Archaeal Signal Transduction: Impact of Protein Phosphatase Deletions on Cell Size, Motility, and Energy Metabolism in Sulfolobus acidocaldarius*

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In this study, the in vitro and in vivo functions of the only two identified protein phosphatases, Saci-PTP and Saci-PP2A, in the crenarchaeal model organism Sulfolobus acidocaldarius were investigated. Biochemical characterization revealed that Saci-PTP is a dual-specific phosphatase (against pSer/pThr and pTyr), whereas Saci-PP2A exhibited specific pSer/pTyr activity and inhibition by okadaic acid. Deletion of saci_pp2a resulted in pronounced alterations in growth, cell shape and cell size, which could be partially complemented. Transcriptome analysis of the three strains (Δsaci_pptp, Δsaci_pp2a and the MW001 parental strain) revealed 155 genes that were differentially expressed in the deletion mutants, and showed significant changes in expression of genes encoding the archaea (archaeal motility structure), components of the respiratory chain and transcriptional regulators. Phosphoproteome studies revealed 801 unique phosphoproteins in total, with an increase in identified phosphopeptides.

highlight the importance of protein phosphorylation in regulating essential cellular processes in the crenarchaeon S. acidocaldarius. Molecular & Cellular Proteomics 12: 10.1074/mcp.M113.027375, 3908–3923, 2013.

Protein phosphorylation is an important post-translational modification (PTM) that has been reported in all three domains of life, Eukarya, Bacteria and Archaea (1–3). The reversible character of the modification allows for subtle and immediate regulation by modulating protein activity in different cellular processes (3–5). The addition of a phosphoryl group is catalyzed by protein kinases and dephosphorylation by their cognate phosphatases. Interestingly, in the Crenarchaeota, one of the phyla of the Archaea, only Ser/Thr/Tyr phosphorylation is predicted by bioinformatics analyses, whereas two-component systems (involving His/Asp phosphorylation) identified in the Euryarchaeota phylum are absent (6–9).

A phosphoproteome study of the thermoacidophilic crenarchaeon Sulfolobus solfataricus P2 revealed a vast amount of Ser/Thr/Tyr-phosphorylated proteins (540 detected in total), in almost all arCOGs categories, with an unexpectedly high number of Tyr phosphorylation. A differential phosphorylation pattern was observed in cells grown on glucose versus tryptone, and a role of protein phosphorylation in regulating the glycolytic flux in Sulfolobus was proposed (10). Further, a phosphoproteome study of the mesophilic euryarchaeon Halobacterium salinarum revealed 69 phosphorylated proteins in total (2).

Although a few archaeal protein kinases and phosphatases have been investigated in more detail (7, 11–19), there is still a knowledge gap regarding signal transduction pathways in Archaea and the impact of Ser/Thr/Tyr phosphorylation on cellular processes. Comparative database searches revealed only two protein phosphatases encoded in the genomes of all...
members of the *Sulfolobales* order, in comparison to eight predicted protein kinases in *S. solfataricus* alone (7). Similarly, in eukaryotes fewer protein phosphatases are encoded in the genome compared with protein kinases, though the phosphatases of the PPP family often act as multimeric proteins with different catalytic, regulatory and core subunits (20, 21).

Protein phosphatases can be categorized into different families. Ser/Thr-specific PPPs contain a 220 amino acid long catalytic domain that includes motif I (GDXXH), motif II (GDXXDRG), and motif III (GNHE) (22), and Mg\(^{2+}\) or Mn\(^{2+}\) dependent Ser/Thr phosphatases (PPMs) contain 11 specific motifs (23, 24). Conversely, the phosphotyrosine phosphatase family (PTP) can be subdivided into the PTPs, which are specific for Tyr dephosphorylation, the dual specific PTPs, which can dephosphorylate Ser/Thr and Tyr, and the low molecular weight PTPs. PTP phosphatase family members share a common amino acid motif, CX\(_2\)R (20).

Only a few archaean Ser/Thr- or Tyr-specific protein phosphatases have been characterized. The Ser/Thr phosphatase PP1-arch1 from *S. solfataricus* displays Mn\(^{2+}\)-dependent protein phosphatase activity (in vitro) (16). The crystal structure of SsoPTP from *S. solfataricus* has been solved and the specificity toward phosphotyrosine determined in vitro (18). Only one additional archaean PTP has been characterized, the Tk-PTP from the euryarchaeon *Thermococcus kodakaraensis* KOD, which exhibits phosphotyrosine as well as phosphoserine, but no phosphothreonine phosphatase activity (17). Finally, the only characterized archaean member of the PPP family is found in *Thermoplasma volcanium* and has a divergent metal-ion dependent dual-specificity toward phosphorylated Ser/Thr as well as Tyr residues (25).

The crenarchaeon *S. acidocaldarius*, which grows optimally at pH 3 and 76 °C, contains two protein phosphatases, the Saci-PTP (phospho Tyr phosphatase), encoded by the *saci0545* gene (*saci_ptp*) and the Saci-PP2A (phospho Ser/Thr phosphatase) encoded by *saci0884* (*saci_pp2a*), which is homologous to the PP2A catalytic subunit of eukaryotic phosphatases. In this study, Saci-PTP and Saci-PP2A were characterized with respect to substrate specificity and kinetic properties. Furthermore, *in vivo* analyses were performed with deletion mutants to investigate the resulting phenotypes, and additional characterization was carried out by RNA-seq and phosphoproteome analysis.

**EXPERIMENTAL PROCEDURES**

**Strains and Growth Conditions—Escherichia coli** K12 DH5\(_{a}\) (Invitrogen, Breda, The Netherlands) and Rosetta (DE3) (Stratagene, La Jolla, CA) were used for cloning and expression studies, respectively. Both strains were grown under standard conditions as reported recently (26).

The markerless in-frame deletion mutants and the uracil auxotrophic parental strain *S. acidocaldarius* MW001 (27) were grown at 76 °C in Brock’s basal medium at pH 3.5 (28). The medium was supplemented with 0.1% (w/v) NZ-amine, 0.2% (w/v) sucrose, and 10 μg/ml uracil (water dissolved). The trans-complementation strains of the deletion mutants were grown without uracil because the uracil auxotrophic marker was encoded by the complementation plasmid.

**Chemicals—**All chemicals were purchased from Sigma-Aldrich (Taufkirchen, Germany), WVR (St. Louis, MO), Carl Roth (Karlsruhe, Germany) or Roche Diagnostics (Mannheim, Germany) in analytical grade. Para-nitrophenylphosphate (pNPP) was purchased from Sigma-Aldrich. The p-Thr-PTP RRA(pT)VA was purchased from Promega (Madison, WI). The p-peptide TEVGKRI(pY)RLVGDKN was synthesized according to the p-ptide of Saci_1938 determined in the phosphoproteome analysis and purchased from PROTEINMODS (Madison, WI). For the characterization of Saci_PTP the p-peptides NIDAIR(pS)NLIMSR and ETTYERW(pI)TTGER derived from the respective p-peptides of Saci_1346 and Saci_1857, respectively, determined in the phospho-proteome analysis were purchased from Thermo Fisher Scientific.

**Heterologous Expression and Protein Purification—**For expression of Saci-PPTP (Saci0545) and Saci-PP2A (Saci0884) the genes were amplified by PCR using specific primers (Table I) and cloned into the PET vector system (PET-15b, or pETDuet-1, Novagen). The constructed plasmids were verified by sequencing of both strands (AGOWA, Berlin and MWG Operon, Ebersberg, Germany).

Heterologous expression of Saci-PTP and Saci-PP2A was performed in *E. coli* Rosetta (DE3)-RIL as expression host strain. For purification of the recombinant enzymes, the resulting *E. coli* crude extracts were diluted 1:1 with 50 mM NaH\(_2\)PO\(_4\), 300 mM NaCl (pH 8.0, RT), and subjected to a heat precipitation for 20 min at 80 °C. After heat precipitation, the samples were cleared by centrifugation (60,000 × g for 30 min at 4 °C).

For Saci-PP2A and Saci-PTP the supernatant was dialyzed overnight against 50 mM NaH\(_2\)PO\(_4\), 300 mM NaCl (pH 8.0, RT), subjected to immobilized metal affinity chromatography (Ni-TED 2000, Macherey-Nagel, pre-equilibrated in 50 mM NaH\(_2\)PO\(_4\), 300 mM NaCl (pH 8.0, RT), and eluted in 50 mM NaH\(_2\)PO\(_4\), 300 mM NaCl, 250 mM imidazole (pH 8.0, RT). Fractions containing the recombinant enzymes (analyzed by SDS-PAGE, enzyme activity) were pooled and concentrated via centrifugal concentrators (Vivaspin 6, Sartorius Ste dim Biotech). From 3.8 g of recombinant cells (wet weight), 1.7 mg Saci-PTP and from 17 g of recombinant cells (wet weight) 0.4 mg Saci-PP2A were obtained.

Saci-PP2A was further purified by ion exchange chromatography and size exclusion chromatography. Briefly, Saci-PP2A was dialyzed overnight against 20 mM Tris/HCl (pH 6.5, 70 °C), subjected to ion exchange chromatography (UNO Q-17, Bio-Rad Laboratories, Hercules, CA) (pre-equilibrated in 20 mM Tris/HCl (pH 6.5, 70 °C), and eluted with a linear salt gradient from 0 to 1 M NaCl). Afterward, the sample was dialyzed overnight against 50 mM Tris/HCl, 300 mM KCl (pH 6.5, 70 °C) and applied to size exclusion chromatography (HiLoad 26/60 Superdex 200 prep grade; Amersham Biosciences) (pre-equilibrated in 50 mM Tris/HCl, 300 mM KCl (pH 6.5, 70 °C)). The fractions containing the recombinant enzymes were analyzed via Coomassie stained SDS-PAGE and enzyme activity tests, pooled and further used for the enzymatic characterization.

**Enzyme Assays—**Phosphatase activity was determined in a continuous enzyme assay at 70 °C with pNPP as substrate. For Saci-PTP the standard assay was performed in 0.1 M HEPES/KOH (pH 6.5) with 7 μg protein, whereas for Saci_0884 the assay was performed in 20 mM Tris/HCl, 300 mM KCl (pH 6.5), 5 mM Mn\(^{2+}\), 1 mM EGTA containing 0.25 μg protein in a reaction volume of 300 μl. Enzymatic activity was determined by monitoring the change in absorbance at 403 nm because of the formation of paranitrophenol (p-NPP = 18 mm/cm). For each assay three independent measurements were performed.

Protein tyrosine phosphatase activity was determined by using the tyrosine phosphatase assay system (Promega) following the manu-
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The artificial p-peptides TEVGKRI(pY)RLVGDKN, NIDAIRA(pS)LNIMSR, and ETTERERV(pT)TITORER were used as substrate. The enzyme activity was determined in a discontinuous enzyme assay at 70 °C in 0.1 n HEPES/KOH (pH 6.5) containing 2 μg Saci-PTP. The phosphate release was measured by using the Serine and Threonine phosphatase malachite-green-based assay system from Promega following the instructions of the manufacturer. For each assay three independent measurements were performed.

Protein serine/threonine phosphatase activity was determined by using the serine/threonine phosphatase assay system (Promega) following the manufacturer’s instruction. The artificial p-peptides RRA(pT)VA and TEVGKRI(pY)RLVGDKN were used as substrate. The enzyme activity was determined in a discontinuous enzyme assay at 70 °C in 20 mM Tris/HCl, 300 mM KCl (pH 6.5), 5 mM Mn2+ containing 0.1 μg Saci-PTP. The phosphate release was measured and visualized by using malachite green. For each assay, three independent measurements were performed.

For enzyme reactions following Michaelis-Menten kinetics the kinetic parameters (Vmax and Km) were calculated by iterative curve-fitting using the program Origin 8.6G 64 bit (Microcal Software, Northampton, MA).

Construction of In-frame Deletion Mutants in S. acidocaldarius—
Plasmids for deletion mutant construction were cloned using the PCR product of the upstream- and downstream-region of the gene of interest (Primers, supplemental Table S5), overlap PCR was performed to fuse both fragments and the overlap product was ligated with the gene targeting plasmid pSVA406 (27). The constructed plasmids were methylated using E. coli ER1821 and transformed into the background strain S. acidocaldarius MW001 (uracil auxotrophic strain) as described below (27). The subsequent steps were performed as described by Lassak et al. 2012 (29). In short, transformants were first selected on gelrite plates lacking uracil and then counterselected on plates with uracil and 5-FOA. The markerless in-frame deletion mutants of both phosphatase genes (saci0545 and saci0884), designated Δsaci_ptp and Δsaci_pp2a, respectively, were identified by colony PCR and confirmed by sequencing (MWG Operon, Ebersberg, Germany).

Flow Cytometry Analysis—Aliquots from growing cultures of strains MW001, Δsaci_ptp and Δsaci_pp2ac were transferred to ice-cold ethanol (final concentration 70% (v/v)). For DNA staining, ~0.5 ml of cell suspension in ethanol was centrifuged at 4 °C for 10 min at 16,000 × g. The pellet was resuspended in 1 ml of 10 mM Tris-buffer (pH 7.4) containing 10 mM MgCl2, centrifuged again, and then resuspended in 70 μl of the same buffer. Equal volumes (65 μl) of cell suspension and DNA-specific stain (0.2 mg/ml mithramycin A and 0.04 mg/ml ethidium bromide, also in Tris-MgCl2 buffer) were mixed. The samples were incubated on ice for 30 min before analysis in an Apogee A40 Analyzer flow cytometer equipped with a 405 nm solid-state laser.

Analysis of Cell Size—Microscopy pictures were taken using an Axio Imager M1 microscope (Zeiss) equipped with a Zeiss Plan Apochromat × 100/1.40 Oil DIC objective and a Cascade:1K CCD camera (Photometric). Cell size was analyzed automatically with ImageJ and ObjectJ using a modified version of the filaments-91.oj project. The results were evaluated with Origin 6.1 (OriginLab Corporation, Northampton, MA, USA).

RNA Isolation and Sample Preparation for RNA-seq Analysis—
S. acidocaldarius MW001, Δsaci_ptp, and Δsaci_pp2a were inoculated in triplicate. Cultures were grown to reach an OD600 of 0.6 (exponential growth phase) and subsequently samples of the three independent cultures of each strain were pooled in equal amounts to generate one mixed sample per strain. Total RNA samples were isolated from 10 ml of exponentially growing shaking cultures. TRIzol reagent (Invitrogen) was used for total RNA isolation following the instructions of the manufacturer. Residual chromosomal DNA present in RNA samples was removed by RNase-free DNase I (Roche) treatment for 2 h at 37 °C. DNA-free RNA samples were confirmed by PCR amplification using saci0574 (secY) primer pairs. Before cDNA synthesis DNA-free RNA samples were fragmented to achieve a molecule size range of 50 to 500 nucleotides. Thus, 6 μg of DNA-free RNA samples were mixed with 4 μl of 5X fragmentation buffer (200 mM Tris acetate, pH 8.2; 500 mM potassium acetate; 150 mM magnesium acetate). The reaction mix was filled up with DEPC H2O up to 20 μl, incubated at 95 °C for 2.5 min and immediately transferred to ice. The reaction was then cleaned up by using sephadex columns (Illustra MicroSpin™ G-25, GE Healthcare). RNA fragmentation range was checked by polyacrylamide gels. Six hundred nanograms of fragmented RNA was used for cDNA synthesis using the SuperScript Double-stranded cDNA Synthesis kit (Invitrogen) and following manufacturer’s instructions. The reaction was cleaned by using phenol/chloroform/isoamyl alcohol and cDNA samples were then precipitated by using 3 M NaOAc/100% ethanol and incubated over night at ~20 °C. Finally, cDNA samples were washed with 500 μl 70% ethanol. Five nanograms of cDNA were used as starting material for the generation of single-end sequencing libraries with the NEBnext DNA sample preparation kit, as described by manufacturer’s protocol. The DNA was ligated to Illumina adapters, which carried a unique four letter barcode sequence. DNA fragments of 150–500 bp were selected for sequencing. To retain strand specificity the DNA was treated with Uracil DNA-Glycosilase (UDGase) before the PCR amplification. The sequencing was performed by an Illumina Genome Analyzer IIX, multiplexed together with a total of eight samples. The reads were partitioned according to their barcode, stripped from adapter sequences and barcodes and mapped, using default settings, to the Sulfolobus acidocaldarius DSM 639 (NC_007181) reference genome, with segemei (30).

Bioinformatic Analysis of RNA-seq Data—For each protein coding and ncRNA gene, according to the NCBI database, the associated reads were counted. An overview where the reads map to, can be found in supplemental Fig. S8A. The overall Illumina sequencing represented 26.6%, 30.2%, 26.5% of genome coverage by the mapped reads for MW001, Δsaci_ptp and Δsaci_pp2a, respectively. To identify differentially expressed genes among the sample conditions, we used DESeq (version 1.5) (31), which allows estimation of the dispersion within one condition. This estimation is based on the assumption that only a few genes are truly differentially expressed; hence variance across different conditions can be seen as a too conservative estimation of the variance within one condition. From the estimated expected variance and the observed fold change in read counts, the statistical significance of altered RNA abundance can be calculated for every gene. Genes with a p value below 0.054 were considered to be significantly altered in their expression, supplemental Figs. S8B and S8C shows the correlation among the observed read counts, the fold change in read count among the sample conditions and the significance of the alteration.

RT-qPCR—First Strand cDNA Synthesis Kit (Fermentas) was used for cDNA synthesis according to the manufacturer’s instructions. The synthesis was thus performed using random hexamer primers and 1 μg of DNA-free RNA as template. Quantitative PCR (qPCR) analysis was carried out using Maxima SYBR Green/ROX qPCR Master Mix (Fermentas) based on the Sybr green detection system (Real-Time 7300 PCR machine; Applied Biosystems, Foster City, CA). The efficiency of each primer pair was calculated from the average slope of a linear regression curve, which resulted from qPCRs using a 10-fold dilution series (10 pg–10 ng) of S. acidocaldarius chromosomal DNA as template. Cq values (quantification cycle) were automatically determined by Real-Time 7300 PCR software (Applied Biosystems) after 40 cycles. Cq values of each transcript of interest were standardized to the Cq value of the reference gene saci0574 (secY) (32). qPCR
results with DNA-free RNA samples as template were performed as control. At least three biological replicates of each assessed condition and two technical replicates per qPCR reaction were performed.

**Immunoblotting Analysis After Nutrient-limited Growth Conditions**—The expression levels of the archaella components FlaB and FlaX were determined as described by Reimann et al. 2012 (33). Therefore the background strain S. acidocaldarius MW001 and both phosphatase deletion mutants (Δsaci_ptp and Δsaci_pp2a) were transferred to nutrient-limited medium. The same amount of cells referring to OD600 with and without starvation stress treatment were lysed and the proteins separated by SDS-PAGE. The expression levels of FlaB and FlaX were detected by immunoblotting using specific antibodies.

**Motility Assay on Semisolid Gelrite Plates**—To compare the motility of the background strain S. acidocaldarius MW001 to the phosphatase deletion strains (Δsaci_ptp and Δsaci_pp2a) the two control strains ΔaapF (hypermotile strain) and ΔaapFΔflaH (nonmotile strain) were used. The strains were spotted on semisolid gelrite plates (0.15% gelrite (w/v)) containing 0.005% (w/v) tryptone, 0.2% (w/v) dextrin and 10 μg/ml uracil as described before (29, 33). Plates were incubated for 5 days in a humid chamber at 75 °C. Swimming behavior of the different S. acidocaldarius strains was analyzed by measuring the swimming radius.

**Protein Sample Preparation for Mass Spectrometry Analysis**—For phosphoproteome analysis S. acidocaldarius MW001, MW010 (Δsaci_pptp), and MW025 (Δsaci_pp2a) were aerobically grown in a 10 liter fermenter with constant rotational mixing of the cultures. At the OD600 0.8 2 liter samples were taken. The cells were harvested by centrifugation at 9000 × g (Beckman Coulter Avanti J-26XP, rotor JLA 6.1) and the pellets immediately frozen in liquid nitrogen.

Cells were washed with cold water and then resuspended in 50 mM phosphate buffer (pH 7.5) and phosphatase inhibitors (5 mM each) comprising 2-glycerol phosphate, sodium fluoride, sodium vanadate, and sodium pyrophosphate (1). Cell extraction was performed using a combination of both ultrasonication and liquid nitrogen (manual grinding with a mortar and pestle). First, cells were sonicated using an ultrasonicator (Branson, Mexico) at a power of 70% for eight times (alternatively 45 s of sonication and 45 s of incubation in ice). Samples were then frozen using liquid nitrogen and ground manually five times. Furthermore, DNase I (100 μg/ml) and N-Octyl-glycoside (a final concentration of 1% (w/v)) were also added to samples to increase solubilization of membrane proteins (1). Samples were then finally sonicated in a water bath containing ice for 10 min before centrifugation at 21,000 × g for 20 min at 4 °C. Protein supernatant was collected and quantified using the RC-DC Protein Quantification Assay (Bio-Rad, UK). All chemicals were purchased from Sigma.

A total of 2 mg protein for each sample was denatured in 8 M urea, reduced with 10 mM DTT in 50 mM ammonium bicarbonate at 56 °C for 1 h, then alkylated with 55 mM iodoacetamide in 50 mM ammonium bicarbonate at 37 °C for 30 min in the dark. Before trypsin digestion was performed, samples were diluted to a final urea concentration less than 1 M by 40 mM ammonium bicarbonate in 9% acetonitrile (ACN) (34). A trypsin/protein ratio of 1:25 was used for trypsin digestion at 37 °C overnight. Samples were then dried in a vacuum concentrator (Eppendorf, Germany) before resuspension in buffer A consisting of 10 mM KH₂PO₄, 30% acetonitrile, and 0.1% trifluoroacetic acid pH 3.0 for fractionation and cleaning of the sample. Strong cation exchange (SCX) was performed using a HPLC system with UV detection (Dionex, UK) consisting of an analytical column (3 mm C₁₈, Dionex-LC Packings) operating at a flow rate of 300 nL/min. Buffer A contained 3% acetonitrile and 0.1% (w/v) formic acid, whereas buffer B contained 97% (w/v) acetonitrile and 0.1% (w/v) formic acid. The 90 min gradient comprised ramping from 3% B to 35% B in the first 70 min, then ramping to 90% B in 5 min, then switching back to 3% of buffer A for 15 min. Peptides eluting from HPLC column were directly submitted to the ion trap.

The PACIFIC technique was applied for analyzing (combined) cleaned peptides. Details of this technique can be found elsewhere (10, 37). Briefly, resuspended peptides were injected multiple times on the ion trap with collision induced association at each of 10 continuous 1.0 m/z intervals across a range of 10 m/z for each LC-MS analysis using a 2.0 m/z width and fragmented ions scanned from 100 to 1600 m/z with a dynamic ion charge threshold of 160,000. Subsequently, a next (forward) new ten ions was started in the same format as the first run, then this process continued until the considered precursors covered the range from 450 to 1100 m/z.

Data from mass spectrometry were then extracted to mgf format using Bruker Data Analysis V4.0 with a MRM script, these were then searched against the S. acidocaldarius database downloaded from NCBI in March 2010 (containing 2223 proteins) using Phenyx V 2.6 (Genebio, Geneva). The searches were performed using parameters as follows: carbamidomethylation of cysteine (fixed modification), oxidation of methionine (variation), and phosphorylation of serine, threonine, and tyrosine (variation), trypsin with two missed cleavages. Furthermore, other parameters such as parent, MS/MS, tolerances were set at 2.0 and 0.8 Da, respectively, whereas minimum peptide length, z-score, p value and AC score were set at 5, 5.5, 10–15, and 5.5 respectively. The spectra for all identified phosphopeptides were then automatically processed using Mathematica 9.0 (Wolfram Research, Inc.) in order orthogonal to confirm (1) the neutral loss of phosphoric acid (H₃PO₄, mass shift = −98) or metaphosphoric acid (HPO₃, mass shift = −80) from the precursor or/and fragment ions and (2) the presence of at least 5 b or y sequencing ion series. For the automatic annotation, peaks of intensity lower than 10% of the maximal intensity were discarded. The annotated spectra were then manually to ensure the quality of the annotations. Moreover, an independent biological duplicate sample was generated for each strain and an identical analysis carried out. Scores of phosphorylation sites are shown in Table S7. Raw spectra have been submitted to the PRIDE database (http://www.ebi.ac.uk/pride/, with ProteomeXchange accession: PXD000289, and are waiting for an accession number) (38).

**RESULTS**

**Characterization of saci_ptp and saci_pp2a Deletion Mutants**—Marker-less gene deletion mutants of the two putative phosphatases, Saci-PTP and Saci-PP2A, were generated using the uracil auxotrophic S. acidocaldarius MW001 strain (supplemental Fig. S1). Δsaci_pp2a showed slow growth (doubling time 9.5 h) and reached the stationary phase after 130 h, whereas MW001 and Δsaci_pptp showed similar growth characteristics (doubling times 5.7 h and 6.3 h, respectively; stationary phase reached after 60 h; supplemental Fig. S2). Notably, the Δsaci_pptp cells appeared normal in cell size (1.3 μm ± 0.55 μm), whereas the Δsaci_pp2a deletion mutant showed significant size variation (1.46 μm ± 1.08 μm) in com-
Compared to MW001 (1.27 μM ± 0.42 μM; supplemental Fig. S3) in microscopy analysis. The larger variation in the cell size distribution of the saci_pp2a deletion mutant, as compared with the parental strain and Δsaci_ptp, was confirmed by flow cytometry analysis (supplemental Fig. S4B). No changes in the DNA content distributions, reflecting the relative lengths of the cell cycle periods (39), could be detected in either deletion mutant, as compared with the parental strain (supplemental Fig. S4A), indicating no major effects on cell cycle regulation or progression through the different cell cycle phases. The aberrant phenotype of the Δsaci_pp2a mutant could be partially restored in a trans-complementation experiment, resulting in a growth curve and cell shape distribution similar to those of the parental strain (supplemental Figs. S5A and S5B).

**Table I**

| Substrate                      | Saci-PTP | Saci-PP2A |
|-------------------------------|----------|-----------|
|                               | $K_m$ (μM) | $V_{max}$ (U/mg) | $K_m$ (μM) | $V_{max}$ (U/mg) |
| pNPP                          | 530      | 3.7       | 2500      | 4.3        |
| TEVGKRI(pY)RLVGDKN            | 530      | 9.2       | No activity | No activity |
| RRA(pT)VA                     | -        | -         | 53.6      | 192.6      |
| NIDAIRA(pS)LNIMSR             | 423      | 0.3       | -         | -          |
| ETTYERW(pT)TITQRER            | 490      | 0.07      | -         | -          |

**Enzymatic Characterization of Saci-PTP and Saci-PP2A—**

Saci-PTP and Saci-PP2A were heterologously expressed (supplemental Fig. S6) in *E. coli*, purified, and analyzed in...
vitro. Both enzymes showed similar phosphatase activity with the pNPP substrate at 70 °C (Table I). For Saci-PTP substrate inhibition was observed at pNPP concentrations above 2 mM (k_inh-value of 0.17 (U/mg)mm⁻¹) (Fig. 1A). For Saci-PP2A cooperativity was observed at pNPP concentrations below 2 mM (Hill coefficient of 4.9) (Fig. 2A).

The substrate specificity of Saci-PTP was addressed with the p-Tyr peptide TEVGKRI(pY)RLVGDKN as substrate (Table I). In addition, low activities were observed with the p-Thr peptide ETTYERW(pT)TITQRER and the p-Ser peptide ETTYERW(pT)TITQRER, indicating dual protein phosphatase specificity (Table I; Fig. 1). However, the maximal velocity (V_max-value) for the p-Tyr peptide was 30- and 131-fold higher as compared with the p-Ser and the p-Thr peptide, respectively (Table I). The affinity (K_m value) toward all three phospho-peptides was similar (Table I), indicating that p-Tyr was the preferred substrate.

Phosphatase activity was only observed with the p-Thr peptide RRA(pT)VA for Saci-PP2A (Table I; Fig. 2). In addition, the analysis revealed that Saci-PP2A requires divalent metal ions for activity. The highest activity was observed with Cu^2+, followed by Mn^2+, Ni^2+, Mg^2+, Cd^2+ and Co^2+, in the presence of EGTA (supplemental Fig. S7). When EGTA was replaced by EDTA, no PP2A activity could be observed. In the absence of divalent metal ions the enzyme was inactive. Activity tests in the presence of okadaic acid, an inhibitor of PPPs, revealed a strong inhibitory effect (56% activity in presence of 10 nM okadaic acid; Fig. 2C).

**FIG. 2. Biochemical characterization of the protein phosphatases Saci-PP2A.** The activity of Saci-PP2A was determined in the presence of pNPP (A), the phospho-Thr peptide RRA(pT)VA (B) as well as in the presence of okadaic acid (p-Thr peptide RRA(pT)VA(C). All experiments were performed at 70 °C in triplicate. Saci_PP2A revealed phosphatase activity with pNPP as well as the p-Thr peptide, whereas no activity was observed with the p-Tyr peptide. Furthermore, Saci_PP2A was significantly inhibited in presence of okadaic acid, a typical inhibitor of PPPs.
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Fig. 3. Transcriptionally regulated genes in Δsaci_ptp and Δsaci_pp2a as compared with the strain MW001. A, The number of differentially regulated gene products in Δsaci_ptp and Δsaci_pp2a are depicted in a Venn diagram. Up and down arrows represent the number of genes which were found to be either up- or down-regulated for each deletion strain. In Δsaci_ptp, most genes were down-regulated, whereas in Δsaci_pp2a most were up-regulated. Eight genes were differentially regulated in both phosphatase deletion strains. B, Heat-map representation of all eight commonly regulated genes in Δsaci_ptp (X) and Δsaci_pp2a (Y). The graph is based on the differential gene expression analysis by RNA-seq, showing all genes that are significantly altered in their expression in both deletion strains. The right side panel shows the functional classification for each gene product.

obtained 1,835,618, 1,472,609, and 1,560,420 qualified Illumina read pairs for MW001, Δsaci_ptp and Δsaci_pp2a, respectively. An average of 82.6% of the reads were mapped to rRNA and tRNA sequences, whereas 14.5% mapped to protein-coding sequences (CDS) and 2.8% to intergenic regions in the S. acidocaldarius genome (supplemental Fig. S8).

A differential gene expression analysis was performed for each mutant strain, as compared with the MW001 parent strain. The transcript levels of 84 and 71 genes were significantly (threshold of twofold change) altered in Δsaci_ptp and Δsaci_pp2a, respectively (Fig. 3A and Fig. 4; supplemental Table S3A and S3B). Although the majority of the changes corresponded to down-regulated genes in Δsaci_ptp (52 of 84 genes), the opposite was observed in Δsaci_pp2a (56 of 71 genes up-regulated; Fig. 3A and Fig. 4). To functionally categorize the regulated genes, an arCOG category analysis was performed (40, 41). A majority of the differentially affected transcripts encoded proteins belonging to the categories unknown function (arCOG S), energy production and conversion (arCOG C), and transcription (arCOG K) (supplemental Table S4).

Only eight common genes were significantly affected in both deletion strains (Fig. 3B). Five of these were affected in the same direction (either up- or down-regulated), whereas three were divergently expressed (down-regulated in Δsaci_ptp whereas up-regulated in Δsaci_pp2a; Fig. 3B and Fig. 4). These three, saci1858, saci1859, and saci1860, corresponded to genes encoding SoxNL complex components (complex III of the respiratory chain; saci1858 and saci1859 encode mono-heme cytochrome b558–566 (CbsA and CbsB), whereas saci1860 encodes a Rieske Fe-S protein (SoxL)) (42). The two downstream genes of the soxNL complex operon, saci1861 (soxN) and saci1862 (odsN), were also up-regulated in Δsaci_pp2a (Fig. 4). Moreover, Δsaci_pp2a displayed augmented transcript levels of components of one of the three terminal oxidases (complex IV; the DoxBCE complex (saci0098 and saci0099); (Fig. 4). Four additional genes belonging to the energy metabolism and conversion arCOG category were up-regulated in Δsaci_pp2a, encoding a cytochrome/quinol oxidase (saci0097), a polyferredoxin (saci1802), a thioredoxin (saci1823), and the sulfocyanine SoxE-like protein (saci2096) (Fig. 4).

Increased transcript levels of three transcriptional regulators (saci1171, saci1180, and saci2193) were detected in Δsaci_pp2a. Interestingly, saci1180 and saci1171 are located up- and downstream of the archaellum operon (formerly archaecal flagellum (43)) (Fig. 4) and have been shown to activate archaella gene expression (44). Three additional genes of the archaellum operon, encoding FlaI, FlaG, and FlaX (saci1173, saci1176, and saci1177), were found to be up-regulated in Δsaci_pp2a cells, in agreement with a regulatory role for protein phosphorylation in archaella expression (33).

In contrast to Δsaci_pp2a, the Δsaci_ptp strain showed a predominant down-regulation of gene expression, including genes encoding components of the SoxABCD oxidase complex (saci2088 (soxB) and saci2086 (soxD)) (Fig. 4) (45). Additionally, transcript levels of ORFs encoding a thioredoxin-like (saci0359) protein, a Rieske-like protein (saci1963) and an isocitrate-lyase (saci2191) were down-regulated. To illustrate the transcription changes, a model of the S. acidocaldarius branched respiratory electron transport chain is depicted in Fig. 5 with the transcriptional response of each phosphatase mutant on the oxidase complexes indicated. Among the few up-regulated genes in Δsaci_ptp, three predicted transcriptional regulators were among the highest expressed genes (saci0065, saci0455, and saci1787). The target genes for these regulators are, unknown.

To confirm the RNA-seq data the expression of all archaellum operon genes, together with selected respiratory chain genes, was analyzed by quantitative RT-PCR. Whereas no significant changes in expression levels could be observed in the Δsaci_ptp deletion strain, all seven genes of the archaellum operon were found to be up-regulated in the Δsaci_pp2a cells (Fig. 6), confirming the result of the analysis (Fig. 4). The
induction of the genes encoding SoxNL complex components and the SoxABCDL terminal oxidase in H9004 saci_pp2a was also confirmed (Fig. 6).

Phosphoproteomic Analysis of MW001, H9004 saci_ptp and H9004 saci_pp2a—Phosphoproteomic analyses (including cytoplasmic and membrane proteins) were performed on the three strains (MW001, H9004 saci_pp2a and H9004 saci_ptp) in late exponential growth phase (OD600 nm of 0.8), as described previously for S. solfataricus P2 (10). Two biological replicates were analyzed for each strain and a false discovery rate (FDR) for each experiment was also estimated. As a result, averaged FDRs of 3.4%, 3.0%, and 3.6% were estimated for MW001, H9004 saci_ptp, and H9004 saci_pp2a, respectively.

Fig. 4. Bar plot representation of the transcriptomic changes in H9004 saci_ptp and H9004 saci_pp2a. A differential gene expression analysis was performed for each deletion strain, H9004 saci_ptp (A) and H9004 saci_pp2a (B), in comparison to the wild-type strain MW001. All genes of whose expression was significantly altered (>twofold) are shown. Red and green bars represent down- and up-regulated genes, respectively. Genes with a p value below 0.054 were considered to be significantly differentially regulated. The annotation of selected genes is indicated to the right of each column.
Overall, 1206 phosphorylated peptides (p-peptides) from 801 unique phosphorylated proteins (phosphoproteins) were identified, with an overall pS/pT/pY %-ratio of 35.6/28.1/36.2. In MW001, 54 phosphopeptides from 54 phosphoproteins (pS/pT/pY %-ratio of 32.9/21.3/45.7) were detected, in Δsaci_pp2a 477 phosphopeptides from 387 phosphoproteins (35.0/27.2/37.7), and in Δsaci_ptp 715 phosphopeptides from 551 phosphoproteins (36.4/29.1/34.5) (supplemental Table S1, Table II). The identified phosphoproteins were categorized according to arCOG functional code (40, 41). Unique phosphoproteins were identified in 21 out of 26 arCOG functional categories (supplemental Table S2), indicating an important role of phosphorylation in most cellular processes.

Fig. 5. Proposed model of *S. acidocaldarius* branched aerobic respiratory chain. The respiratory chain model is based on separate biochemical characterization studies. Electrons from NADH or succinate oxidation presumably enter electron transport chain via NADH: quinone oxidoreductases or succinate:quinone oxidoreductases (not shown). Reduced quinones (CQH2) then deliver electrons to cytochrome oxidase bc1 complex SoxLN-CbsAB-OdsN (42) and subsequently to one of the three terminal oxidase complexes. These terminal complexes are SoxABCD-SoxL (45, 54, 55), the bb3 terminal oxidase complex SoxEFGHIM (56–58), and the DoxBCE (59). Components whose transcript levels were altered in each deletion strain are depicted either in red or green for down- and up-regulation, respectively. Gene identifiers for each component are shown below.

Fig. 6. Differential expression levels of archaella operon genes and respiratory chain genes in Δsaci_pp2a and Δsaci_ptp. Total RNA isolated from *S. acidocaldarius* MW001, Δsaci_pp2a and Δsaci_ptp cultures were used for cDNAs synthesis. qRT-PCR analysis was performed using specific primers for each archaella component (A) and components of the terminal oxidase complexes (B). Relative transcript levels of each gene were normalized to an internal control gene secY. The values reflect the fold change compared with cDNA prepared from MW001. The means and standard deviations of three biological replicates are shown. Up-regulation of all archaella and the terminal oxidase genes in the saci_pp2a deletion strain reflect the RNA-seq results. * significant (p value ≤ 0.05), ** highly significant (p value ≤ 0.01).
Target Proteins for Phosphorylation—The phosphoproteome data set was specifically screened for proteins belonging to the aerobic respiration chain and the archaellum (Table III). Six proteins of the arCOG family C (energy production and conversion) were found to be phosphorylated, including SoxC (Saci2087) and SoxI (Saci2258), which belong to different terminal oxidase complexes (Fig. 5). Furthermore, three von Willebrand domain-containing proteins (Saci0977, Saci1209, and Saci1211) involved in archaellum synthesis were phosphorylated at different residues. One of these, ArnB (Saci1211), has previously been shown to be involved in negative regulation of the archaellum in vivo (33). Also the membrane-associated FlaJ protein (Saci1172), involved in archaella assembly, contained four phosphorylation sites and was identified in both phosphatase deletion mutants. The predicted positive transcriptional regulator ArnR1 (Saci1171)(44) located directly adjacent to the archaellum operon was also phosphorylated. Seventeen additional predicted transcriptional regulators also contained phosphorylation sites, demonstrating the importance of this PTM in transcriptional regulation in Archaea. Five predicted serine/threonine kinases were also identified as phosphorylated in the deletion strains.

Regulation of Cell Motility Via Reversible Protein Phosphorylation—The up-regulation of the seven archaella genes in /H9004 sacci_pp2a (Figs. 4 and 6), was investigated at the protein phosphorylation level. The parental strain and both deletion strains were cultivated in rich medium and then transferred to medium without supplemented nutrients for 5 h, a treatment that has been shown to induce archaella biosynthesis (29). Samples before and after induction were analyzed by immunoblotting with specific antibodies against the archaellin protein FlaB and the accessory protein FlaX. Both proteins were induced by nutrient limitation stress in the parent strain MW001, and to the same extent in /H9004 sacci_ptp. In contrast, /H9004 sacci_pp2a showed a FlaB signal already before induction, and highly increased levels of FlaB and FlaX after nutrient limitation as compared with MW001 and /H9004 sacci_ptp (Fig. 7A).

### Table II

| Organism               | Genome size (Mb) | Strain     | P-proteins (no.) | P-peptides (no.) | P-sites (no.) | pSer (%) | pThr (%) | pTyr (%) |
|------------------------|------------------|------------|------------------|------------------|---------------|----------|----------|----------|
| S. acidocaldarius Total| 2.2              | MW001     | 801              | 1246             | 1855          | 35.6     | 28.1     | 36.2     |
| S. acidocaldarius      |                  | Δsacci_pp2a| 387              | 477              | 734           | 35.0     | 27.2     | 37.7     |
|                        |                  | Δsacci_ptp | 551              | 715              | 1044          | 36.4     | 29.1     | 34.5     |
| S. solfataricus Total  | 2.9              | P2        | 540              | 690              | 1318          | 25.8     | 20.6     | 53.6     |
| S. solfataricus        |                  | P2        | 311              | 343              | 580           | 25.2     | 18.5     | 55.3     |
|                        |                  |           | 311<sub>Tryp</sub>| 384<sub>Tryp</sub>| 810<sub>Tryp</sub> | 25.1<sub>Tryp</sub> | 21.5<sub>Tryp</sub> | 53.4<sub>Tryp</sub> |
| H. salinarum (2)       | 2.7              | R1        | 69               | 90               | 81            | 86       | 12       | 1        |
| Lactococcus lactis (60)| 2.4              | III403    | 63               | 102              | 73            | 46.5     | 50.6     | 2.7      |
| Bacillus subtilis (1)  | 4.2              | 168       | 78               | 103              | 78            | 69.2     | 20.5     | 10.3     |
| Escherichia coli K12(61)| 4.6            | K12-MG1655| 79               | 105              | 81            | 67.9     | 23.5     | 8.6      |
| Zea mays leaf (62)    | 2.065            | B73       | 125              | 149              | 157           | 89.8     | 9.6      | 0.6      |
| Synechococcus sp. (63) | 3.4              | PCC 7002  | 245              | 280              | 410           | 43.9     | 42.4     | 13.6     |
| Pseudomonas putida(51) | 5.9              | PNL-K25   | 40               | 56               | 53            | 52.8     | 39.6     | 7.5      |
| Pseudomonas aeruginosa| 6.3              | PAO1      | 23               | 57               | 55            | 52.7     | 32.7     | 14.5     |
| Campylobacter jejuni (64)| 1.6            | NCTC11168| 36               | 58               | 35            | -        | -        | -        |
| Streptococcus pneumonia (65)| 2.1          | D39       | 84               | 102              | 163           | 47.2     | 43.8     | 9        |
| Streptomyces coelicolor (66) | 9.1        | A3 (2)    | 40               | 44               | 44            | 34.1     | 52.3     | 13.6     |
| Streptomyces coelicolor (67)| 9.1        | M145      | 127              | 260              | 289           | 46.8     | 48.3     | 5.2      |
| Klebsiella pneumonia (68)| 5.5          | NTHU-K2044| 81               | 117              | 93            | 31.2     | 15.1     | 25.8     |
| Mycoplasma pneumoniae (69)| 0.8         | M129      | 63               | 16               | 16            | 53.3     | 46.7     | 0        |
| Helibacter pylori (70)| 1.7              | 26695    | 67               | 80               | 126           | 42.8     | 38.7     | 18.5     |
| Mycobacterium tuberculosis (71)| 4.4        | H37Rv     | 301              | 381              | 516           | 60       | 40       | 0        |
| Listeria monocytogenes (72)| 2.9          | EGD<e>   | 112              | 155              | 143           | 65       | 30.1     | 4.9      |
| Clostridium acetobutylicum (73)| 23.3       | ATCC824  | 61               | 82               | 107           | 42       | 47.7     | 10.3     |
| Plasmidium falciparum (74)| 176.4        | ATCC30236| 82               | 93               | 103           | 71.8     | 21.4     | 6.8      |
| Thermus thermophilus (76)| 2.1           | HB8      | 48               | 52               | 50            | 64       | 26       | 10       |
| Rhodopseudomonas palustris (77)| 5.5          | CGA010   | 54<sub>CH</sub>  | 100<sub>CH</sub>| 60<sub>CH</sub>| 63.3<sub>CH</sub> | 16.1<sub>CH</sub> | 19.4<sub>CH</sub> |
| Trypanosoma cruzi (78)| 67               | DM28c    | 755              | -                | 2572          | 84.1     | 14.9     | 1.0      |
| Toxoplasma gondii (79)| 80               | RH       | 2793<sup>i</sup>| 11,822<sup>i</sup>| 12,793<sup>i</sup>| 87.2     | 12.6     | 0.25     |
| Plasmodium falciparum (79)| 23            | 3D7      | 1673             | 7835             | 8463          | 89.1     | 10.4     | 0.51     |
| Arabidopsis thaliana (80)| 119.7         | -        | 598              | 962              | 609           | 86       | 14       | 0.16     |

Crenarchaeal Signal Transduction
Crenarchaeal Signal Transduction

The swimming ability of the wild-type strain and both deletion mutants was investigated using a motility assay. MW001, Δsaci_pp2a, Δsaci_pp2a, a hypermotile positive control (Δsaci_pp2a) and a nonmotile negative control (ΔaapFΔflaH) were spotted on semi-solid Gelrite-plates containing low amounts of nutrients (29, 46). After 5 days of incubation at 76 °C, MW001 Δsaci_pp2a showed only slight motility. In contrast, the motility of Δsaci_pp2a was comparable to the hypermotile ΔaapF, which lacks the Aap pili (Archaeal adhesive pili) (Fig. 7B), indicating a major impact of protein phosphorylation on cell motility in *S. acidocaldarius*.

**DISCUSSION**

We report a detailed characterization of two protein phosphatases of *S. acidocaldarius*, Saci-PTP and Saci-PP2A. Saci-PTP displayed dual-specific Ser/Thr and Tyr phosphatase activity, whereas Saci-PP2A was specific for Ser/Thr dephosphorylation. Differences in phosphorylation and gene expression patterns were observed in deletion strains, suggesting roles for both phosphatases in signal transduction pathways. The changes observed when the gene encoding Saci-PP2A was deleted were similar to responses observed when *S. acidocaldarius* cultures experience severe nutrient limitation. Up-regulation of archaellar and respiratory chain components may enable the organism to evade unfavorable conditions by promoting archaellar assembly and providing energy for archaella-driven motility.

Comparison of the phosphoproteome of *S. acidocaldarius* with that of *S. solfataricus* (40) revealed an overlap of 94 proteins, although the different experimental approaches (deletion mutants compared with growth on different carbon sources) should be taken into account. Conserved phosphorylation sites were identified in 13 of the 94 overlapping proteins. A single conserved site was identified in ten of these, whereas two sites were detected in Saci0940, Saci0381, and Saci2134. None of the 15 sites had a score higher than 0.5.
Crenarchaeal Signal Transduction

The phosphoproteome of the hyperthermophilic bacterium *Thermus thermophilus* was recently analyzed, revealing 48 phosphoproteins with 50 phosphorylation sites and a pSer/pThr/pTyr %-ratio of 64/26/10. The amount of pTyr should be the same. Nonetheless, the enzymatic properties of SerB were not investigated until now, the data imply that the protein is not specific for pSer.

Bioinformatic analysis revealed that SerB shares no similarity with Saci-PTP or Saci-PP2A. For *S. acidocaldarius*, the unexpected change of the pSer/pThr/pTyr %-ratio could partly be explained by the biochemical properties of Saci-PTP and Saci-PP2a, as the Saci-PTP displays dual activity. It is also possible that deletion of the protein phosphatases induces other PTMs to compensate for the loss of regulation.

The high amount of Tyr phosphorylation in both *Sulfolobus* species appears to be a crenarchaeal feature, rather than a thermoadaptation property or a general archaeal feature. The phosphoproteome of the hyperthermophilic bacterium *Thermus thermophilus* was recently analyzed, revealing 48 phosphoproteins with 50 phosphorylation sites and a pSer/pThr/pTyr %-ratio of 64/26/10. The amount of pTyr is in accordance with amounts identified in other prokaryotic phosphoproteome studies (Table II), supporting the conclusion that Tyr phosphorylation is a crenarchaeal feature rather than a thermoadaptation. Both *T. thermophilus* and *H. salinarum* contain two-component phosphorylation systems, which are absent in crenarchaeas. It is thus possible that the extensive Tyr phosphorylation in *Sulfolobus* species provides a means to compensate for the absence of two-component systems in these organisms.

The expression of several transcription factors was highly affected in the deletion mutants. However, in the absence of information about target genes, the physiological consequences remain unknown. In contrast, an explanation can be provided for the effects of deletion of Saci-PP2A on archaellum and respiration chain genes. In *S. acidocaldarius* two regulatory systems have been shown to be involved in control of archaella biosynthesis, the ArnR (Saci1180) and ArnR1 (Saci1171) positive regulators...
of flaB expression (See model in Fig. 8) (44). ArnR1 and ArnR, were up-regulated in the Δsaci_pp2a strain and, in addition, ArnR1 was found to be phosphorylated in the Δsaci_pp2a strain. Further, in the biochemical characterization, ArnA and ArnB were phosphorylated by the Saci1193 kinase and dephosphorylated by the Saci-PP2A phosphatase (35). Probably another, yet unknown, factor is repressing the flaB promoter. When S. acidocaldarius cells enter stationary phase and undergo starvation, or in the Δsaci_pp2a deletion mutant (B), the ArnA/B complex exists only in its phosphorylated form and repression is released. Moreover, the expression of ArnR (Saci1180) and ArnR1 (Saci1171) is induced and these factors then drive high expression of FlaB which leads to the assembly of archaea on the cell surface and therefore to increased motility.

Thioredoxin (Saci1823), thioredoxin reductase (Saci1169) and peroxiredoxin (Saci2227) encoding genes were up-regulated in the Δsaci_pp2a strain. These were also up-regulated in a study on oxidative stress in S. solfataricus, but no archaea gene up-regulation was found (49). In all three domains of life, these proteins are involved in disulfide bond shuffling during oxidative stress conditions. The thioredoxin reductase gene is located immediately adjacent to the archaea genes and the ArnR1 regulator and the protein was, in addition, found to be phosphorylated. Hence, nutrient limitation leads both to archaea expression and to oxidative stress responses to avoid cellular damage. In summary, the ArnA/ArnB and ArnR/R1 regulatory systems of archaea expression constitute the first complex regulatory network involving protein phosphorylation to be investigated in crenarchaea.

Surface structure-driven motility is an energy-consuming process and archaea assembly therefore has to be tightly controlled. Several genes involved in energy conversion, including those encoding the SoxNL, SoxABCDL, and DoxBCE respiratory chain complexes, were differentially regulated in the phosphatase deletion strains as compared with the parental strain (Fig. 4). Recently, an association of the flagellar switch-motor complex to the NADH-ubiquinone oxidoreductase and the F$_{0}$F$_{1}$ ATP synthase was demonstrated in the bacterium Escherichia coli, and predicted to function as a local energy supply for flagellar movement (50). The up-regulation of archaea genes in S. acidocaldarius may similarly be connected to the up-regulation of genes involved in the res-
piratory chain. Possible interactions of energy generating complexes with the archaea l basal body should therefore be investigated. Phosphorylation of respiratory chain components has also been reported in phosphoproteomic studies of *Pseudomonas* spp (51), *Saccharomyces cerevisiae* mitochondria (52), human muscle mitochondria (53) and *S. solfataricus* (10).

This study provides the first insights into the *in vitro* and *in vivo* functions of the two *S. acidocaldarius* protein phosphatases, Saci-PTP and Saci-PP2A. The RNA-seq and phosphoproteomics studies of the deletion mutants shed light on the multitude of cellular processes affected by reversible protein phosphorylation. In particular, we demonstrate that SaciPP2A is involved in the complex signal transduction pathway regulating the expression of the *S. acidocaldarius* archaellum, and also plays a major role in the control of aerobic respiration and oxidative stress responses.

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