phoA supE44 thi-1 gyrA96 relA1	Invitrogen
DL739	E. coli MC4100 recA dam-13::Tn9	48
H26	H. volcanii ΔpyrE2	19
H53	H. volcanii ΔpyrE2 ΔtrpA	19
FH27	H53/pTA963::hvo_0405-6×His	12
FH28	H53 Δ <i>artA</i> /pTA963:: <i>hvo_0405</i> -6×His	12
FH55	H53 ΔhvpssA/pTA963	This work
FH56	H53 ΔhvpssA/pTA963::hvpssA-6×His	This work
FH57	H53 ΔhvpssA/pTA963::hvo_0405-6×His	This work
FH69	H53 ΔhvpssD/pTA963	This work
FH70	H53 ΔhvpssD/pTA963::hvpssD-6×His	This work
FH71	H53 ΔhvpssD/pTA963::hvo_0405-6×His	This work
FH77	H53 Δhvo_0405/pTA963::N-term-hvo_0405-6×His	This work
aBL128	ΔpyrE2 artA::artA-msfGFP-pyrE2	This work
aBL129	ΔpyrE2 hvpssD::hvpssD-msfGFP-pyrE2	This work
aBL183	ΔpyrE2 hvpssA::hvpssA-msfGFP-pyrE2	This work
aBL131	ΔpyrE2 ftsZ1::ftsZ1-msfGFP-pyrE2	This work
aBL184	ΔpyrE2/pTA962::msfGFP(SW)	This work
aBL118	ΔpyrE2/pTA962::csg-msfGFP(SW)	This work

<sup>&</sup>lt;sup>a</sup>Amp<sup>r</sup>, ampicillin resistant.

FW\_pssD\_KO\_Xbal and RV\_pssD\_KO\_Xhol, followed by cloning into pTA131 digested with Xbal and Xhol. The insertion of the correct DNA fragment into the cloning site of the recombinant plasmid was verified by sequencing using the same oligonucleotides. The final plasmid construct, pFH43, contained upstream and downstream *hvpssD* flanking regions and was transformed into the parental *H. volcanii* strain H53. Confirmation of *hvpssD* deletion on the chromosome was screened by PCR using the FW\_pssD\_KO\_Xbal and RV\_pssD\_KO\_Xhol oligonucleotides. The *hvpssD* deletion mutant generated in strain H53 was designated FH63 (Table 1).

**Construction of expression plasmids for HvPssA and HvPssD.** To construct a tryptophan-inducible *H. volcanii hvpssA* gene with C-terminal His tag, its coding region was amplified by PCR using the oligonucleotides FW\_pssA\_OE\_Ndel and RV\_pssA\_OE\_EcoRl\_His (Table S1). Meanwhile, for the construction of the *H. volcanii hvpssD* gene with a C-terminal His tag, its coding region was amplified by PCR using the oligonucleotides FW\_pssD\_OE\_Ndel and RV\_pssD\_OE\_EcoRl\_His (Table S1). The PCR product was cloned into the expression vector pTA963 that had been digested with Ndel and EcoRl. This places the *hvpssA* or *hvpssD* gene under the control of the inducible tryptophanase promoter (p.tna). The recombinant pTA963 carrying the *hvpssA* gene was designated pFH39, and the pTA963 carrying *hvpssD* was designated pFH44. To complement the Δ*hvpssA* mutant strain FH38, this strain was transformed with plasmid pFH39 to result in FH56. For complementation of the Δ*hvpssD* mutant strain FH63, this strain was transformed with plasmid pFH44 to result in FH70. The H53, Δ*hvpssA* mutant, and Δ*hvpssD* mutant strains were also transformed with the empty expression vector pTA963, which was used as a control.

Construction of expression plasmid for SLG-msfGFP and msfGFP. To construct a tryptophan-inducible H.  $volcanii\ csg$  gene tagged with msfGFP, a sandwich fusion was created by intercalating an msfGFP gene product amplified from a synthetic fragment (oligonucleotides oHV81 and oHV82) between the first 102 bp (the secretion signal sequence of SLG, oligonucleotides oAB500 and oAB501) and the rest of the csg coding region (oligonucleotides oHV83 and oAB502). The 3 fragments were then assembled by Gibson Assembly (42) together with the pTA962 plasmid (43), digested with Nhel and BamHI, and transformed into DH5 $\alpha$  cells. The same process was followed for the cloning of msfGFP alone (oligonucleotides oHV101 and oHM68). The clones were subjected to validation by PCR and Sanger sequencing (Table S1).

**Generation of chromosomal msfGFP fusions in H26.** C-terminal translational fusion constructs, including a pyrE2 cassette for selection, were generated by direct transformation of PCR fragment ensembles generated by Gibson Assembly (42), transformed directly in *H. volcanii* H26 cells, and selected by growth in the absence of uracil. PCR fragments from *ftsZ1* (oligonucleotides oHV3 and oHV4 for the upstream region and oHV8 and oHV9 for the downstream region), *artA* (oligonucleotides oHM91 and

Abdul-Halim et al. mBio

oHM92 for the upstream region and oHM93 and oHM94 for the downstream region), *hvpssA* (oligonucleotides oHV156 and oHV157 for the upstream region and oHV158 and oHV159 for the downstream region), and *hvpssD* (oligonucleotides oHV126 and oHV127 for the upstream region and oHV128 and oHV129 for the downstream region) were assembled to msfGFP (oligonucleotides oHM34 and oHM6) and the *pyrE2* cassette (oligonucleotides oHV6 and oHV7). Chromosomal replacements were confirmed by PCR and Sanger sequencing (Table S1).

Immunoblotting. Liquid cultures were grown until mid-log phase (optical density at 600 nm [OD<sub>600</sub>], 0.2 to 0.5), and the cells were harvested by centrifugation at 3,800  $\times$  q for 5 min at room temperature. Cell pellets were resuspended and lysed in 1% (vol/vol) NuPAGE lithium dodecyl sulfate (LDS) sample buffer supplemented with 100 mM dithiothreitol (DTT) and stored at -20°C. Samples were electrophoresed on 4 to 12% Bis-Tris polyacrylamide gels (Invitrogen) with NuPAGE 3-(N-morpholino)propanesulfonic acid (MOPS)-SDS running buffer (Invitrogen). Proteins were then transferred to polyvinylidene difluoride (PVDF) membranes (Millipore) using a semidry transfer apparatus at 15 V for 30 min (Bio-Rad). Subsequently, the membrane was washed twice in phosphate-buffered saline (PBS), blocked for 1 h in 3% bovine serum albumin (BSA) in PBS, and washed twice in PBS with 1% Tween 20 and once with PBS. For detection of the poly-His tag, the mouse anti-penta-His antibody (catalog no. 34660; Qiagen) was used at a 1:2,000 dilution in 3% BSA in PBS with sodium azide. For the secondary antibody, horseradish peroxidase (HRP)-conjugated Amersham ECL anti-mouse IgG from sheep (GE) was used at a 1:20,000 dilution in 10% nonfat milk in PBS. For the detection of HVO 0405 Nterm, the rabbit anti-HVO 0405-N-term serum (12) was used as the primary antibody at a 1:10,000 dilution in 3% BSA in PBS with sodium azide. For the secondary antibody, HRP-conjugated Amersham ECL anti-rabbit IgG from donkey (GE) was used at a 1:60,000 dilution in 10% nonfat milk in PBS.

**Lipid radiolabeling.** The H53 parent strain carrying the vector control pTA963 and the Δ*hvpssA* mutant strain carrying either the *hvpssA* expression plasmid pFH39 or the vector control pTA963 were grown in 5 ml liquid CA medium. Upon reaching mid-log phase  $(OD_{600}, \sim 0.5)$ , 20  $\mu$ l of each culture was transferred into 1 ml of fresh liquid CA medium supplemented with [14C]mevalonic acid (resuspended in ethanol) at a final concentration of 1  $\mu$ Ci/ml. *Haloferax volcanii* cultures were harvested after reaching mid-log phase, and proteins were precipitated from 1-ml cultures with 10% trichloroacetic acid (TCA), followed by a delipidation step to remove noncovalently linked lipid, as described previously (44, 45). The delipidated proteins were separated by 7% Tris-acetate (TA) LDS-PAGE gels. For analysis of the samples, the gel was dried onto blotting paper using a gel dryer (model 583; Bio-Rad), exposed to a phosphor screen (Molecular Dynamics) for 3 weeks, and analyzed using a Typhoon imager (Amersham Biosciences).

**Motility assays.** The motility assays of the *H. volcanii* H53 (parent) and the  $\Delta artA$ ,  $\Delta hvpssA$ , and  $\Delta hvpssD$  mutant strains carrying the plasmid expressing the complementary gene (or pTA963 as control) were performed on 0.35% agar in CA medium supplemented with tryptophan, as described previously (41). A toothpick was used to stab inoculate the agar, followed by incubation at 45°C. Halo sizes around the stab-inoculation site were measured after 3 to 5 days of incubation.

**Growth curves.** Growth curves were measured using a BioTek PowerWaveX2 microplate spectrophotometer. *H. volcanii* H53 (parent) and  $\Delta artA$ ,  $\Delta hvpssA$ , and  $\Delta hvpssD$  mutant strains carrying the plasmid expressing the complementary gene (or pTA963 as control) were first incubated in 5-ml liquid cultures in CA medium supplemented with tryptophan with continuous shaking at 45°C until suitable OD<sub>600</sub> values (0.2 to 0.5) were reached. Approximately 6  $\mu$ l of each culture (adjusted to correct for OD<sub>600</sub> differences) was then transferred into 194  $\mu$ l of fresh CA medium supplemented with tryptophan (50  $\mu$ g ml<sup>-1</sup> final concentration) and grown to stationary phase, with OD<sub>600</sub> recordings taken every 30 min.

**Light microscopy.** The *H. volcanii* strains H53 (parent) and  $\Delta artA$ ,  $\Delta hvpssA$ , and  $\Delta hvpssD$  mutant strains carrying the plasmid expressing the complementary gene (or pTA963 as control) were inoculated from colony to 5 ml CA liquid medium and grown until they reached mid-log phase (OD<sub>600</sub>, ~0.4 to 0.5). Serial liquid-to-liquid subinoculations were carried out by transferring 10  $\mu$ l of the liquid culture to 5 ml fresh liquid CA medium with up to two transfers. Subsequently, 1 ml of each culture was concentrated by centrifugation at 4,911 × g for 1 min, and the pellets were resuspended in 10  $\mu$ l of liquid CA medium. Then, 10  $\mu$ l of the concentrated cells was transferred to under a 0.5% agarose pad with CA medium and observed using a Nikon Eclipse TiE inverted total internal reflection fluorescence (TIRF) microscope. ArtA-msfGFP time-lapse images were acquired by culturing *H. volcanii* cells inside Millipore ONIX CellASIC microfluidic plates, as previously described (35). Images were taken under 5-min intervals for 12 h in both phase-contrast and 488-nm laser channels. *H. volcanii* protoplasts were generated within microfluidic channels by the addition of 1 mg/ml of proteinase K (Invitrogen) in YPC medium until cells lost shape. Subsequently, the cells were washed with fresh YPC, and time-lapse images were recorded with 10-min intervals for 12 h.

**Image analysis.** All image processing, unless otherwise specified, was performed in Fiji. Automated segmentation of cells and subsequent aspect ratio measurements were performed using the MicrobeJ plugin (46). Finally, statistics and plots were generated using ggplot2 in RStudio.

**Gene synteny analysis.** We used the SyntTax server (https://archaea.i2bc.paris-saclay.fr/synttax/) (47) in order to evaluate if the orthologs of *artA*, *hvpssA*, and *hvpssD* are genomically clustered in other haloarchaea. According to the inspection of more than 70 haloarchaeal genomes, the genes for orthologs of *artA*, *hvpssA*, and *hvpssD* are not clustered.

### SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

MOVIE S1, MOV file, 15 MB.

MOVIE S2, MOV file, 2.7 MB.

FIG S1, TIF file, 2.3 MB.

**FIG S2**, TIF file, 1.5 MB.

**FIG S3**, TIF file, 2.8 MB. **FIG S4**, TIF file, 2.3 MB.

rid 34, fir file, 2.3 Mib

**FIG S5**, TIF file, 2 MB.

TABLE S1, DOCX file, 0.1 MB.

#### **ACKNOWLEDGMENTS**

We thank Howard Goldfine and the Pohlschroder, Daldal, and Garner labs for helpful discussions. We also thank Henry Miziorko, Ethan Garner, and Jenny Zheng for access to equipment and reagents.

M.P., A.D., and M.F.A.-H. were supported by the National Science Foundation grant 1817518. A.W.B.F. was supported by his personal startup fund from Brandeis University. S.S. was supported by the German Research Foundation Postdoctoral Fellowship (DFG grant 398625447).

#### **REFERENCES**

- Bisson-Filho AW, Zheng J, Garner E. 2018. Archaeal imaging: leading the hunt for new discoveries. Mol Biol Cell 29:1675–1681. https://doi.org/10 .1091/mbc.E17-10-0603.
- de Pedro MA, Quintela JC, Höltje JV, Schwarz H. 1997. Murein segregation in Escherichia coli. J Bacteriol 179:2823–2834. https://doi.org/10.1128/jb.179.9.2823-2834.1997.
- May JW, Mitchison JM. 1986. Length growth in fission yeast cells measured by two novel techniques. Nature 322:752–754. https://doi.org/10.1038/322752a0.
- Brown PJB, de Pedro MA, Kysela DT, Van der Henst C, Kim J, De Bolle X, Fuqua C, Brun YV. 2012. Polar growth in the alphaproteobacterial order Rhizobiales. Proc Natl Acad Sci U S A 109:1697–1701. https://doi.org/10 .1073/pnas.1114476109.
- Sipiczki M, Takeo K, Grallert A. 1998. Growth polarity transitions in a dimorphic fission yeast. Microbiology 144:3475–3485. https://doi.org/10 .1099/00221287-144-12-3475.
- Aaron M, Charbon G, Lam H, Schwarz H, Vollmer W, Jacobs-Wagner C. 2007. The tubulin homologue FtsZ contributes to cell elongation by guiding cell wall precursor synthesis in Caulobacter crescentus. Mol Microbiol 64: 938–952. https://doi.org/10.1111/j.1365-2958.2007.05720.x.
- Rodrigues-Oliveira T, Belmok A, Vasconcellos D, Schuster B, Kyaw CM. 2017. Archaeal S-layers: overview and current state of the art. Front Microbiol 8:2597. https://doi.org/10.3389/fmicb.2017.02597.
- Pohlschroder M, Pfeiffer F, Schulze S, Halim M. 2018. Archaeal cell surface biogenesis. FEMS Microbiol Rev 42:694–717. https://doi.org/10 .1093/femsre/fuy027.
- Eun Y-J, Ho P-Y, Kim M, LaRussa S, Robert L, Renner LD, Schmid A, Garner E, Amir A. 2018. Archaeal cells share common size control with bacteria despite noisier growth and division. Nat Microbiol 3:148–154. https://doi.org/10.1038/s41564-017-0082-6.
- Abdul Halim MF, Pfeiffer F, Zou J, Frisch A, Haft D, Wu S, Tolić N, Brewer H, Payne SH, Paša-Tolić L, Pohlschroder M. 2013. *Haloferax volcanii* archaeosortase is required for motility, mating, and C-terminal processing of the S-layer glycoprotein. Mol Microbiol 88:1164–1175. https://doi .org/10.1111/mmi.12248.
- Abdul Halim MF, Karch KR, Zhou Y, Haft DH, Garcia BA, Pohlschroder M. 2015. Permuting the PGF signature motif blocks both archaeosortasedependent C-terminal cleavage and prenyl lipid attachment for the Haloferax volcanii S-layer glycoprotein. J Bacteriol 198:808–815. https:// doi.org/10.1128/JB.00849-15.
- Abdul Halim MF, Stoltzfus JD, Schulze S, Hippler M, Pohlschroder M. 2017. ArtA-dependent processing of a Tat substrate containing a conserved tripartite structure that is not localized at the C terminus. J Bacteriol 199:e00802-16. https://doi.org/10.1128/JB.00802-16.
- Abdul Halim MF, Rodriguez R, Stoltzfus JD, Duggin IG, Pohlschroder M. 2018. Conserved residues are critical for *Haloferax volcanii* archaeosortase catalytic activity: implications for convergent evolution of the catalytic mechanisms of non-homologous sortases from archaea and bacteria. Mol Microbiol 108:276–287. https://doi.org/10.1111/mmi.13935.
- 14. Haft DH, Payne SH, Selengut JD. 2012. Archaeosortases and exosortases

- are widely distributed systems linking membrane transit with posttranslational modification. J Bacteriol 194:36–48. https://doi.org/10.1128/JB .06026-11.
- 15. Morii H, Koga Y. 2003. CDP-2,3-di-O-geranylgeranyl-sn-glycerol:1-serine O-archaetidyltransferase (archaetidylserine synthase) in the methanogenic archaeon Methanothermobacter thermautotrophicus. J Bacteriol 185:1181–1189. https://doi.org/10.1128/jb.185.4.1181-1189.2003.
- Dutt A, Dowhan W. 1981. Characterization of a membrane-associated cytidine diphosphate-diacylglycerol-dependent phosphatidylserine synthase in bacilli. J Bacteriol 147:535–542. https://doi.org/10.1128/JB.147 .2.535-542.1981.
- 17. Okada M, Matsuzaki H, Shibuya I, Matsumoto K. 1994. Cloning, sequencing, and expression in Escherichia coli of the Bacillus subtilis gene for phosphatidylserine synthase. J Bacteriol 176:7456–7461. https://doi.org/10.1128/jb.176.24.7456-7461.1994.
- Nishibori A, Kusaka J, Hara H, Umeda M, Matsumoto K. 2005. Phosphatidylethanolamine domains and localization of phospholipid synthases in Bacillus subtilis membranes. J Bacteriol 187:2163–2174. https://doi.org/ 10.1128/JB.187.6.2163-2174.2005.
- Allers T, Ngo H-P, Mevarech M, Lloyd RG. 2004. Development of additional selectable markers for the halophilic archaeon *Haloferax volcanii* based on the leuB and trpA genes. Appl Environ Microbiol 70:943–953. https://doi.org/10.1128/aem.70.2.943-953.2004.
- 20. Banerjee A, Tsai C-L, Chaudhury P, Tripp P, Arvai AS, Ishida JP, Tainer JA, Albers S-V. 2015. FlaF is a  $\beta$ -sandwich protein that anchors the archaellum in the archaeal cell envelope by binding the S-layer protein. Structure 23:863–872. https://doi.org/10.1016/j.str.2015.03.001.
- Tsai C-L, Tripp P, Sivabalasarma S, Zhang C, Rodriguez-Franco M, Wipfler RL, Chaudhury P, Banerjee A, Beeby M, Whitaker RJ, Tainer JA, Albers S-V. 2020. The structure of the periplasmic FlaG-FlaF complex and its essential role for archaellar swimming motility. Nat Microbiol 5:216–225. https://doi.org/10.1038/s41564-019-0622-3.
- Duggin IG, Aylett CHS, Walsh JC, Michie KA, Wang Q, Turnbull L, Dawson EM, Harry EJ, Whitchurch CB, Amos LA, Löwe J. 2015. CetZ tubulin-like proteins control archaeal cell shape. Nature 519:362–365. https://doi .org/10.1038/nature13983.
- Dinh T, Bernhardt TG. 2011. Using superfolder green fluorescent protein for periplasmic protein localization studies. J Bacteriol 193:4984–4987. https://doi.org/10.1128/JB.00315-11.
- Gadwal S, Johnson TL, Remmer H, Sandkvist M. 2018. C-terminal processing of GlyGly-CTERM containing proteins by rhombosortase in Vibrio cholerae. PLoS Pathog 14:e1007341. https://doi.org/10.1371/journal.ppat.1007341.
- Daiyasu H, Kuma K-I, Yokoi T, Morii H, Koga Y, Toh H. 2005. A study of archaeal enzymes involved in polar lipid synthesis linking amino acid sequence information, genomic contexts and lipid composition. Archaea 1:399–410. https://doi.org/10.1155/2005/452563.
- Kellermann MY, Yoshinaga MY, Valentine RC, Wörmer L, Valentine DL.
   2016. Important roles for membrane lipids in haloarchaeal bioenerget-

- ics. Biochim Biophys Acta 1858:2940–2956. https://doi.org/10.1016/j.bbamem.2016.08.010.
- Kandiba L, Guan Z, Eichler J. 2013. Lipid modification gives rise to two distinct *Haloferax volcanii* S-layer glycoprotein populations. Biochim Biophys Acta 1828:938–943. https://doi.org/10.1016/j.bbamem.2012.11 023
- 28. Irihimovitch V, Eichler J. 2003. Post-translational secretion of fusion proteins in the halophilic archaea *Haloferax volcanii*. J Biol Chem 278: 12881–12887. https://doi.org/10.1074/jbc.M210762200.
- Zhang C, Phillips APR, Wipfler RL, Olsen GJ, Whitaker RJ. 2018. The essential genome of the crenarchaeal model Sulfolobus islandicus. Nat Commun 9:4908. https://doi.org/10.1038/s41467-018-07379-4.
- Zhang C, Wipfler RL, Li Y, Wang Z, Hallett EN, Whitaker RJ. 2019. Cell structure changes in the hyperthermophilic crenarchaeon Sulfolobus islandicus lacking the S-layer. mBio 10:e01589-19. https://doi.org/10 .1128/mBio.01589-19.
- Dion MF, Kapoor M, Sun Y, Wilson S, Ryan J, Vigouroux A, van Teeffelen S, Oldenbourg R, Garner EC. 2019. Bacillus subtilis cell diameter is determined by the opposing actions of two distinct cell wall synthetic systems. Nat Microbiol 4:1294–1305. https://doi.org/10.1038/s41564-019-0439-0.
- 32. Pérez-Núñez D, Briandet R, David B, Gautier C, Renault P, Hallet B, Hols P, Carballido-López R, Guédon E. 2011. A new morphogenesis pathway in bacteria: unbalanced activity of cell wall synthesis machineries leads to coccus-to-rod transition and filamentation in ovococci. Mol Microbiol 79:759–771. https://doi.org/10.1111/j.1365-2958.2010.07483.x.
- Pereira AR, Hsin J, Król E, Tavares AC, Flores P, Hoiczyk E, Ng N, Dajkovic A, Brun YV, VanNieuwenhze MS, Roemer T, Carballido-Lopez R, Scheffers D-J, Huang KC, Pinho MG. 2016. FtsZ-dependent elongation of a coccoid bacterium. mBio 7:e00908-16. https://doi.org/10.1128/mBio.00908-16.
- 34. Li Z, Kinosita Y, Rodriguez-Franco M, Nußbaum P, Braun F, Delpech F, Quax TEF, Albers S-V. 2019. Positioning of the motility machinery in halophilic archaea. mBio 10:e00377-19. https://doi.org/10.1128/mBio.00377-19.
- Walsh JC, Angstmann CN, Bisson-Filho AW, Garner EC, Duggin IG, Curmi P. 2019. Division plane placement in pleomorphic archaea is dynamically coupled to cell shape. Mol Microbiol 112:785–799. https://doi.org/10 .1111/mmi.14316.
- Walsby AE. 1980. A square bacterium. Nature 283:69 –71. https://doi.org/ 10.1038/283069a0.
- Comerci CJ, Herrmann J, Yoon J, Jabbarpour F, Zhou X, Nomellini JF, Smit J, Shapiro L, Wakatsuki S, Moerner WE. 2019. Topologically-guided

- continuous protein crystallization controls bacterial surface layer self-assembly. Nat Commun 10:2731. https://doi.org/10.1038/s41467-019-10650-x.
- Oatley P, Kirk JA, Ma S, Jones S, Fagan RP. 2018. Spatial organization of Clostridium difficile S-layer biogenesis. bioRxiv https://www.biorxiv.org/ content/10.1101/405993v2.full.
- 39. Bharat TAM, Kureisaite-Ciziene D, Hardy GG, Yu EW, Devant JM, Hagen WJH, Brun YV, Briggs JAG, Löwe J. 2017. Structure of the hexagonal surface layer on Caulobacter crescentus cells. Nat Microbiol 2:17059. https://doi.org/10.1038/nmicrobiol.2017.59.
- Dyall-Smith M. 2009. The Halohandbook: protocols for halobacterial genetics v.7.3. https://haloarchaea.com/wp-content/uploads/2018/10/ Halohandbook\_2009\_v7.3mds.pdf.
- 41. Tripepi M, Imam S, Pohlschröder M. 2010. *Haloferax volcanii* flagella are required for motility but are not involved in PibD-dependent surface adhesion. J Bacteriol 192:3093–3310. https://doi.org/10.1128/JB.00133-10.
- Gibson DG, Young L, Chuang R-Y, Venter JC, Hutchison CA, Smith HO. 2009. Enzymatic assembly of DNA molecules up to several hundred kilobases. Nat Methods 6:343–345. https://doi.org/10.1038/nmeth.1318.
- Allers T, Barak S, Liddell S, Wardell K, Mevarech M. 2010. Improved strains and plasmid vectors for conditional overexpression of His-tagged proteins in *Haloferax volcanii*. Appl Environ Microbiol 76:1759–1769. https:// doi.org/10.1128/AEM.02670-09.
- Kikuchi A, Sagami H, Ogura K. 1999. Evidence for covalent attachment of diphytanylglyceryl phosphate to the cell-surface glycoprotein of Halobacterium halobium. J Biol Chem 274:18011–18016. https://doi.org/10 .1074/jbc.274.25.18011.
- 45. Konrad Z, Eichler J. 2002. Protein glycosylation in *Haloferax volcanii*: partial characterization of a 98-kDa glycoprotein. FEMS Microbiol Lett 209:197–202. https://doi.org/10.1111/j.1574-6968.2002.tb11131.x.
- Ducret A, Quardokus EM, Brun YV. 2016. MicrobeJ, a tool for high throughput bacterial cell detection and quantitative analysis. Nat Microbiol 1:16077. https://doi.org/10.1038/nmicrobiol.2016.77.
- 47. Oberto J. 2013. SyntTax: a Web server linking synteny to prokaryotic taxonomy. BMC Bioinformatics 14:4. https://doi.org/10.1186/1471-2105-14-4.
- Blattner FR, Williams BG, Blechl AE, Denniston-Thompson K, Faber HE, Furlong L, Grunwald DJ, Kiefer DO, Moore DD, Schumm JW, Sheldon EL, Smithies O. 1977. Charon phages: safer derivatives of bacteriophage lambda for DNA cloning. Science 196:161–169. https://doi.org/10.1126/ science.847462.