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2 3 4 5 6	Natronocalculus amylovorans gen. nov., sp. nov., and Natranaeroarchaeum aerophilus sp. nov., dominant culturable amylolytic natronoarchaea from hypersaline soda lakes in southwestern Siberia.
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18 19 20 21 22 23 24 25 26 27 28 29	Running title: <i>Natronocalculus amylovorans</i> gen. nov., sp. nov. and <i>Natranaeroarchaeum aerophilus</i> sp. nov.
30	The draft genome sequences of strains AArc-St1-1 <sup>T</sup> and AArc-St2 <sup>T</sup> are deposited in the GenBank
31	under the numbers JAKRVY000000000 and JAKRVX000000000, respectively

#### Abstract

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Several pure cultures of alkaliphilic haloaloarchaea were enriched and isolated from hypersaline soda lakes in southwestern Siberia using amylopectin and fructans as substrates. Phylogenomic analysis placed the isolates into two distinct groups within the class *Halobacteria*. Four isolates forming group 1 were closely related to a recently described Natranaeroarchaeum sulfidigenes and the other three strains forming group 2 represent a novel genus-level phylogenetic lineage. All isolates are saccharolytic archaea growing with various starch-like alpha-glucans including soluble starch, amylopectin, dextrin, glycogen, pullulane and cyclodextrin. In addition, group 1 can also use levan while group 2 - inulin (plant storage beta-fructans). Group 1 strains can also grow anaerobically with either glucose or maltose using elemental sulfur as the electron acceptor. Both groups are moderately alkaliphilic with a pH range for growth from 7.2 to 9.3 (optimum between 8.0-8.8) and low Mg-demanding extreme halophiles growing optimally at 4 M total Na<sup>+</sup>. The major respiratory menaquinone is MK-8:8 and the core biphytanyl lipids are dominated by archaeol (C<sub>20</sub>-C<sub>20</sub>) and a less abundant extended archaeol (C<sub>20</sub>-C<sub>25</sub>) with PG and PGP-Me as polar groups. The four isolates of group 1 are suggested to be classified into a new species as Natranaeroarchaeum aerophilus sp. nov. (type strain AArc-St1-1<sup>T</sup>=JCM 32519<sup>T</sup>=UQM 41458<sup>T</sup>). The three isolates of group 2 are proposed to form a new genus and species for which the name Natronocalculus amylovorans gen. nov., sp. nov. is suggested (type strain AArc-St2<sup>T</sup>=JCM 32475<sup>T</sup>=UQM 41459<sup>T</sup>).

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Key words: hypersaline soda lakes, natronoarchaea, amylolytic, starch, fructans,

53 Natranaeroarchaeum, Halobacteria

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#### Introduction

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Most of the known species of aerobic extremely halophilic and haloalkaliphilic euryarchaea (natronoarchaea) were enriched and isolated from hypersaline alkaline lakes on unspecific media containing rich soluble organic substrates, such as peptone, yeast extract or simple sugars [1-5]. Yet, these extremophilic, organoheterotrophic archaea are definitely more important in organic matter mineralization, in particular organic polymers, in hypersaline systems than is widely recognized. For example, a test for amylolytic activity with soluble starch is included into the minimal standards for taxonomy characterization of haloarchaea [6] but starch is rarely used for targeted isolation of amylolytic haloarchaea which would not only hydrolyze the polymer but also able to utilize it as growth substrate. And this situation is also true for other polysaccharides. So far, only a few examples of haloarchaea specialized on utilization of recalcitrant polysaccharides have been described in pure culture, which is particularly true for natronoarchaea living in alkaline hypersaline (soda) lakes. Our recent targeted enrichments from such lakes using insoluble forms of cellulose and chitin resulted in isolation of several groups of natronoarchaea highly specialized on utilization of either various forms of cellulose and xylan (genera Natronobiforma and Natronolimnobius) or chitin (genus Natrarchaeobius) [7-9]. Following further in this direction, we used insoluble starch (amylopectin) or beta-fructans as substrates to enrich for amylolytic or fructanolytic natronoarchaea from hypersaline soda lakes. So far, we are aware only of a single amylolytic natronarchaeon, Natronococcus amylolyticus, specifically isolated from the hypersaline soda lake Magadi in Kenya using starch as a growth substrate [10-11], while no natronoarchaea, growing on fructans are currently known.

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Here we describe phenotypic, phylogenetic and genomic properties of two novel taxa of amylolytic natronoarchaea enriched from hypersaline soda lakes in southwestern Siberia, which specialized on utilizing various alpha-glucan and beta-fructan polysaccharides as growth substrates.

#### **Materials and Methods**

Enrichment and cultivation conditions

The sources of inocula were mixed surface (0-3 cm deep) aerated sediments and the near bottom brines obtained from four hypersaline soda lakes in Kulunda Steppe (Tanatar and Bitter lake systems, Altai region). The lakes brines total salt concentration varied from 20 to 40%, the total soluble carbonate alkalinity - from 2.5 to 5 M and the pH - from 10.2 to 11.0. The top flocculant sediment layer together with the near bottom brines was sucked into 50 ml syringe through silicon tubing and placed into a sterile 50 ml Falcon tube, resulting in an approximate volumetric ratio between the solid and liquid fractions of 1:5. After transportation to the laboratory the samples were separated onto clean brine top layer and concentrated sediment, all of which then were kept at 4°C. Before inoculation, the two fractions from each of four lakes were combined in equal proportions to make two master mixes (brine and sediments) used as inocula at 1% final (v/v).

For the enrichment and further cultivation of pure cultures two basic mineral media were used, both containing 4 M total Na<sup>+</sup>. The neutral 4 M NaCl base medium included (g l<sup>-1</sup>): NaCl, 240; K<sub>2</sub>HPO<sub>4</sub> 2.5 g l<sup>-1</sup>; NH<sub>4</sub>Cl 0.5 g l<sup>-1</sup>, KCl 5 g l<sup>-1</sup>, 20 mg l<sup>-1</sup> yeast extract and was adjusted to pH 7 with 10% KH<sub>2</sub>PO<sub>4</sub>. The alkaline sodium carbonate base contained (g l<sup>-1</sup>): Na<sub>2</sub>CO<sub>3</sub> 190, NaHCO<sub>3</sub> 45, NaCl 16, KCl 5, K<sub>2</sub>HPO<sub>4</sub> 1 and 20 mg l<sup>-1</sup> yeast extract (pH 10). After sterilization, the basic media were supplemented with 1 ml l<sup>-1</sup> of trace metal and vitamin solutions [12] and 1 mM MgSO<sub>4</sub>. NH<sub>4</sub>Cl (4 mM) was also added from 1 M sterile stock solution to the carbonate base after sterilization. To prepare final medium with a certain pH/alkalinity, the two ready to use bases were mixed in different proportions resulting in pH range from 8 to 10. For the enrichments from salt lakes, the neutral base medium was used as it is, while for the soda lake enrichments the neutral and alkali base media were mixed 3:1 with the final pH 9.5. For the pH range from 7 to 8, the NaCl base was supplied with various amounts of 1 M filter-sterilized NaHCO<sub>3</sub>, while for the pH below 7

it was titrated with sterile 1 M KH<sub>2</sub>PO<sub>4</sub>. Carbon and energy substrtates were added from sterile 10% stock solution.

For isolation of pure cultures, several rounds of enrichments were repeated at dilutions 1:100 and finally the sediment-free enrichments were plated onto solid medium obtained by mixing the liquid alkaline medium and 4% washed agar at 50°C in ratio 3:2. To compensate for the decreased salinity, solid NaCl was added to the liquid medium before mixing with melted agar to bring the final salt concentration back to 4 M total Na<sup>+</sup>. The isolation of pure cultures was achieved from separate colonies which grew back in liquid medium with the target polysaccharide and the purity was confirmed by 16S rRNA gene and genome sequencing.

### Pure culture characterization

Cell morphology was examined by using phase contrast microscopy (Zeiss Axioplane Imaging 2, Germany). Substrate utilization profiles were performed in medium containing 1 part of the alkaline base and 7 parts of the NaCl base (final pH 9.0). For the pH profiling, the two media were mixed in various proportions as described above and soluble starch served as the substrate. The growth was measured by increase of OD<sub>600</sub> with pH monitoring at each point. Anaerobic cultivation was performed as described previously [13]. Catalase and oxidase activity were tested with 3% (v/v) H<sub>2</sub>O<sub>2</sub> and 0.1% *N,N,N,N* tetramethyl-p-phenylenediamine hydrochloride, respectively, using cell-free extract (obtained by sonication) from cells of type strains. The protease, esterase/lipase activities were tested on plates spotted with fully grown liquid cultures: using casein/gelatin (hydrolyzis zones after flooding with 10% TCA) and emulsified tributyrin/olive oil (turbidity clearance), respectively. Antibiotic sensitivity of type strains AArc-St1 and AArc-St2 was tested in liquid medium at pH 9 with starch as substrate.

The intact polar lipids (IPLs) and respiratory quinones were analyzed as described previously [14]. Briefly, the lipid fraction was extracted from freeze-dried cells with sonication in

methanol:dichloromethane:phosphate buffer (2:1:0.8, v:v), followed by phase separation by adjusting the solvent mix to a ratio 1:1:0.9. The lipids and quinones were analyzed by normal phase, high performance liquid chromatography-ion trap mass spectrometry (HPLC-ITMS) and identified by masses and mass spectral fragmentation according to literature [14-15].

#### Genome sequencing

Genomic DNA from the type strains AArc-St1-1 and AArc-St2 was extracted using DNeasy PowerLyzer Microbial Kit (Qiagen) according to manufacturer instructions. Quality and quantity of the DNA samples were measured with Trinean Xpose spectrophotometer (PLT Scientific Instruments) and Qubit 2.0 fluorometer (Thermo Fisher Scientific) DNA libraries were prepared using KAPA HyperPlus kit (KAPAbiosystems) according to manufacturer recommendations. Paired-end sequencing (2x100bp) was performed using Illumina NextSeq. Obtained reads were filtered (quality and length) with CLC Genomics Workbench v.10. Genomes of the strains were assembled using SPAdes v.3.15.2 [16] --isolate mode with --trusted-contigs option (contigs obtained from Unicycler v.0.4.9 [17] were used as trusted contigs). Contigs with length  $\leq$  500bp or with low coverage were eliminated. Genome assemblies statistics were checked with Quast v.5.0.2 [18-19]. Completeness and contamination levels were detected using CheckM v.1.1.2 [20] with archaea-specific marker set.

### Phylogenetic and genomic analyses

For 16S rRNA gene sequence-based phylogenetic analysis 16S rRNA gene sequences of the seven isolates were aligned with the sequences of type species of all genera within *Halobacteria* (as well as *Archaeoglobus fulgidus* VC-16, *Methanocella paludicola* SANAE, *Methanothermobacter thermautotrophicus* Delta H used as the outgroup). Multiple sequence alignment and phylogenetic tree construction were performed as described earlier [9]. For phylogenomic analysis based on the

"ar122" set of conserved single copy archaeal proteins [21] the protein sequences were identified and aligned in *in silico* proteomes of type species of all genera within *Halobacteria* (nontype species were taken for *Halalkalicoccus*, *Halorbellus*, *Natronoarchaeum* and *Halohasta* genera because the genomes of type spices are not available) using the GTDB-tk v.1.7.0 with reference data v.202 [22]. The phylogenomic tree was constructed in the RAxML v.8.2.12 [23] with the PROTGAMMAILG model of amino acid substitution; local support values were 1000 rapid bootstrap replications. Phylogenetic trees were visualized using iTOL v.6.5.2 [24]. The whole genome-based comparisons were done as described by Sorokin et al. [25].

For functional genome analysis, genes encoding carbohydrate-active enzymes (CAZymes) were searched in genomes of AArc-St2 and AArc-St1-1 strains using dbCAN v.3.0.2 [26]. Further manual checking of the specificity of discovered glycosidases and other CAZymes were performed using BLAST against Swiss-Prot/PDB database.

# **Results and discussion**

Enrichment and isolation of pure cultures

The primary enrichments for amylolytic and fructan-utilizing natronoarchaea were performed with amylopectin (insoluble starch) or levan/inulin, respectively, in the presence of 200 mg 1<sup>-1</sup> streptomycin to suppress growth of bacteria. The enrichments from sediment fraction showed visible growth (after removal of the sediment particles by low speed centrifugation) after one week of incubation, while the brine enrichments became turbid and pinkish after 10-14 days of incubation. After several 1:100 transfers, the liquid cultures were plated and individual pink-colored colonies were transferred back to the corresponding liquid media. This procedure was repeated 2 more times to ensure the homogeneity of colony morphology. The purity and identity of

obtained cultures were verified by the 16S rRNA gene sequencing. The list of isolates is shown in **Table 1**.

Cell morphology of the isolates grown at pH 9 and 4 M total Na<sup>+</sup> with soluble starch is shown on **Fig. 1**. Cultures of all strains were dominanated by nonmotile (with only a few occasionally showing slow motility) flattish cocci with irregular contour, typical for haloarchaea, with a small fraction of flat roads ("boards"). The colonies of all isolates were colored from orange to red and concentrated cell pellets were bright red, what is also typical for aerobic haloarchaea.

# Chemotaxonomy

Membrane polar lipids and respiratory lipoquinones were analyzed in two strains, one of which represents a group 1 (AArc-St1-1<sup>T</sup>), while another a group 2 (AArc-St2<sup>T</sup>). In both, the core lipids were dominated by archaeol (AR; C<sub>20</sub>-C<sub>20</sub>) with a smaller proportion of extended archaeol (Ext-AR; C<sub>20</sub>-C<sub>25</sub>). The polar head groups of the intact polar lipids were phosphatidylglycerophosphate methylether (PGP-Me) and phosphatidylglycerol (PG). Both the core lipids and the polar head groups are very similar to the closest phylogenetic relatives of the amylolytic natronoarchaea (see below in comparative tables). The major difference is the absence of glycolipids and sulfolipids (such as phosphatidylglycerosulfate and sulfated glycosyl diethers) in natronoarchaea, which are more common in neutrophilic haloarchaea. The only respiratory lipoquinone species detected in both strains was the fully saturated MK-8:8, one of the most common in haloarchaea [15].

### Phylogenetic and genomic analyses

The genome of AArc-St2 was assembled to 20 scaffolds including one circular plasmid, while the genome of AArct-St1-1 was assembled into 32 scaffolds with no plasmids in it (Suppl. **Table S1**). Genome size was 3.26 Mbp (GC content was 51.5%) for strain AArc-St2 and 3.29 Mbp (GC content was 61%) for strain AArc-St1-1. Completeness and contamination levels for AArc-St2

genome were 100% / 0% and for AArc-St1-1 – 99.07% / 1.87%. Genomic sequences are available in NCBI GenBank database with accession numbers JAKRVX000000000 (AArc-St2) and JAKRVY000000000 (AArc-St1-1).

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Seven natronoarchaeal isolates formed two clusters on 16S rRNA gene sequence-based phylogenetic tree. Four strains of the group 1 clustered with the recently described facultatively anaerobic sulfur-reducing natronarchaeon Natranaeroarchaeum sulfidigenes which can also use starch as a substrate for growth either aerobically or anaerobically [27-28]. The three closely related isolates from the group 2 formed a novel genus-level lineage with the genera "Halalkalirubrum" and Halohasta as the nearest neighbors (around 93 and 92 % sequence identity to its type species, respectively). This potentially new genus lineage also includes multiple uncharacterized isolates from various hypersaline habitats whose 16S rRNA gene sequences were deposited recently in the GenBank. However, the 16S rRNA gene sequence-based phylogenetic tree (Fig. 2a) had relatively low bootstrap support of its inner nods within the class *Halobacteria*. Phylogenomic tree based on 122 archaeal conserved single-copy protein markers of the strains AArc-St1-1 and AArc-St2 and other haloarchaea supported the branching, obtained 16S rRNA gene sequence-based tree but showed a better resolution of the inner nods (Fig. 2b). Strain AArct-St2 formed a novel-genus branch in a distinct cluster containing genera "Halalkalirubrum", Halohasta and Halonotius. The latter three genera are currently classified in the order Haloferacales, family Halorubraceae [29]. Strain AArc-St1-1 was closely related to Natranaeroarchaeum sulfidigenum AArc-S and formed a potential new species in the genus which is currently classified within the family Natranoarchaeacea [28]. Strains AArct-St2 and AArc-St1-1 were proposed to be type species within their lineages.

For additional support of phylogenetic and phylogenomic analyses, ANI and AAI values were calculated for the genomes of AArc-St1-1 and AArc-St2 and the nearest relatives (Suppl. **Table S2 and S3**). The AAI values between strain AArc-St2 and the species of three related

haloarchaeal genera with available genome sequences ranged from 62.7% to 70.1%. These values are below the level of AAI for the representatives of the majority of genera, for which AAI results were compared [30], but it is similar to the intergenera level within the whole cluster. The ANI values varied from 70.7% to 73%. Same calculations for strain AArc-St1-1 and *Natranaeroarchaeum sulfidigenum* AArc-S showed AAI and ANI values of 90.1% and 88.8%, respectively, confirming the separate species status of AArc-St1-1 within the genus *Natranaeroarchaeum*. Although, according to the 16S rRNA gene sequence phylogeny, strain AArc-St1-2 might be sufficiently distant from the other three members of the group 1 and the type strain AArc-S<sup>T</sup> of *Natranaeroarchaeum sulfidigenes* (98.6-98.8 and 98.5%, respectively), the proposed type species of the genus *Natranaeroarchaeum*, its phenotypic properties were quite similar to the other group 1 strains implying it would be more practical to classify all four isolates of this group in a single species.

# Metabolic properties

All isolates were capable of utilizing alpha-1,4/1,6 glucans as growth substrates, including soluble starch, amylopectin (insoluble starch), glycogen, dextrin, cyclodextrin and pullulan. Furthermore, the group 1 isolates can also grow with levan (polyfructose with beta-2,6 backbond) while the group 2 strains utilized another fructan - inulin (polyfructose with beta-2,1 backbond). Two other alpha-bonded polysaccharides tested, including dextran from *Leuconostoc* and arabinan were not utilized by any of the seven isolates, as well as various beta-glucans (amorphous cellulose, xylan, xyloglucan, chitin, mannan, glucomannan, galactomannan, lichenan, laminarin and galactan). All strains can also grow on three sugar dimers including maltose,  $\alpha$ , $\alpha$ -trehalose and cellobiose. In addition, the group 2 strains were able to grow with glycerol. None of the other tested substrates, except for a weak growth with mannose, gave positive results (glucose, sucrose, galactose, arabinose, rhamnose, raffinose, aminosugars, uronic acids, xylose and arabinose, sugar alcohols,

C2-C6 organic acids, pepton). Such limited substrate profile characterize the isolates as narrow-specialized saccharolytics. Anaerobic fermentative growth with either starch, maltose or arginine was not observed. None of the isolates were capable of anaerobic respiratory growth with soluble starch or maltose as substrates, using thiosulfate, DMSO, fumarate or nitrate as the electron acceptors. However, all four group 1 isolates grew anaerobically with maltose as the electron donor and carbon source and elemental sulfur as the electron acceptor, similar to the closely related type species of the genus *Natranaeroarchaeum*. In 15 d incubation (4 M Na<sup>+</sup>, pH 9, 30°C) the following amount of sulfide was produced: 15.6 mM by AArc-St1-1, 5.8 mM by AArc-St1-2; 19 mM by AArc-St1-3 and 10.5 mM by AArc-St-lev1. In comparison, *Natranaeroarchaeum sulfidigenes* formed 30 mM sulfide in 6 days of cultivation at pH 9.5.

Two type strains tested positive for catalase reaction and TMPD-oxidase. The protease, esterase and lipase activities were negative in all strains in the spot-plate tests. Ammonium and yeast extract (but not nitrate) can serve as the N-source in cultures grown with soluble starch for both strains. Urea was only utilized by strain AArc-St1-1 consistent with the presence of the *ureABCDEFG* urease operon in the genome. Indole formation from tryptophan (Kovac's reagent test) showed a weak positive result only for strain AArc-St2. The type strains grown in liquid culture at pH 9 with soluble starch were insensitive to streptomycin, penicillin G, ampicillin, kanamycin, vancomycin and gentamicin up to 200 mg l<sup>-1</sup>. Rifampicin and chloramphenicol inhibited growth at 50 mg l<sup>-1</sup>, and tetracyclin - at 100 mg l<sup>-1</sup>.

All isolates grew well at as low Mg concentration as 1 mM, while in their sodium requirement they are typical extreme halophiles, growing optimally at 4 M total Na<sup>+</sup> and within the range from 3 to 5 M (tested at pH 8.8). The cells of isolates in both groups lyzed at salinity downshift below 2 M total Na<sup>+</sup>. The pH profiling of four cultures at 4 M total Na<sup>+</sup> showed that they are moderate alkaliphiles with optimal growth within a pH range from 8.0 to 8.8 (**Fig. 3**). The

maximum growth temperature of type strains grown with soluble starch at pH 8.5 and 4 M total Na $^+$  was 50 $^{\circ}$ C for the group 1 strains and 48 $^{\circ}$ C for the group 2 strains.

Comparative properties of the group 1 and group 2 isolates with their nearest phylogenetic relatives are shown in **Tables 2** and **3.** 

The main difference of the two groups from each other was in utilization of two different fructans and the phylogeny. The key difference of the group 1 isolates from the type species of the genus *Natrananaeroaechaeum* was their inability for anaerobic growth and inability to grow at extremely high pH values above 9.3. The main difference of the group 2 isolates from the nearest related genera is that they are the only ones isolated from soda lakes. The two out of three related genera (*Halohasta* and *Halonotius*) are definitely neutrophiles, while, despite the reported ability of "Halalkalirubrum salinum" to grow up to pH 10.5, there is a doubt about it. First, the organism is isolated from a salt lake with pH 8.5, thus being only a slightly alkaline salt (but not soda) lake. Even natronoarchaea isolated from hypersaline soda lakes with permanent pH above 10, seldom grow above pH 10.2. Secondly, the final pH values were apparently not measured during the pH profiling, which makes the reported values for the maximum pH unverified. Hence, the newly isolated amylolytic strains from soda lakes can still be considered as first obligate alkaliphilic (albeit only moderate) representatives of this group of related genera. They also differs from the other three genera in their alpha-glucan/fructan substrate specialization and the absence of glycoand sulfo-lipids in their membranes.

# Genomic analysis

The genome search the two type strains (dbCAN) identified a set of genes typically encoding alpha-amylases and alpha-glucosidases (GH13 and 15 families) in both representatives of two groups, although in AArc-St1-1 the total number and the fraction of putative extracellular amylases are much more abundant (Suppl. **Table S4**, consensus results from HHMER/DIAMOND

tools). This is also in agreement with the growth and amylase activity results (Suppl. **Fig. 1**). These enzymes would allow utilization of a spectrum of alpha-linked glucans such as amylopectin, soluble starch, dextrin, glycogen, pullulan, maltose and trehalose as sole carbon and energy sources. In this respect strain AArc-St1-1 is highly similar to the type species of the genus *Natranaeroarchaeum* [16-17]. Furthermore genomes of both type strains contain genes coding for beta-fructosidases of the GH families 32 and 68, which is in agreement with their ability to use fructans as growth substrate. As for the other glucanases encoded in two genomes, such as the beta-endo-1,3/1,4-glucanases of GH16 and 81 families (in AArc-St1-1), and pectin lyase (PL family) in AArc-St2, none of the tested potential polysaccharide substrates for these hydrolases supported growth (laminarin, lichenan, xylan, beta-glucan, glucomannan, mannan, curdlan, pachyman or pectin).

In respect to the osmoprotection and pH homeostasis, both genomes encode a range of typical potassium import complexes (but variable in copy numbers) and a multysubunit Na $^+$ /H $^+$  antiporter of the Mrp family. Both genomes lacks genes for organic osmolyte import and synthesis, indicating that the organisms rely solely on the potassium accumulation strategy. Both strains produce catalase/peroxidase and have a haem-copper family cytochrome c terminal oxidase of the  $aa_3$  type. In addition, strain AArc-St2 has another terminal oxidase of the  $ba_3$  type (Supplementary **Table S5**). These also agree with the positive tests for catalase and oxidase in both strains.

A major difference between the two type strains was found in the presence of two types of other respiratory complexes. Strain AArc-St2 genome contains a locus apparently coding for the aerobic type of CO-dehydrogenase (Cox, most probably of the type II) lacking in AArc-St1-1. Although the capacity to oxidize CO at low concentration has been demonstrated for several haloarchaeal species [31-32], the physiological role of such potential is still unclear. One of the possibilities is CO detoxification.

The genome of AArc-St1-1 contains loci apparently encoding enzymes for sulfurdependent anaerobic respiration, which are lacking in AArc-St2: two of them are highly homologous to the PsrABCD/SseA and PhsABCD complexes encoded in the genome of type species of genus Natranaeroarchaeum responsible for sulfur- and thiosulfate-dependent anaerobic respiration, respectively [27]. In addition, there is a second encoded PsrABCD lacking sulfur transferase more homologous to the one present in sulfur-respiring Halalkaliarchaeum desulfuricum (Supplementary Table S5). In our experience, such genomic potential must enable the anaerobic sulfur respiration in strain AArc-St1-1, similar to the type species Natranaeroarchaeum sulfidigenes. And this, indeed, is directly confirmed by the growth experiments, although the activity of sulfur reduction, in general, was lower in the novel isolates than in the type species of the genus Natranaeroarchaeum. Furthermore, none of the novel group 1 isolates were capable of thiosulfate-dependent anaerobic respiration, which was a prominent trait in the type species. On the other hand, the amylolytic isolates grew much better at fully aerobic conditions, while N. sulfidigenes needed transition via microaerophilic conditions before it started to grow at aerobic conditions. This reflects the difference in enrichment conditions used to isolate these closely related but still differentially specialized species of the same genus.

Overall, on the basis of distinct phenotypic, phylogenetic and genomic features, the group 1 isolates from hypersaline soda lakes are proposed to be classified in a novel species within the genus *Natranaeroarchaeum* as *Natranaeroarchaeum aerophilus* sp. nov. (type strain AArc-St1-1), while the group 2 isolates are forming a new species in a new genus for which the name *Natronocalculus amylovorans* gen. nov., sp. nov. (type strain AArc-St2). The protologues for the new taxa are presented in **Tables 4** and **5**.

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#### **Conflict of interests**

358 The authors declare that there is no conflict of interests.

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#### References

- 363 1. Oren, A. (2011) Diversity of halophiles. In: Horikoshi, K. (Ed.) Extremophiles Handbook. Springer, Tokyo, pp. 309-325.
- Andrei, A.Ş., Banciu, H.L., Oren, A. (2012) Living with salt: Metabolic and phylogenetic diversity of archaea inhabiting saline ecosystems. FEMS Microbiol. Lett. 330, 1-9.
- 367 3. Oren, A. (2013) Life at high salt concentrations. In: Rosenberg, E., DeLong, E.F., Lory, S., Stackebrandt, E., Thompson, F., (Eds.), The Prokaryotes: Prokaryotic Communities and Ecophysiology, Springer Berlin Heidelberg, Berlin, Heidelberg, pp. 421–40.
- 370 4. Oren, A. (2015) Halophilic microbial communities and their environments. Curr Opin Biotechnol 371 33, 119-24.
- 5. Grant, W.D., Jones, B.E. (2016) Bacteria, archaea and viruses of soda lakes. In: Schagerl, M., (Ed.), Soda Lakes of East Africa, Springer International Publishing, Cham, pp. 97–147.
- 6. Oren, A., Ventosa, A., Grant, W.D. (1997) Proposed minimal standards for description of new taxa in the order *Halobacteriales*. Int J Syst Bacteriol 47, 233–238.
- Sorokin, D.Y., Toshchakov, S.V., Kolganova, T.V., Kublanov, I.V. (2015) Halo(natrono)archaea
   isolated from hypersaline lakes utilize cellulose and chitin as growth substrates. Frontiers Microbiol
   6, 942.
- 379 8. Sorokin, D.Y., Khijniak, T.V., Kostrikina, N.A., Elcheninov, A.G., Toshchakov, S.V., Bale, N.J., 380 Sinnighe Damsté, J.S., Kubalnov, I.V. (2018) *Natronobiforma cellulositropha* gen. nov., sp. nov., a novel haloalkaliphilic member of the family *Natrialbaceae* (class *Halobacteria*) from hypersaline
- alkaline lakes. Syst Appl Microbiol 41, 355–362.
- Sorokin, D.Y., Elcheninov. A.G., Toshchakov, S.V., Bale, N.J., Sinninghe Damsté, J.S., Khijniak,
   T.V., Kublanov, I.V. (2019) *Natrarchaeobius chitinivorans* gen. nov., sp. nov., and
   *Natrarchaeobius halalkaliphilus* sp. nov., alkaliphilic, chitin-utilizing haloarchaea from hypersaline
   alkaline lakes. Syst Appl Microbiol 42, 309-318.
- 387 10. Kobayashi, T., Kanai H., Hayashi T., Akiba T., Akaboshi R., Horikoshi K. (1992) Haloalkaliphilic 388 maltotriose-forming a-amylase from the archaebacterium *Natronococcus* sp. strain Ah-36. J 389 Bacteriol 174, 3439-3444.
- Kana<u>i</u>, H., Yashi, K., T, Aono, R., Kudo, T. (1995) *Natronococcus amylolyticus* sp. nov., a
   haloalkaliphilic archaeon. Int J Syst Bacteriol 45, 762-766.
- 392 12. Pfennig, N., Lippert, K.D. (1966) Über das Vitamin B12-Bedürfnis phototropher Schwefelbakterien. Arch Mikrobiol 55, 245-256.

- 394 13. Sorokin, D.Y., Yakimov, M.M., Messina, E., Merkel, A.Y., Koenen, M., Bale, N.J., Sinninghe
- Damsté, J.S. (2021) Halapricum desulfuricans sp. nov., carbohydrate-utilizing sulfur-reducing
- haloarchaea from hypersaline lakes. Syst Appl Microbiol 44, 126249.
- 397 14. Bale, N.J., Sorokin, D.Y., Hopmans, E.C., Koenen, M., Rijpstra, W.I.C., Villanueva, L., Wienk, H.,
- 398 Sinninghe Damsté, J.S. (2019) New insights into the polar lipid composition of halophilic
- euryarchaea from hyper saline lakes. Front Microbiol 10, 377.
- 400 15. Elling, F.J., Becker, K.W., Könneke, M., Schröder J.M., Kellermann, M.Y., Thomm, M., Hinrichs,
- 401 K.-U. (2016) Respiratory quinones in Archaea: phylogenetic distribution and application as
- biomarkers in the marine environment. Environ. Microbiol 18, 692–707.
- 403 16. Bankevich, A., Nurk, S., Antipov, D., Gurevich, A.A., Dvorkin, M., Kulikov, A.S., Lesin, V.M.,
- Nikolenko, S.I., Pham, S., Prjibelski, A.D., Pyshkin, A.V., Sirotkin, A.V., Vyahhi, N., Tesler, G.,
- Alekseyev, M.A., Pevzner, P.A. (2012) SPAdes: A New Genome Assembly Algorithm and Its
- 406 Applications to Single-Cell Sequencing. J Comput Biol 19, 455–477.
- 407 17. Wick, R.R., Judd, L.M., Gorrie, C.L., Holt, K.E. (2017) Unicycler: Resolving bacterial genome
- 408 assemblies from short and long sequencing reads. PLoS Comput Biol 13, 1-22.
- 409 18. Gurevich, A., Saveliev, V., Vyahhi, N., Tesler, G. (2013) QUAST: Quality assessment tool for
- genome assemblies. Bioinformatics 29, 1072–1075.
- 411 19. Mikheenko, A., Prjibelski, A., Saveliev, V., Antipov, D., Gurevich, A. (2018) Versatile genome
- 412 assembly evaluation with QUAST-LG. Bioinformatics 34, i142–50.
- 413 20. Parks, D.H., Imelfort, M., Skennerton, C.T., Hugenholtz, P., Tyson, G.W. (2015) CheckM:
- Assessing the quality of microbial genomes recovered from isolates, single cells, and metagenomes.
- 415 Genome Res 25, 1043–55.
- 416 21. Rinke C, Chuvochina M, Mussig AJ, P-A Chaumeil, Davín AA, Waite DW, Whitman WB, Parks
- DH, Hugenholtz P. (2021) A standardized archaeal taxonomy for the Genome Taxonomy Database.
- 418 Nat Microbiol 6, 946-959.
- 419 22. Parks, D.H., Chuvochina, M., Waite, D.W., Rinke, C., Skarshewski, A., Chaumeil, P.-A.,
- Hugenholtz, P. (2018) A standardized bacterial taxonomy based on genome phylogeny substantially
- revises the tree of life. Nat Biotechnol 36(10), 996–1004.
- 422 23. Stamatakis, A. (2014) RAxML version 8: A tool for phylogenetic analysis and post-analysis of
- 423 large phylogenies. Bioinformatics 30, 1312–1313.
- 424 24. Letunic, I., Bork, P. (2019) Interactive Tree Of Life (iTOL) v4: recent updates and new
- developments. Nucleic Acids Res 47, W256-W259.
- 426 25. Sorokin, D.Y., Elcheninov, A.G., Khijniak, T. V., Zaharycheva, A.P., Boueva, O. V., Ariskina, E.
- V., Bunk, B., Spröer, C., Evtushenko, L.I., Kublanov, I. V., Hahnke, R.L. (2022)
- Natronosporangium hydrolyticum gen. nov., sp. nov., a haloalkaliphilic polyhydrolytic
- 429 actinobacterium from a soda solonchak soil in Central Asia. Syst Appl Microbiol 45, 126307.

- 430 26. Zhang, H., Yohe, T., Huang, L., Entwistle, S., Wu, P., Yang, Z., Busk, P.K., Xu, Y., Yin, Y. (2018)
- dbCAN2: a meta server for automated carbohydrate-active enzyme annotation. Nucleic Acids Res
- 432 46(W1), W95–101.
- 433 27. Sorokin, D.Y., Messina, E., Smedile, F., La Cono, V., Hallsworth, J.E. (2021) Carbohydrate-
- dependent sulfur respiration in halo(alkali)philic euryarchaea from hypersaline lakes. Environ
- 435 Microbiol 23, 3779-3808.
- 436 28. Sorokin, D.Y., Yakimov, M.M., Messina, E., Merkel, A.Y., Koenen, M., Bale N.J., Sinninghe
- Damsté J.S. (2022) Natranaerarchaeum desulfuricum gen. nov., sp. nov., carbohydrate-utilizing
- sulfur-respiring haloarchaeon from hypersaline lakes and proposal of a new family
- Natronoarchaeaceae fam. nov. in the order Halobacteriales. Int J Syst Evol Microbiol 72,
- 440 10.1099/ijsem.0.005332.
- 441 29. Gupta, R.S., Naushad, S., Fabros, R., Adeolu, M. (2016) A phylogenomic reappraisal of family-
- level divisions within the class *Halobacteria*: proposal to divide the order *Halobacteriales* into the
- families Halobacteriaceae, Haloarculaceae fam. nov., and Halococcaceae fam. nov., and the order
- 444 Haloferacales into the families, Haloferacaceae and Halorubraceae fam nov. Antonie van
- Leeuwenhoek 109, 565-587.
- 446 30. Luo, C., Rodriguez-R, L.M., Konstantinidis, K.T. (2014) MyTaxa: An advanced taxonomic
- classifier for genomic and metagenomic sequences. Nucleic Acids Res 42, e73.
- 448 31. King, G.M. (2015) Carbon monoxide as a metabolic energy source for extremely halophilic
- microbes: Implications for microbial activity in Mars regolith. Proc Nat Ac Sci USA 112, 4465-
- 450 4470.
- 451 32. Myers, M.R., King, G.M. (2020) Halobacterium bonnevillei sp. nov., Halobaculum saliterrae sp.
- nov. and *Halovenus carboxidivorans* sp. nov., three novel carbon monoxide-oxidizing Halobacteria
- from saline crusts and soils. Int J Syst Evol Microbiol 70, 4261-4268.
- 454 33. Shimane, Y., Hatada, Y., Minegishi, H., Mizuki, T., Echigo, A. (2010) Natronoarchaeum
- 455 mannanilyticum gen. nov., sp. nov., an aerobic, extremely halophilic archaeon isolated from
- 456 commercial salt. Int J Syst Evol Microbiol 60, 2529–2534.
- 457 34. Zhenqiang Zuo . Dahe Zhao . Jian Zhou . Jing Han . Hua Xiang (2021) Halalkalirubrum salinum
- gen. nov., sp. nov., a halophilic archaeon isolated from a saline lake. Ant van Leeuwenhoek 114,
- 459 83–94.
- 460 35. Mou, Y.-Z., Qiu, X.-X., Zhao, M.-L., Cui, H.-L., Oh, D., Dyall-Smith, M.L. (2012) Halohasta
- litorea gen. nov. sp. nov., and Halohasta litchfieldiae sp. nov., isolated from the Daliang
- aguaculture farm, China and from Deep Lake, Antarctica, respectively. Extremophiles 16, 895-901.
- 463 36. Durán-Viseras, A., Andrei A.-S., Ghai, R., Sánchez-Porro, C., Ventosa, A. (2019) New Halonotius
- species provide genomics-based insights into cobalamin synthesis in haloarchaea. Front Microbiol
- 465 10, 1928.

 **Table 1**. List of pure cultures of natronoarchaea enriched and isolated from hypersaline soda lakes in southwestern Siberia with amylopecting and fructans

<u> </u>			<b>7</b> 1	C
Strain	Source	Enrichment	Phylogenetic	Closest relative
		substrate	group	
AArc-St1-1 <sup>T</sup>	Sediments	Amylopectin	Group 1	Natranaeroarchaeum
AArc-St1-2	Brines			sulfidigenes
AArc-St1-3	Brines			
AArc-lev1	Sediments	Levan		
AArc-St2 <sup>T</sup>	Sediments	Amylopectin	Group 2	"Halalkalirubrum
AArc-St3	Brines			halophilum"
AArc-in2	Sediments	Inulin		

**Table 2**. Comparative properties of amylolytic natronoarchaea of group 1 with the type species of the nearest phylogenetically related genera [28, 33]

Property	"Natranaer <mark>o</mark> archaeum	Natranaer <u>o</u> archaeum	Natronoarchaeum
	aerophilu <u>s</u> "	sulfidigenes	mannanilyticum
	(4 strains)	JCM 34033 <sup>T</sup>	JCM 16328 <sup>T</sup>
Cell morphology	flat pleomorphic,	flat pleomorphic,	pleomorphic,
	motility not observed	motile	nonmotile
Pigmentation	red	red (aerobic); pink	red
		(anaerobic)	
PHA accumulation	-	-	-
Aerobic growth	+	+	+
Anaerobic growth by:			
sugar fermentation	-	-	-
sulfur/thiosulfate respiration	+ (with sulfur)	+ (with sulfur and $S_2O_{32-}$ )	-
sulfoxide respiration	-	-	-
Number of Psr/Phs operons	3	2	0*
in genomes			
<i>e</i> -donors for anaerobic	glucose, maltose	sugars, starch, glycerol	-
growth		_	
Substrates for aerobic growth	starch-like alpha-glucans,	sugars, starch,	lactose, raffinose, sucrose,
	levan, maltose, cellobiose,	yeast extract	maltose, cellobiose, starch,
	trehalose		galactomannan, pyruvate, lactate,
A 1			glutamate, yeast extract, peptone
Amylase	+	+	+ (T. 90)
Esterase/lipase Protease	-	- (tributyrin/olive oil)	- (Tween 80)
Protease Catalase/oxidase	- +/+	- (gelatin, casein)	- (gelatin)
	+/+	+/+(w)	-/ +(w) +
Indole from tryptophane Salinity range (opt.) (M Na <sup>+</sup> )	3.0-5.0 (4.0)	2.5-4.5 (3.5)	1.6-4.2 (2.5-3.2)
pH range (opt.)	7.2-9.3 (8.0-8.8)	8.5-10.2 (9.5-9.7)	6.0-9.5 (8.5-9.0)
Temperature max (°C)	50 (at pH 8.5)	45 (at pH 9)	55
Core lipids	C <sub>20</sub> -C <sub>20</sub> , C <sub>20</sub> -C <sub>25</sub> DGE	C <sub>20</sub> -C <sub>20</sub> , C <sub>20</sub> -C <sub>25</sub> DGE	NR
Intact membrane polar lipids:	220 C20, C20 C23 DGL	220 C20, C20 C23 DGL	
phospholipids	PG, PGP-Me	PG, PGP-Me	PG, PGP-Me, PGP,
glycolipids/sulfolipids	-	-	S <sub>2</sub> -DGDE
Respiratory lipoquinones	MK-8:8	MK-8:8	NR
DNA G+C	61.0 (genome)	60.8% (genome)	63.0 (mol%)
Type of hypersaline habitat		e soda lakes	Marine solar saltern

NR, not reported; (v) - variable property in different species of the same genus; w (weak); Psr/Phs - polysulfide/thiosulfate reductase; \*genome of *N. phillipinensis*. Lipids: (PG) phosphatidylglycerol, phosphatidylglycero-phosphate (PGP), (PGP-Me) phosphatidylglycerophosphate methyl ester, disulfated diglycosyl diether (S2-DGDE), (DGE) - dialkyl glycerol ether.

**Table 3**. Comparative properties of group 2 isolates with nearest phylogenetic relatives [34-36].

Property	"Natranocalculus amylovorans" (3 isolates)	"Halalkalirubrum halophilu <u>m</u> "	Halohasta (2 species)	Halonotius (4 species)
Cell morphology	flat pleomorphic nonmotile	pleomorphic, nonmotile	rods, motile	polymorphic rods, motility (V)
Pigmentation	red	red	red	red
PHA accumulation	-	NR	NR	
Anaerobic growth by: sugar fermentation	-	NR	NR	NR
sulfur respiration	-	NR	NR	NR
DMSO respiration	-	-	-	+(1 species)
Growth substrates				
carbohydrates:	starch-like alpha-	glucose, maltose,	glucose, sucrose;	glucose, arabinose,
•	glucans, inulin,	fructose, sorbose,	mannose, galactose,	fructose, galactose,
	maltose, cellobiose,	lactose, xylose,	lactose, maltose (all	sucrose, maltose,
	trehalose, glycerol	mannitol, sorbitol	V)	raffinose, xylose,
				mannitol, sorbitol, glycerol, (all V)
organic acids:	none	acetate, pyruvate, lactate, fumarate, succinate, citrate	pyruvate, lactate, succinate, malate, fumarate, citrate (V);	pyruvate, citrate, tartrate (all V)
Amylase	+ (soluble starch)	- (soluble starch)	- (soluble starch)	- (soluble starch)
Esterase/lipase	- (tributyrin/olive oil)	- (Tweens)	- (Tween 80)	- (Tween 80)
Protease	- (gelatin, casein)	- (gelatin, casein)	- (gelatin, casein)	- (gelatin, casein)
Catalase/oxidase	+/+	+/+	+/V	V/V
Indole from tryptophane	+(w)	+	-	NR
Salinity range (opt.) M Na <sup>+</sup>	3-5 (4.0)	1.9-4.2 (2.5)	2.0-4.7 (2.5-3.0)	2.5-6.0**(3.0-4.0)
Mg <sup>2+</sup> demand	low	low	high	high
pH range (opt.)	7.2-9.3 (8.5-8.8)	7.0-10.5* (8.5-9.5)	5.5-9.0* (7.0-7.5)	5.0-9.0 (7.0-7.5)
Temperature max (°C)	48 (at pH 8.5)	42	45-50	45-50
Core lipids	C <sub>20</sub> -C <sub>20</sub> , C <sub>20</sub> -C <sub>25</sub> DGE	NR	NR	NR
Intact membrane polar lipids:				
phospholipids:	PG, PGP-Me	PG, PGP-Me	PG, PGP-Me, PA	PG, PGP-Me
glycolipids:	-	1, unidentified	S-DGD-1	S-DGD-1
sulfolipids:	-	PGS	ND.	PGS (V)
Respiratory lipoquinones	MK-8:8	NR	NR	NR
DNA G+C (% genomic)	<u>5</u> 1. <u>5</u> (type strain)	58.4 (type strain)	58.8 (type species)	59.7-62.7 (4 species)
Type of hypersaline habitat	soda lakes	salt lake	solar saltern	solar saltern, saline soils
Havitat				Samle Sons

NR, not reported; (V), variable property in different species of the same genus; \* actual final pH values were not measured; \*\* reported for the type species, but not verified in any further research; PA, phosphatidic acid; PGS, phosphatidylglycerol sulfate; S-DGD-1, sulfated mannosyl glucosyl diether; other abbreviations (see **Table 2**).

Table 4. Natranaeroarchaeum aerophilus: protologue

Table 4. Natranaeroarchaeum aerophilus: protologue			
Parameter	Species: Natranaeroarchaeum aerophilus sp. nov.		
Author	Dimitry Y. Sorokin		
Species name	aerophilu <u>s</u>		
Genus name	Natranaeroarchaeum		
Specific epithet Species status (SPST)	sp. nov.		
Etymology	a.e.ro'phi.lus Gr. masc. n. aer, air; N.L. masc. adj. philus (from Gr. masc. adj. philos), friend,		
Etymology	loving; N.L. masc. adj. aerophilus, air-loving.		
Description of the new taxon  Authors	The cells are angular, flat, polymorphyc coccoids or rods, mostly nonmotile, variable in size from 1 to 3 µm. The cells lyze in hypotonic solutions below 1 M NaCl. produces red carotenoids. The core membrane diether lipids are composed of C <sub>20</sub> -C <sub>20</sub> DGE (extended archaeol). The polar lipid head groups include phosphatidylglycerolphosphate methyl ester (PGP-Me) and phosphatidylglycerol (PG). The dominant respiratory quinone is MK-8:8. Saccharolytic with limited substrate profile including several starch-like alpha-glucans, levan, maltose, trehalose and cellobiose. Capable of anaerobic sulfogenic growth with glucose and maltose as electron donors and sulfur as electron acceptor. Ammonium, urea and yeast extract serve as the N-source. Oxidase and catalase positive. Maximum growth temperature is 50°C. Extremely halophilic with a range of total Na <sup>+</sup> for growth from 3 to 5 M (optimum at 4 M) and moderately alkaliphilic, with a pH range for growth from 7.2 to 9.3 (optimum at 8.0-8.8). The G + C content of the DNA is 61.0% (genome of the type strain). Habitat - aerobic sediments and brines of hypersaline soda lakes. The type strain (AArc-St1-1 <sup>T</sup> 2CM 32519 <sup>T</sup> =UQM 41458 <sup>T</sup> ) was isolated aerobic sediments of hypersaline soda lakes in Kulunda Steppe (Altai, Russia). The species also includes other three closely related strains isolated from the same area. The draft genome of type strain is deposited in the GenBank under accession number JAKRVY0000000000.		
Title	Dimitry Y. Sorokin, Alexander G. Elcheninov, Tatjana V. Khizhniak, Michel Koenen, Nicole J. Bale, Jaap S. Sinninghe Damsté, Ilya V. Kublanov  Natronocalculus amylovorans gen. nov., sp. nov., and Natranaeroarchaeum aerophilus sp.		
Title	nov., dominant culturable amylolytic natronoarchaea from hypersaline soda lakes in southwestern Siberia.		
Journal	Systematic and Applied Microbiology		
Corresponding author	Dimitry Y. Sorokin		
E-mail of corresponding author	soroc@inmi.ru; d.sorokin@tudelft		
Designation of the type strain	AArc-St1-1		
Strain collection numbers  16S rRNA gene accession numbers	JCM 32519 <sup>T</sup> =UQM 41458 <sup>T</sup> MG584707- MG584709; ON003450		
Genome accession numbers	JAKRVY00000000 (type strain)		
Genome status	Draft		
G+C, %	61.0 (genome of type strain)		
Country of origin	Russian Federation		
Region of origin	Altai region		
Date of isolation	2016 Surface conchine and imports from hymercaline and a later		
Source of isolation Sampling dates	Surface aerobic sediments from hypersaline soda lakes 2015-07-07		
Geographic location	S-W Siberia, Kulunda Steppe; southern Russia		
Latitude	51°39' N; 49°10' N; 48°14' N		
Longtitude	79°48' E; 46°39' E; 46°35' E		
Depth	0-2 cm		
Temperature of the sample	20°C		
pH of the sample	10-11		
Salinity of the sample Number of strains in study	18-36% 4		
Source of isolation of non-type strains	Surface aerobic sediments and brines from hypersaline soda lakes, S-W Siberia, Kulunda		
Growth medium, incubation conditions	Steppe; southern Russia  4 M total Na <sup>+</sup> , pH 9; incubation - 37°C; starch as substrates; aerobic		
Conditions of preservation	Deep freezing in 15% glycerol (v/v)		
Gram stain	Negative		
Cell shape	Pleomorphic flat coccoids		
Cell size	0.8-2 µm in diameter		
Motility (MOTY)	Mostly nonmotile		
Sporulation Colony morphology	None Flat, compact, max. 2 mm, red		
Temperature range for growth	nd		
Lowest temperature for growth	nd		
Highest temperature for growth	50		
Optimal temperature for growth	37-40		
Lowest pH for growth	7.2		
Highest pH for growth	9.3		
Optimum pH for growth pH category	8-8.8 Moderately alkaliphilic		
Lowest Na <sup>+</sup> concentration for growth	3.0 M		
Highest Na <sup>+</sup> concentration for growth	5.0 M		
Optimum salt concentration for growth	4.0 M total Na <sup>+</sup>		
Other salts important for growth	KCl; Na-carbonates		
Salinity category	Extremely halophilic		
Relation to oxygen	Facultatively anaerobic		
O <sub>2</sub> conditions for strain testing Carbon source used (class)	Fully aerobic Carbohydrates		
Carbon source used (Class)	Caronjanuo		

Specific compounds	Starch-like alpha-glucans, levan
Nitrogen source	Ammonium, urea, yeast extract
Terminal electron acceptor	$O_2$ and $S_8$
Energy metabolism	Chemoorganotrophic
Phospholipids	Core membrane lipids are C <sub>20</sub> -C <sub>20</sub> DGE (archaeol) and C <sub>20</sub> -C <sub>25</sub> DGE (extended archaeol). Polar head groups are phosphatidylglycerophosphate methylester (PGP-Me) and phosphatidylglycerol (PG)
Respiratory lipoquinones	MK-8:8
Glycolipids (GLYC)	-
Habitat (HABT)	Hypersaline soda lakes
Extraordinary feautures (EXTR)	Narrowly specialized amylolytics

 Table 5. Natronocalculus amylovorans: protologue

Table 5. Natronocalculus amylo	ı Ç		
Parameter	Genus: Natronocalculus gen. nov.	Species: Natronocalculus amylovorans sp. nov.	
Author (AUTE)	Dimitry Y. Sorokin	Ι ,	
Species name (SPNA)	Natura	amylo <u>voran</u> s	
Genus name (GENA) Specific epithet (SPEP)	Natronocalculus -	amylovorans	
Species status (SPST)	_	sp. nov.	
Etymology (GETY/SPTY)	Na.tro.no.cal'cu.lus N.L. neut. n. natron, arbitrarily derived from the Arabic n. natrun or natron soda; L. masc. n. calculus, pebble, gravel; N.L. masc. n. Natronocalculus, soda loving pebble-shaped cells	a.my.lo.vo' <u>rans</u> . Gr. neut. n. <i>amylon</i> , starch; L. inf. v. <i>vorare</i> , to devour; N.L. part. adj. <i>amylovorans</i> , <u>eating</u> starch	
Type species of the genus (TYPE)	Natronocalculus amylovorans	yes	
Description of new taxon	Obligately aerobic and organotrophic member of <i>Halobacteriales</i> narrowly specialized on utilization of starch-like polymers. Core lipids are dominated by archaeol and extended archaeol with PGP-Me and PG polar groups. Glycolipids are not present. MK-8:8 is the dominant lipoquinone. Extremely halophilic and moderately alkaliphilic inhabitans of hypersaline soda lakes. The three-letter abbreviation is Ncl.	The cells are angular, flat, polymorphyc coccoids or rods, nonmotile, from 0.8 to 2.0 μm. The cells lyze in hypotonic solutions below 1 M NaCl. Colonies are orange-red. The core membrane diether lipids include of C <sub>20</sub> -C <sub>20</sub> DGE (archaeol) and C <sub>20</sub> -C <sub>25</sub> DGE (extended archaeol). The polar lipid head groups consists of hosphatidylglycerolphosphate methyl ester (PGP-Me) and phosphatidylglycerol (PG). The dominant respiratory menaquinone is MK-8:8. Obligately aerobic saccharolytic with limited substrate profile including several starch-like alpha-glucans, inulin, maltose, trehalose, cellobiose and glycerol. Ammonium and yeast extract serve as the N-source. Oxidase and catalase positive. Maximum growth temperature is 48°C. Extremely halophilic with a range of total Na <sup>+</sup> for growth from 3 to 5 M (optimum at 4 M) and moderately alkaliphilic, with a pH range for growth from 7.2 to 9.3 (optimum at 8.5-8.8). The G + C content of the DNA is 51.5% ( genome of the type strain). Habitat - aerobic sediments and brines of hypersaline soda lakes. The type strain (AArc-St2 <sup>T</sup> =JCM 32475 <sup>T</sup> =UQM 41459 <sup>T</sup> ) was isolated from aerobic sediments of hypersaline soda lakes in Kulunda Steppe (Altai, Russia). The species also includes 2 other closely related strains isolated from the same area. The draft genome of type strain is deposited in the GenBank under accession number JAKRVX0000000000.	
Authors (AUT)	Dimitry Y. Sorokin, Alexander G. Elcheninov, Tatjana V. Khizhniak, Michel Koenen, Nicole J.		
Title (TITL)	Bale, Jaap S. Sinninghe Damsté, Ilya V. Kublanov  Natronocalculus amylovorans gen. nov., sp. nov., and Natranaeroarchaeum aerophilus sp. nov., dominant culturable amylolytic natronoarchaea from hypersaline soda lakes in southwestern Siberia.		
Journal (JOUR)	Systematic and Applied Microbiology		
Corresponding author (COAU) E-mail of corresponding author (EMAU)	Dimitry Y. Sorokin d.sorokin@tudelft; soroc@inmi.ru		
Strain collection numbers (COLN)	-	JCM 32475; UQM 41459	
16S rRNA gene accession number (16 SR)	-	MG584710; ON000203; ON000205	
Genome accession numbers		JAKRVX000000000	
Genome status (GSTA)		Draft	
GC mol % (GGCM)	-	<u>5</u> 1. <u>5</u> (genome type strain)	
Country of origin (COUN)	Russian Federation	Russian Federation	
Region of origin (REGI)	-	Altai region	
Date of isolation (DATI)	-	2016	
Source of isolation (SOUR)	Hypersaline soda lakes	Surface sediments and brines of hypersaline soda lakes in southwestern Siberia	
Sampling dates (DATS)	2015	2015	
Geographic location (GEOL)	S-W Siberia	S-W Siberia	
Latitude (LATI)	-	51°39' N; 49°10' N; 48°14' N	
Longtitude (LONG)	-	79°48′ E; 46°39′ E; 46°35′ E	
Depth (DEPT)		0-2 cm	
Temperature of the sample (TEMS)		20°C	
pH of the sample (PHSA)		10-11	
Salinity of the sample (SALS)		18-36%	
Number of strains in study (NSTR)	3	3	
Source of isolation of non-type strains (SAMP)	-	Surface sediments and brines of hypersaline soda lakes in southwestern Siberia	
Growth medium, incubation conditions (CULT)		4 M total Na <sup>+</sup> , pH 9; incubation - 37°C; starch as substrates; aerobic	
Conditions of preservation (PRES)	Deep freezing in 15% glycerol (v/v)		
Gram stain (GRAM)	negative		
Cell shape (CSHA)	Pleomorphic, from flat irregular coccoids		
Cell size (CSZI)	-	0.8-2 μm in diameter	
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Motility (MOTY)	- n	nonmotile	
Sporulation (SPOR)	none		
Colony morphology (COLM)	P	Pink-orange, up to 2 mm, flat	
Temperature range for growth (TEMR)		-	
Lowest temperature for growth (TEML)			
Highest temperature for growth(TEMH)	4	8 (at pH 8.5)	
Optimal temperature for growth	3	37-40 °C	
(TEMO)			
Lowest pH for growth (PHLO)		1.2	
Highest pH for growth (PHHI)	9	0.3	
Optimum pH for growth (PHOP)	1 -	3.5-8.8	
pH category (PHCA)	alkaliphile (optimum > 8.5)		
Lowest NaCl concentration for growth	3.0 M total Na <sup>+</sup>		
(SALL)			
Highest NaCl concentration for growth	5 M total Na <sup>+</sup>		
(SALH)			
Optimum salt concentration for growth	4.0 M total Na <sup>+</sup>		
(SALO)	Sodium carbonates		
Other salts important for growth			
Salinity category (SALC)	extremely halophilic		
Relation to oxygene (OREL)	aerobe		
O <sub>2</sub> conditions for strain testing (OCON)	aerobic		
Carbon source used (class) (CSUC)	carbohydrates		
Specific compounds (CSUC)	Starch-like alpha glucans and inulin		
Nitrogen source (NSOU)	Ammonium, yeast extract		
Terminal electron acceptor (ELAC)	02		
Energy metabolism (EMET)	chemoorganotrophic		
Phospholipids (PHOS)	Core membrane lipids are archaeol (C <sub>20</sub> -C <sub>20</sub> DGE) and extended archaeol (C <sub>20</sub> -C <sub>25</sub> DGE)		
	Polar lipids are phosphatidylglycerophosphate methyl ester (PGP-Me) and phosphatidylglycerol (PG)		
Glycolipids (GLYC)	-		
Respiratory lipoquinones	MK8:8		
Habitat (HABT)	Hypersaline soda lakes		

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# Legends to the figures

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- 501 Fig. 1. Cell morphology (phase contrast microphotograps) of starch-utilizing natronoarchaea growing
- 502 aerobically at 4 M total Na<sup>+</sup>, pH 9 and 37°C. (a-d), group 1, including strains AArc-St1-1<sup>T</sup>, AArc-St1-
- 503 2, AArc-St1-3 and AArc-lev11, respectively. (e-f), group 2, including strains AArc-St2<sup>T</sup> and AArc-
- 504
- 505 Figure 2. Phylogeny of amylolytic natronoarchaea.
- 506 (a) Figure X. (a) 16S rRNA gene sequence-based maximum-likelihood phylogenetic tree, showing the position 507 of AArc-St2<sup>T</sup> and AArc-St1-1<sup>T</sup> (in bold) within the *Halobacteria* class. The black circles at nodes indicate that 508 the percentage of corresponding support values was above 50. Archaeoglobus fulgidus VC-16<sup>T</sup>, Methanocella 509 paludicola SANAE<sup>T</sup>, Methanothermobacter thermautotrophicus Delta H<sup>T</sup> were used as an outgroup (not shown). 510 Species in clusters: I (Halodesulfurarchaeum, Halanaeroarchaeum, Halarchaeum, Halobacterium, 511 Salarchaeum, Halocalculus); II (Halomicroarcula, Haloarcula, Halorientalis, Halorhabdus, Halococcoides, 512 Halapricum, Salinirussus, Halovenus); III (Haloglomus, Natronomonas, Halosegnis, Salinirubellus, 513 Halomarina, Halocatena); IV (Halopelagius, Haloferax, Halogeometricum, Haloquadratum, Halobellus); V 514 (Halococcus, Halalkalicoccus, Haloarchaeobius, Halorubellus); VI (Halorussus, Halomicrococcus, 515 Haladaptatus); VII (Saliphagus, Natribaculum, Halovarius, Natronococcus, Halovivax, Natronobiforma, Halostagnicola, Natronobacterium, Halopiger, Halobiforma, Natrarchaeobaculum, Natronolimnohabitans, Natronolimnobius, Natronorubrum, Natrinema, Haloterrigena, Natrialba, Natrarchaeobius, Salinadaptatus); VIII (Halosimplex, Salinibaculum, Halosiccatus, Halomicrobium); IX (Salinigranum, Haloplanus, Halobium,
- 516 517 518 519 520 521 Halegenticoccus, Halogranum, Haloprofundus, Halolamina); X (Halalkaliarchaeum, Halopenitus, Halorubrum, Haloparvum). (b) Maximum likelihood phylogenetic tree based on concatenated alignment of 122 conserved archaeal proteins and showing position of strains AArc-St2<sup>T</sup> and AArc-St1-1<sup>T</sup> (in bold) within the class *Halobacteria*. The branch
- 522 523 524 525 526 527 lengths correspond to the number of substitutions per site with corrections associated with the models. The black circles at nodes indicate that the percentage of corresponding support values was above 50. Archaeoglobus fulgidus VC-16<sup>T</sup>, Methanocella paludicola SANAE<sup>T</sup> and Methanothermobacter thermautotrophicus Delta H<sup>T</sup> were used as an outgroup (not shown). Species in clusters: I (Halanaeroarchaeum, Halodesulfurarchaeum, Halarchaeum, Halobacterium, Salarchaeum, Halocalculus); II (Halalkalicoccus, Halorussus, Halomicrococcus, 528 529 Haladaptatus); III (Haloarchaeobius, Halorubellus, Halovivax, Saliphagus, Natronobiforma, Halostagnicola, Natrarchaeobius, Natrarchaeobaculum, Salinadaptatus, Halopiger, Natronolimnobius, Natronobacterium, 530 Halobiforma, Natrialba, Natronococcus, Natrinema, Haloterrigena, Natronorubrum, Natronolimnohabitans); 531 IV (Halococcus, Halocatena, Halomarina, Natronomonas, Haloglomus, Halosegnis, Halorientalis, Halapricum, 532 Salinibaculum, Halovenus, Halosimplex, Halococcoides, Halorhabdus, Halomicrobium, 533 Halosiccatus, Halomicroarcula, Haloarcula); V (Haloplanus, Salinirubrum, Haloprofundus, Halegenticoccus,

534 Halogranum, Salinigranum, Haloferax, Halopelagius, Halogeometricum, Halobellus, Haloquadratum).

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- Fig. 3. pH profiles for growth with soluble starch in amylolytic natronoarchaea at 4 M total Na<sup>+</sup> and
- 538 37°C. Actual final pH are shown. The results are average from two parallel incubations.

Natronocalculus amylovorans gen. nov., sp. nov., and Natranaeroarchaeum aerophilus sp. nov., dominant culturable amylolytic natronoarchaea from hypersaline soda lakes in southwestern Siberia. Dimitry Y. Sorokin<sup>a,b\*</sup>, Alexander G. Elcheninov<sup>a</sup>, Tatjana V. Khizhniak<sup>a</sup>, Michel Koenen<sup>c</sup>, Nicole J. Bale<sup>c</sup>, Jaap S. Sinninghe Damsté<sup>c</sup> and Ilya V. Kublanov<sup>a</sup> <sup>a</sup>Winogradsky Institute of Microbiology, Research Centre of Biotechnology, Russian Academy of Sciences, Moscow, Russia <sup>b</sup>Department of Biotechnology, Delft University of Technology, The Netherlands <sup>c</sup> NIOZ Royal Netherlands Institute for Sea Research, Department of Marine Microbiology and Biogeochemistry, Texel, The Netherlands \*Author for correspondence: D.Y. Sorokin; e-mail: soroc@inmi.ru; d.sorokin@tudelft.nl Running title: Natronocalculus amylovorans gen. nov., sp. nov. and Natranaeroarchaeum aerophilus sp. nov. The draft genome sequences of strains AArc-St1-1<sup>T</sup> and AArc-St2<sup>T</sup> are deposited in the GenBank under the numbers JAKRVY000000000 and JAKRVX000000000, respectively 

#### Abstract

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Several pure cultures of alkaliphilic haloaloarchaea were enriched and isolated from hypersaline soda lakes in southwestern Siberia using amylopectin and fructans as substrates. Phylogenomic analysis placed the isolates into two distinct groups within the class *Halobacteria*. Four isolates forming group 1 were closely related to a recently described Natranaeroarchaeum sulfidigenes and the other three strains forming group 2 represent a novel genus-level phylogenetic lineage. All isolates are saccharolytic archaea growing with various starch-like alpha-glucans including soluble starch, amylopectin, dextrin, glycogen, pullulane and cyclodextrin. In addition, group 1 can also use levan while group 2 - inulin (plant storage beta-fructans). Group 1 strains can also grow anaerobically with either glucose or maltose using elemental sulfur as the electron acceptor. Both groups are moderately alkaliphilic with a pH range for growth from 7.2 to 9.3 (optimum between 8.0-8.8) and low Mg-demanding extreme halophiles growing optimally at 4 M total Na<sup>+</sup>. The major respiratory menaquinone is MK-8:8 and the core biphytanyl lipids are dominated by archaeol (C<sub>20</sub>-C<sub>20</sub>) and a less abundant extended archaeol (C<sub>20</sub>-C<sub>25</sub>) with PG and PGP-Me as polar groups. The four isolates of group 1 are suggested to be classified into a new species as Natranaeroarchaeum aerophilus sp. nov. (type strain AArc-St1-1<sup>T</sup>=JCM 32519<sup>T</sup>=UQM 41458<sup>T</sup>). The three isolates of group 2 are proposed to form a new genus and species for which the name Natronocalculus amylovorans gen. nov., sp. nov. is suggested (type strain AArc-St2<sup>T</sup>=JCM 32475<sup>T</sup>=UQM 41459<sup>T</sup>).

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Key words: hypersaline soda lakes, natronoarchaea, amylolytic, starch, fructans,

Natranaeroarchaeum, Halobacteria

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#### Introduction

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Most of the known species of aerobic extremely halophilic and haloalkaliphilic euryarchaea (natronoarchaea) were enriched and isolated from hypersaline alkaline lakes on unspecific media containing rich soluble organic substrates, such as peptone, yeast extract or simple sugars [1-5]. Yet, these extremophilic, organoheterotrophic archaea are definitely more important in organic matter mineralization, in particular organic polymers, in hypersaline systems than is widely recognized. For example, a test for amylolytic activity with soluble starch is included into the minimal standards for taxonomy characterization of haloarchaea [6] but starch is rarely used for targeted isolation of amylolytic haloarchaea which would not only hydrolyze the polymer but also able to utilize it as growth substrate. And this situation is also true for other polysaccharides. So far, only a few examples of haloarchaea specialized on utilization of recalcitrant polysaccharides have been described in pure culture, which is particularly true for natronoarchaea living in alkaline hypersaline (soda) lakes. Our recent targeted enrichments from such lakes using insoluble forms of cellulose and chitin resulted in isolation of several groups of natronoarchaea highly specialized on utilization of either various forms of cellulose and xylan (genera Natronobiforma and Natronolimnobius) or chitin (genus Natrarchaeobius) [7-9]. Following further in this direction, we used insoluble starch (amylopectin) or beta-fructans as substrates to enrich for amylolytic or fructanolytic natronoarchaea from hypersaline soda lakes. So far, we are aware only of a single amylolytic natronarchaeon, Natronococcus amylolyticus, specifically isolated from the hypersaline soda lake Magadi in Kenya using starch as a growth substrate [10-11], while no natronoarchaea, growing on fructans are currently known.

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Here we describe phenotypic, phylogenetic and genomic properties of two novel taxa of amylolytic natronoarchaea enriched from hypersaline soda lakes in southwestern Siberia, which specialized on utilizing various alpha-glucan and beta-fructan polysaccharides as growth substrates.

# **Materials and Methods**

Enrichment and cultivation conditions

The sources of inocula were mixed surface (0-3 cm deep) aerated sediments and the near bottom brines obtained from four hypersaline soda lakes in Kulunda Steppe (Tanatar and Bitter lake systems, Altai region). The lakes brines total salt concentration varied from 20 to 40%, the total soluble carbonate alkalinity - from 2.5 to 5 M and the pH - from 10.2 to 11.0. The top flocculant sediment layer together with the near bottom brines was sucked into 50 ml syringe through silicon tubing and placed into a sterile 50 ml Falcon tube, resulting in an approximate volumetric ratio between the solid and liquid fractions of 1:5. After transportation to the laboratory the samples were separated onto clean brine top layer and concentrated sediment, all of which then were kept at 4°C. Before inoculation, the two fractions from each of four lakes were combined in equal proportions to make two master mixes (brine and sediments) used as inocula at 1% final (v/v).

For the enrichment and further cultivation of pure cultures two basic mineral media were used, both containing 4 M total Na<sup>+</sup>. The neutral 4 M NaCl base medium included (g l<sup>-1</sup>): NaCl, 240; K<sub>2</sub>HPO<sub>4</sub> 2.5 g l<sup>-1</sup>; NH<sub>4</sub>Cl 0.5 g l<sup>-1</sup>, KCl 5 g l<sup>-1</sup>, 20 mg l<sup>-1</sup> yeast extract and was adjusted to pH 7 with 10% KH<sub>2</sub>PO<sub>4</sub>. The alkaline sodium carbonate base contained (g l<sup>-1</sup>): Na<sub>2</sub>CO<sub>3</sub> 190, NaHCO<sub>3</sub> 45, NaCl 16, KCl 5, K<sub>2</sub>HPO<sub>4</sub> 1 and 20 mg l<sup>-1</sup> yeast extract (pH 10). After sterilization, the basic media were supplemented with 1 ml l<sup>-1</sup> of trace metal and vitamin solutions [12] and 1 mM MgSO<sub>4</sub>. NH<sub>4</sub>Cl (4 mM) was also added from 1 M sterile stock solution to the carbonate base after sterilization. To prepare final medium with a certain pH/alkalinity, the two ready to use bases were mixed in different proportions resulting in pH range from 8 to 10. For the enrichments from salt lakes, the neutral base medium was used as it is, while for the soda lake enrichments the neutral and alkali base media were mixed 3:1 with the final pH 9.5. For the pH range from 7 to 8, the NaCl base was supplied with various amounts of 1 M filter-sterilized NaHCO<sub>3</sub>, while for the pH below 7

it was titrated with sterile 1 M KH<sub>2</sub>PO<sub>4</sub>. Carbon and energy substrtates were added from sterile 10% stock solution.

For isolation of pure cultures, several rounds of enrichments were repeated at dilutions 1:100 and finally the sediment-free enrichments were plated onto solid medium obtained by mixing the liquid alkaline medium and 4% washed agar at 50°C in ratio 3:2. To compensate for the decreased salinity, solid NaCl was added to the liquid medium before mixing with melted agar to bring the final salt concentration back to 4 M total Na<sup>+</sup>. The isolation of pure cultures was achieved from separate colonies which grew back in liquid medium with the target polysaccharide and the purity was confirmed by 16S rRNA gene and genome sequencing.

### Pure culture characterization

Cell morphology was examined by using phase contrast microscopy (Zeiss Axioplane Imaging 2, Germany). Substrate utilization profiles were performed in medium containing 1 part of the alkaline base and 7 parts of the NaCl base (final pH 9.0). For the pH profiling, the two media were mixed in various proportions as described above and soluble starch served as the substrate. The growth was measured by increase of OD<sub>600</sub> with pH monitoring at each point. Anaerobic cultivation was performed as described previously [13]. Catalase and oxidase activity were tested with 3% (v/v) H<sub>2</sub>O<sub>2</sub> and 0.1% *N,N,N,N* tetramethyl-p-phenylenediamine hydrochloride, respectively, using cell-free extract (obtained by sonication) from cells of type strains. The protease, esterase/lipase activities were tested on plates spotted with fully grown liquid cultures: using casein/gelatin (hydrolyzis zones after flooding with 10% TCA) and emulsified tributyrin/olive oil (turbidity clearance), respectively. Antibiotic sensitivity of type strains AArc-St1-1 and AArc-St2 was tested in liquid medium at pH 9 with starch as substrate.

The intact polar lipids (IPLs) and respiratory quinones were analyzed as described previously [14]. Briefly, the lipid fraction was extracted from freeze-dried cells with sonication in

methanol:dichloromethane:phosphate buffer (2:1:0.8, v:v), followed by phase separation by adjusting the solvent mix to a ratio 1:1:0.9. The lipids and quinones were analyzed by normal phase, high performance liquid chromatography-ion trap mass spectrometry (HPLC-ITMS) and identified by masses and mass spectral fragmentation according to literature [14-15].

#### Genome sequencing

Genomic DNA from the type strains AArc-St1-1 and AArc-St2 was extracted using DNeasy PowerLyzer Microbial Kit (Qiagen) according to manufacturer instructions. Quality and quantity of the DNA samples were measured with Trinean Xpose spectrophotometer (PLT Scientific Instruments) and Qubit 2.0 fluorometer (Thermo Fisher Scientific) DNA libraries were prepared using KAPA HyperPlus kit (KAPAbiosystems) according to manufacturer recommendations. Paired-end sequencing (2x100bp) was performed using Illumina NextSeq. Obtained reads were filtered (quality and length) with CLC Genomics Workbench v.10. Genomes of the strains were assembled using SPAdes v.3.15.2 [16] --isolate mode with --trusted-contigs option (contigs obtained from Unicycler v.0.4.9 [17] were used as trusted contigs). Contigs with length  $\leq$  500bp or with low coverage were eliminated. Genome assemblies statistics were checked with Quast v.5.0.2 [18-19]. Completeness and contamination levels were detected using CheckM v.1.1.2 [20] with archaea-specific marker set.

### Phylogenetic and genomic analyses

For 16S rRNA gene sequence-based phylogenetic analysis 16S rRNA gene sequences of the seven isolates were aligned with the sequences of type species of all genera within *Halobacteria* (as well as *Archaeoglobus fulgidus* VC-16, *Methanocella paludicola* SANAE, *Methanothermobacter thermautotrophicus* Delta H used as the outgroup). Multiple sequence alignment and phylogenetic tree construction were performed as described earlier [9]. For phylogenomic analysis based on the

"ar122" set of conserved single copy archaeal proteins [21] the protein sequences were identified and aligned in *in silico* proteomes of type species of all genera within *Halobacteria* (nontype species were taken for *Halalkalicoccus*, *Halorbellus*, *Natronoarchaeum* and *Halohasta* genera because the genomes of type spices are not available) using the GTDB-tk v.1.7.0 with reference data v.202 [22]. The phylogenomic tree was constructed in the RAxML v.8.2.12 [23] with the PROTGAMMAILG model of amino acid substitution; local support values were 1000 rapid bootstrap replications. Phylogenetic trees were visualized using iTOL v.6.5.2 [24]. The whole genome-based comparisons were done as described by Sorokin et al. [25].

For functional genome analysis, genes encoding carbohydrate-active enzymes (CAZymes) were searched in genomes of AArc-St2 and AArc-St1-1 strains using dbCAN v.3.0.2 [26]. Further manual checking of the specificity of discovered glycosidases and other CAZymes were performed using BLAST against Swiss-Prot/PDB database.

# **Results and discussion**

Enrichment and isolation of pure cultures

The primary enrichments for amylolytic and fructan-utilizing natronoarchaea were performed with amylopectin (insoluble starch) or levan/inulin, respectively, in the presence of 200 mg 1<sup>-1</sup> streptomycin to suppress growth of bacteria. The enrichments from sediment fraction showed visible growth (after removal of the sediment particles by low speed centrifugation) after one week of incubation, while the brine enrichments became turbid and pinkish after 10-14 days of incubation. After several 1:100 transfers, the liquid cultures were plated and individual pink-colored colonies were transferred back to the corresponding liquid media. This procedure was repeated 2 more times to ensure the homogeneity of colony morphology. The purity and identity of

obtained cultures were verified by the 16S rRNA gene sequencing. The list of isolates is shown in **Table 1**.

Cell morphology of the isolates grown at pH 9 and 4 M total Na<sup>+</sup> with soluble starch is shown on **Fig. 1**. Cultures of all strains were dominanated by nonmotile (with only a few occasionally showing slow motility) flattish cocci with irregular contour, typical for haloarchaea, with a small fraction of flat roads ("boards"). The colonies of all isolates were colored from orange to red and concentrated cell pellets were bright red, what is also typical for aerobic haloarchaea.

# Chemotaxonomy

Membrane polar lipids and respiratory lipoquinones were analyzed in two strains, one of which represents a group 1 (AArc-St1-1<sup>T</sup>), while another a group 2 (AArc-St2<sup>T</sup>). In both, the core lipids were dominated by archaeol (AR; C<sub>20</sub>-C<sub>20</sub>) with a smaller proportion of extended archaeol (Ext-AR; C<sub>20</sub>-C<sub>25</sub>). The polar head groups of the intact polar lipids were phosphatidylglycerophosphate methylether (PGP-Me) and phosphatidylglycerol (PG). Both the core lipids and the polar head groups are very similar to the closest phylogenetic relatives of the amylolytic natronoarchaea (see below in comparative tables). The major difference is the absence of glycolipids and sulfolipids (such as phosphatidylglycerosulfate and sulfated glycosyl diethers) in natronoarchaea, which are more common in neutrophilic haloarchaea. The only respiratory lipoquinone species detected in both strains was the fully saturated MK-8:8, one of the most common in haloarchaea [15].

# Phylogenetic and genomic analyses

The genome of AArc-St2 was assembled to 20 scaffolds including one circular plasmid, while the genome of AArct-St1-1 was assembled into 32 scaffolds with no plasmids in it (Suppl. **Table S1**). Genome size was 3.26 Mbp (GC content was 51.5%) for strain AArc-St2 and 3.29 Mbp (GC content was 61%) for strain AArc-St1-1. Completeness and contamination levels for AArc-St2

genome were 100% / 0% and for AArc-St1-1 – 99.07% / 1.87%. Genomic sequences are available in NCBI GenBank database with accession numbers JAKRVX000000000 (AArc-St2) and JAKRVY000000000 (AArc-St1-1).

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Seven natronoarchaeal isolates formed two clusters on 16S rRNA gene sequence-based phylogenetic tree. Four strains of the group 1 clustered with the recently described facultatively anaerobic sulfur-reducing natronarchaeon Natranaeroarchaeum sulfidigenes which can also use starch as a substrate for growth either aerobically or anaerobically [27-28]. The three closely related isolates from the group 2 formed a novel genus-level lineage with the genera "Halalkalirubrum" and Halohasta as the nearest neighbors (around 93 and 92 % sequence identity to its type species, respectively). This potentially new genus lineage also includes multiple uncharacterized isolates from various hypersaline habitats whose 16S rRNA gene sequences were deposited recently in the GenBank. However, the 16S rRNA gene sequence-based phylogenetic tree (Fig. 2a) had relatively low bootstrap support of its inner nods within the class *Halobacteria*. Phylogenomic tree based on 122 archaeal conserved single-copy protein markers of the strains AArc-St1-1 and AArc-St2 and other haloarchaea supported the branching, obtained 16S rRNA gene sequence-based tree but showed a better resolution of the inner nods (Fig. 2b). Strain AArct-St2 formed a novel-genus branch in a distinct cluster containing genera "Halalkalirubrum", Halohasta and Halonotius. The latter three genera are currently classified in the order Haloferacales, family Halorubraceae [29]. Strain AArc-St1-1 was closely related to Natranaeroarchaeum sulfidigenum AArc-S and formed a potential new species in the genus which is currently classified within the family Natranoarchaeacea [28]. Strains AArct-St2 and AArc-St1-1 were proposed to be type species within their lineages.

For additional support of phylogenetic and phylogenomic analyses, ANI and AAI values were calculated for the genomes of AArc-St1-1 and AArc-St2 and the nearest relatives (Suppl. **Table S2 and S3**). The AAI values between strain AArc-St2 and the species of three related

haloarchaeal genera with available genome sequences ranged from 62.7% to 70.1%. These values are below the level of AAI for the representatives of the majority of genera, for which AAI results were compared [30], but it is similar to the intergenera level within the whole cluster. The ANI values varied from 70.7% to 73%. Same calculations for strain AArc-St1-1 and *Natranaeroarchaeum sulfidigenum* AArc-S showed AAI and ANI values of 90.1% and 88.8%, respectively, confirming the separate species status of AArc-St1-1 within the genus *Natranaeroarchaeum*. Although, according to the 16S rRNA gene sequence phylogeny, strain AArc-St1-2 might be sufficiently distant from the other three members of the group 1 and the type strain AArc-S<sup>T</sup> of *Natranaeroarchaeum sulfidigenes* (98.6-98.8 and 98.5%, respectively), the proposed type species of the genus *Natranaeroarchaeum*, its phenotypic properties were quite similar to the other group 1 strains implying it would be more practical to classify all four isolates of this group in a single species.

# Metabolic properties

All isolates were capable of utilizing alpha-1,4/1,6 glucans as growth substrates, including soluble starch, amylopectin (insoluble starch), glycogen, dextrin, cyclodextrin and pullulan. Furthermore, the group 1 isolates can also grow with levan (polyfructose with beta-2,6 backbond) while the group 2 strains utilized another fructan - inulin (polyfructose with beta-2,1 backbond). Two other alpha-bonded polysaccharides tested, including dextran from *Leuconostoc* and arabinan were not utilized by any of the seven isolates, as well as various beta-glucans (amorphous cellulose, xylan, xyloglucan, chitin, mannan, glucomannan, galactomannan, lichenan, laminarin and galactan). All strains can also grow on three sugar dimers including maltose,  $\alpha$ , $\alpha$ -trehalose and cellobiose. In addition, the group 2 strains were able to grow with glycerol. None of the other tested substrates, except for a weak growth with mannose, gave positive results (glucose, sucrose, galactose, arabinose, rhamnose, raffinose, aminosugars, uronic acids, xylose and arabinose, sugar alcohols,

C2-C6 organic acids, pepton). Such limited substrate profile characterize the isolates as narrow-specialized saccharolytics. Anaerobic fermentative growth with either starch, maltose or arginine was not observed. None of the isolates were capable of anaerobic respiratory growth with soluble starch or maltose as substrates, using thiosulfate, DMSO, fumarate or nitrate as the electron acceptors. However, all four group 1 isolates grew anaerobically with maltose as the electron donor and carbon source and elemental sulfur as the electron acceptor, similar to the closely related type species of the genus *Natranaeroarchaeum*. In 15 d incubation (4 M Na<sup>+</sup>, pH 9, 30°C) the following amount of sulfide was produced: 15.6 mM by AArc-St1-1, 5.8 mM by AArc-St1-2; 19 mM by AArc-St1-3 and 10.5 mM by AArc-St-lev1. In comparison, *Natranaeroarchaeum sulfidigenes* formed 30 mM sulfide in 6 days of cultivation at pH 9.5.

Two type strains tested positive for catalase reaction and TMPD-oxidase. The protease, esterase and lipase activities were negative in all strains in the spot-plate tests. Ammonium and yeast extract (but not nitrate) can serve as the N-source in cultures grown with soluble starch for both strains. Urea was only utilized by strain AArc-St1-1 consistent with the presence of the *ureABCDEFG* urease operon in the genome. Indole formation from tryptophan (Kovac's reagent test) showed a weak positive result only for strain AArc-St2. The type strains grown in liquid culture at pH 9 with soluble starch were insensitive to streptomycin, penicillin G, ampicillin, kanamycin, vancomycin and gentamicin up to 200 mg l<sup>-1</sup>. Rifampicin and chloramphenicol inhibited growth at 50 mg l<sup>-1</sup>, and tetracyclin - at 100 mg l<sup>-1</sup>.

All isolates grew well at as low Mg concentration as 1 mM, while in their sodium requirement they are typical extreme halophiles, growing optimally at 4 M total Na<sup>+</sup> and within the range from 3 to 5 M (tested at pH 8.8). The cells of isolates in both groups lyzed at salinity downshift below 2 M total Na<sup>+</sup>. The pH profiling of four cultures at 4 M total Na<sup>+</sup> showed that they are moderate alkaliphiles with optimal growth within a pH range from 8.0 to 8.8 (**Fig. 3**). The

maximum growth temperature of type strains grown with soluble starch at pH 8.5 and 4 M total Na $^+$  was 50 $^{\circ}$ C for the group 1 strains and 48 $^{\circ}$ C for the group 2 strains.

Comparative properties of the group 1 and group 2 isolates with their nearest phylogenetic relatives are shown in **Tables 2** and **3.** 

The main difference of the two groups from each other was in utilization of two different fructans and the phylogeny. The key difference of the group 1 isolates from the type species of the genus *Natrananaeroaechaeum* was their inability for anaerobic growth and inability to grow at extremely high pH values above 9.3. The main difference of the group 2 isolates from the nearest related genera is that they are the only ones isolated from soda lakes. The two out of three related genera (*Halohasta* and *Halonotius*) are definitely neutrophiles, while, despite the reported ability of "Halalkalirubrum salinum" to grow up to pH 10.5, there is a doubt about it. First, the organism is isolated from a salt lake with pH 8.5, thus being only a slightly alkaline salt (but not soda) lake. Even natronoarchaea isolated from hypersaline soda lakes with permanent pH above 10, seldom grow above pH 10.2. Secondly, the final pH values were apparently not measured during the pH profiling, which makes the reported values for the maximum pH unverified. Hence, the newly isolated amylolytic strains from soda lakes can still be considered as first obligate alkaliphilic (albeit only moderate) representatives of this group of related genera. They also differs from the other three genera in their alpha-glucan/fructan substrate specialization and the absence of glyco-and sulfo-lipids in their membranes.

## Genomic analysis

The genome search the two type strains (dbCAN) identified a set of genes typically encoding alpha-amylases and alpha-glucosidases (GH13 and 15 families) in both representatives of two groups, although in AArc-St1-1 the total number and the fraction of putative extracellular amylases are much more abundant (Suppl. **Table S4**, consensus results from HHMER/DIAMOND

tools). This is also in agreement with the growth and amylase activity results (Suppl. **Fig. 1**). These enzymes would allow utilization of a spectrum of alpha-linked glucans such as amylopectin, soluble starch, dextrin, glycogen, pullulan, maltose and trehalose as sole carbon and energy sources. In this respect strain AArc-St1-1 is highly similar to the type species of the genus *Natranaeroarchaeum* [16-17]. Furthermore genomes of both type strains contain genes coding for beta-fructosidases of the GH families 32 and 68, which is in agreement with their ability to use fructans as growth substrate. As for the other glucanases encoded in two genomes, such as the beta-endo-1,3/1,4-glucanases of GH16 and 81 families (in AArc-St1-1), and pectin lyase (PL family) in AArc-St2, none of the tested potential polysaccharide substrates for these hydrolases supported growth (laminarin, lichenan, xylan, beta-glucan, glucomannan, mannan, curdlan, pachyman or pectin).

In respect to the osmoprotection and pH homeostasis, both genomes encode a range of typical potassium import complexes (but variable in copy numbers) and a multysubunit Na $^+$ /H $^+$  antiporter of the Mrp family. Both genomes lacks genes for organic osmolyte import and synthesis, indicating that the organisms rely solely on the potassium accumulation strategy. Both strains produce catalase/peroxidase and have a haem-copper family cytochrome c terminal oxidase of the  $aa_3$  type. In addition, strain AArc-St2 has another terminal oxidase of the  $ba_3$  type (Supplementary **Table S5**). These also agree with the positive tests for catalase and oxidase in both strains.

A major difference between the two type strains was found in the presence of two types of other respiratory complexes. Strain AArc-St2 genome contains a locus apparently coding for the aerobic type of CO-dehydrogenase (Cox, most probably of the type II) lacking in AArc-St1-1. Although the capacity to oxidize CO at low concentration has been demonstrated for several haloarchaeal species [31-32], the physiological role of such potential is still unclear. One of the possibilities is CO detoxification.

The genome of AArc-St1-1 contains loci apparently encoding enzymes for sulfurdependent anaerobic respiration, which are lacking in AArc-St2: two of them are highly homologous to the PsrABCD/SseA and PhsABCD complexes encoded in the genome of type species of genus Natranaeroarchaeum responsible for sulfur- and thiosulfate-dependent anaerobic respiration, respectively [27]. In addition, there is a second encoded PsrABCD lacking sulfur transferase more homologous to the one present in sulfur-respiring Halalkaliarchaeum desulfuricum (Supplementary Table S5). In our experience, such genomic potential must enable the anaerobic sulfur respiration in strain AArc-St1-1, similar to the type species Natranaeroarchaeum sulfidigenes. And this, indeed, is directly confirmed by the growth experiments, although the activity of sulfur reduction, in general, was lower in the novel isolates than in the type species of the genus Natranaeroarchaeum. Furthermore, none of the novel group 1 isolates were capable of thiosulfate-dependent anaerobic respiration, which was a prominent trait in the type species. On the other hand, the amylolytic isolates grew much better at fully aerobic conditions, while N. sulfidigenes needed transition via microaerophilic conditions before it started to grow at aerobic conditions. This reflects the difference in enrichment conditions used to isolate these closely related but still differentially specialized species of the same genus.

Overall, on the basis of distinct phenotypic, phylogenetic and genomic features, the group 1 isolates from hypersaline soda lakes are proposed to be classified in a novel species within the genus Natranaeroarchaeum as Natranaeroarchaeum aerophilus sp. nov. (type strain AArc-St1-1), while the group 2 isolates are forming a new species in a new genus for which the name Natronocalculus amylovorans gen. nov., sp. nov. (type strain AArc-St2). The protologues for the new taxa are presented in **Tables 4** and **5**.

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#### 357 Conflict of interests

358 The authors declare that there is no conflict of interests.

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#### References

- 363 1. Oren, A. (2011) Diversity of halophiles. In: Horikoshi, K. (Ed.) Extremophiles Handbook. Springer, Tokyo, pp. 309-325.
- Andrei, A.Ş., Banciu, H.L., Oren, A. (2012) Living with salt: Metabolic and phylogenetic diversity of archaea inhabiting saline ecosystems. FEMS Microbiol. Lett. 330, 1-9.
- 367 3. Oren, A. (2013) Life at high salt concentrations. In: Rosenberg, E., DeLong, E.F., Lory, S., Stackebrandt, E., Thompson, F., (Eds.), The Prokaryotes: Prokaryotic Communities and
- Ecophysiology, Springer Berlin Heidelberg, Berlin, Heidelberg, pp. 421–40.
- 370 4. Oren, A. (2015) Halophilic microbial communities and their environments. Curr Opin Biotechnol 33, 119-24.
- 5. Grant, W.D., Jones, B.E. (2016) Bacteria, archaea and viruses of soda lakes. In: Schagerl, M., (Ed.), Soda Lakes of East Africa, Springer International Publishing, Cham, pp. 97–147.
- Oren, A., Ventosa, A., Grant, W.D. (1997) Proposed minimal standards for description of new taxa in the order *Halobacteriales*. Int J Syst Bacteriol 47, 233–238.
- Sorokin, D.Y., Toshchakov, S.V., Kolganova, T.V., Kublanov, I.V. (2015) Halo(natrono)archaea
   isolated from hypersaline lakes utilize cellulose and chitin as growth substrates. Frontiers Microbiol
   6, 942.
- Sorokin, D.Y., Khijniak, T.V., Kostrikina, N.A., Elcheninov, A.G., Toshchakov, S.V., Bale, N.J.,
   Sinnighe Damsté, J.S., Kubalnov, I.V. (2018) *Natronobiforma cellulositropha* gen. nov., sp. nov., a
   novel haloalkaliphilic member of the family *Natrialbaceae* (class *Halobacteria*) from hypersaline
- 382 alkaline lakes. Syst Appl Microbiol 41, 355–362.
- 383 9. Sorokin, D.Y., Elcheninov. A.G., Toshchakov, S.V., Bale, N.J., Sinninghe Damsté, J.S., Khijniak,
- T.V., Kublanov, I.V. (2019) Natrarchaeobius chitinivorans gen. nov., sp. nov., and
- 385 *Natrarchaeobius halalkaliphilus* sp. nov., alkaliphilic, chitin-utilizing haloarchaea from hypersaline alkaline lakes. Syst Appl Microbiol 42, 309-318.
- 387 10. Kobayashi, T., Kanai H., Hayashi T., Akiba T., Akaboshi R., Horikoshi K. (1992) Haloalkaliphilic
- 388 maltotriose-forming a-amylase from the archaebacterium Natronococcus sp. strain Ah-36. J
- 389 Bacteriol 174, 3439-3444.
- 390 11. Kanai, H., Yashi, K., T, Aono, R., Kudo, T. (1995) *Natronococcus amylolyticus* sp. nov., a haloalkaliphilic archaeon. Int J Syst Bacteriol 45, 762-766.
- 392 12. Pfennig, N., Lippert, K.D. (1966) Über das Vitamin B12-Bedürfnis phototropher Schwefelbakterien. Arch Mikrobiol 55, 245-256.

- 394 13. Sorokin, D.Y., Yakimov, M.M., Messina, E., Merkel, A.Y., Koenen, M., Bale, N.J., Sinninghe
- Damsté, J.S. (2021) Halapricum desulfuricans sp. nov., carbohydrate-utilizing sulfur-reducing
- haloarchaea from hypersaline lakes. Syst Appl Microbiol 44, 126249.
- 397 14. Bale, N.J., Sorokin, D.Y., Hopmans, E.C., Koenen, M., Rijpstra, W.I.C., Villanueva, L., Wienk, H.,
- 398 Sinninghe Damsté, J.S. (2019) New insights into the polar lipid composition of halophilic
- euryarchaea from hyper saline lakes. Front Microbiol 10, 377.
- 400 15. Elling, F.J., Becker, K.W., Könneke, M., Schröder J.M., Kellermann, M.Y., Thomm, M., Hinrichs,
- 401 K.-U. (2016) Respiratory quinones in Archaea: phylogenetic distribution and application as
- biomarkers in the marine environment. Environ. Microbiol 18, 692–707.
- 403 16. Bankevich, A., Nurk, S., Antipov, D., Gurevich, A.A., Dvorkin, M., Kulikov, A.S., Lesin, V.M.,
- Nikolenko, S.I., Pham, S., Prjibelski, A.D., Pyshkin, A.V., Sirotkin, A.V., Vyahhi, N., Tesler, G.,
- Alekseyev, M.A., Pevzner, P.A. (2012) SPAdes: A New Genome Assembly Algorithm and Its
- 406 Applications to Single-Cell Sequencing. J Comput Biol 19, 455–477.
- 407 17. Wick, R.R., Judd, L.M., Gorrie, C.L., Holt, K.E. (2017) Unicycler: Resolving bacterial genome
- 408 assemblies from short and long sequencing reads. PLoS Comput Biol 13, 1-22.
- 409 18. Gurevich, A., Saveliev, V., Vyahhi, N., Tesler, G. (2013) QUAST: Quality assessment tool for
- genome assemblies. Bioinformatics 29, 1072–1075.
- 411 19. Mikheenko, A., Prjibelski, A., Saveliev, V., Antipov, D., Gurevich, A. (2018) Versatile genome
- 412 assembly evaluation with QUAST-LG. Bioinformatics 34, i142–50.
- 413 20. Parks, D.H., Imelfort, M., Skennerton, C.T., Hugenholtz, P., Tyson, G.W. (2015) CheckM:
- Assessing the quality of microbial genomes recovered from isolates, single cells, and metagenomes.
- 415 Genome Res 25, 1043–55.
- 416 21. Rinke C, Chuvochina M, Mussig AJ, P-A Chaumeil, Davín AA, Waite DW, Whitman WB, Parks
- DH, Hugenholtz P. (2021) A standardized archaeal taxonomy for the Genome Taxonomy Database.
- 418 Nat Microbiol 6, 946-959.
- 419 22. Parks, D.H., Chuvochina, M., Waite, D.W., Rinke, C., Skarshewski, A., Chaumeil, P.-A.,
- Hugenholtz, P. (2018) A standardized bacterial taxonomy based on genome phylogeny substantially
- revises the tree of life. Nat Biotechnol 36(10), 996–1004.
- 422 23. Stamatakis, A. (2014) RAxML version 8: A tool for phylogenetic analysis and post-analysis of
- 423 large phylogenies. Bioinformatics 30, 1312–1313.
- 424 24. Letunic, I., Bork, P. (2019) Interactive Tree Of Life (iTOL) v4: recent updates and new
- developments. Nucleic Acids Res 47, W256-W259.
- 426 25. Sorokin, D.Y., Elcheninov, A.G., Khijniak, T. V., Zaharycheva, A.P., Boueva, O. V., Ariskina, E.
- V., Bunk, B., Spröer, C., Evtushenko, L.I., Kublanov, I. V., Hahnke, R.L. (2022)
- Natronosporangium hydrolyticum gen. nov., sp. nov., a haloalkaliphilic polyhydrolytic
- 429 actinobacterium from a soda solonchak soil in Central Asia. Syst Appl Microbiol 45, 126307.

- 430 26. Zhang, H., Yohe, T., Huang, L., Entwistle, S., Wu, P., Yang, Z., Busk, P.K., Xu, Y., Yin, Y. (2018)
- dbCAN2: a meta server for automated carbohydrate-active enzyme annotation. Nucleic Acids Res
- 432 46(W1), W95–101.
- 433 27. Sorokin, D.Y., Messina, E., Smedile, F., La Cono, V., Hallsworth, J.E. (2021) Carbohydrate-
- dependent sulfur respiration in halo(alkali)philic euryarchaea from hypersaline lakes. Environ
- 435 Microbiol 23, 3779-3808.
- 436 28. Sorokin, D.Y., Yakimov, M.M., Messina, E., Merkel, A.Y., Koenen, M., Bale N.J., Sinninghe
- Damsté J.S. (2022) Natranaerarchaeum desulfuricum gen. nov., sp. nov., carbohydrate-utilizing
- sulfur-respiring haloarchaeon from hypersaline lakes and proposal of a new family
- Natronoarchaeaceae fam. nov. in the order Halobacteriales. Int J Syst Evol Microbiol 72,
- 440 10.1099/ijsem.0.005332.
- 441 29. Gupta, R.S., Naushad, S., Fabros, R., Adeolu, M. (2016) A phylogenomic reappraisal of family-
- level divisions within the class *Halobacteria*: proposal to divide the order *Halobacteriales* into the
- families Halobacteriaceae, Haloarculaceae fam. nov., and Halococcaceae fam. nov., and the order
- 444 Haloferacales into the families, Haloferacaceae and Halorubraceae fam nov. Antonie van
- Leeuwenhoek 109, 565-587.
- 446 30. Luo, C., Rodriguez-R, L.M., Konstantinidis, K.T. (2014) MyTaxa: An advanced taxonomic
- classifier for genomic and metagenomic sequences. Nucleic Acids Res 42, e73.
- 448 31. King, G.M. (2015) Carbon monoxide as a metabolic energy source for extremely halophilic
- microbes: Implications for microbial activity in Mars regolith. Proc Nat Ac Sci USA 112, 4465-
- 450 4470.
- 451 32. Myers, M.R., King, G.M. (2020) Halobacterium bonnevillei sp. nov., Halobaculum saliterrae sp.
- nov. and *Halovenus carboxidivorans* sp. nov., three novel carbon monoxide-oxidizing Halobacteria
- from saline crusts and soils. Int J Syst Evol Microbiol 70, 4261-4268.
- 454 33. Shimane, Y., Hatada, Y., Minegishi, H., Mizuki, T., Echigo, A. (2010) Natronoarchaeum
- 455 mannanilyticum gen. nov., sp. nov., an aerobic, extremely halophilic archaeon isolated from
- 456 commercial salt. Int J Syst Evol Microbiol 60, 2529–2534.
- 457 34. Zhenqiang Zuo . Dahe Zhao . Jian Zhou . Jing Han . Hua Xiang (2021) Halalkalirubrum salinum
- gen. nov., sp. nov., a halophilic archaeon isolated from a saline lake. Ant van Leeuwenhoek 114,
- 459 83–94.
- 460 35. Mou, Y.-Z., Qiu, X.-X., Zhao, M.-L., Cui, H.-L., Oh, D., Dyall-Smith, M.L. (2012) Halohasta
- litorea gen. nov. sp. nov., and Halohasta litchfieldiae sp. nov., isolated from the Daliang
- aguaculture farm, China and from Deep Lake, Antarctica, respectively. Extremophiles 16, 895-901.
- 463 36. Durán-Viseras, A., Andrei A.-S., Ghai, R., Sánchez-Porro, C., Ventosa, A. (2019) New Halonotius
- species provide genomics-based insights into cobalamin synthesis in haloarchaea. Front Microbiol
- 465 10, 1928.

 **Table 1**. List of pure cultures of natronoarchaea enriched and isolated from hypersaline soda lakes in southwestern Siberia with amylopecting and fructans

<u> </u>			<b>7</b> 1	C
Strain	Source	Enrichment	Phylogenetic	Closest relative
		substrate	group	
AArc-St1-1 <sup>T</sup>	Sediments	Amylopectin	Group 1	Natranaeroarchaeum
AArc-St1-2	Brines			sulfidigenes
AArc-St1-3	Brines			
AArc-lev1	Sediments	Levan		
AArc-St2 <sup>T</sup>	Sediments	Amylopectin	Group 2	"Halalkalirubrum
AArc-St3	Brines			halophilum"
AArc-in2	Sediments	Inulin		

**Table 2**. Comparative properties of amylolytic natronoarchaea of group 1 with the type species of the nearest phylogenetically related genera [28, 33]

Property	"Natranaeroarchaeum	Natranaeroarchaeum	Natronoarchaeum
	aerophilus"	sulfidigenes	mannanilyticum
	(4 strains)	JCM 34033 <sup>T</sup>	JCM 16328 <sup>T</sup>
Cell morphology	flat pleomorphic,	flat pleomorphic,	pleomorphic,
	motility not observed	motile	nonmotile
Pigmentation	red	red (aerobic); pink	red
		(anaerobic)	
PHA accumulation	-	-	-
Aerobic growth	+	+	+
Anaerobic growth by:			
sugar fermentation	-	-	-
sulfur/thiosulfate respiration	+ (with sulfur)	+ (with sulfur and $S_2O_{32}$ -)	-
sulfoxide respiration	-	-	-
Number of Psr/Phs operons	3	2	$0^*$
in genomes			
<i>e</i> -donors for anaerobic	glucose, maltose	sugars, starch, glycerol	-
growth			
Substrates for aerobic growth	starch-like alpha-glucans,	sugars, starch,	lactose, raffinose, sucrose,
	levan, maltose, cellobiose,	yeast extract	maltose, cellobiose, starch,
	trehalose		galactomannan, pyruvate, lactate,
			glutamate, yeast extract, peptone
Amylase	+	+	+
Esterase/lipase	-	- (tributyrin/olive oil)	- (Tween 80)
Protease	-	- (gelatin, casein)	- (gelatin)
Catalase/oxidase	+/+	+/+(w)	-/ +(w)
Indole from tryptophane	2050(40)	2.5.4.5.(2.5)	+
Salinity range (opt.) (M Na <sup>+</sup> )	3.0-5.0 (4.0)	2.5-4.5 (3.5)	1.6-4.2 (2.5-3.2)
pH range (opt.)	7.2-9.3 (8.0-8.8)	8.5-10.2 (9.5-9.7)	6.0-9.5 (8.5-9.0) 55
Temperature max (°C) Core lipids	50 (at pH 8.5) C <sub>20</sub> -C <sub>20</sub> , C <sub>20</sub> -C <sub>25</sub> DGE	45 (at pH 9) C <sub>20</sub> -C <sub>20</sub> , C <sub>20</sub> -C <sub>25</sub> DGE	NR
Intact membrane polar lipids:	C20-C20, C20-C25 DGE	C20-C20, C20-C25 DGE	IVK
phospholipids	PG, PGP-Me	PG, PGP-Me	PG, PGP-Me, PGP,
glycolipids/sulfolipids	- 1 G, 1 G1 -IVIE	- 1 G, 1 G1 -1vie	S <sub>2</sub> -DGDE
Respiratory lipoquinones	MK-8:8	MK-8:8	NR
DNA G+C	61.0 (genome)	60.8% (genome)	63.0 (mol%)
Type of hypersaline habitat		e soda lakes	Marine solar saltern
ND not reported (v) verichle			

NR, not reported; (v) - variable property in different species of the same genus; w (weak); Psr/Phs - polysulfide/thiosulfate reductase; \*genome of *N. phillipinensis*. Lipids: (PG) phosphatidylglycerol, phosphatidylglycero-phosphate (PGP), (PGP-Me) phosphatidylglycerophosphate methyl ester, disulfated diglycosyl diether (S<sub>2</sub>-DGDE), (DGE) - dialkyl glycerol ether.

Table 3. Comparative properties of group 2 isolates with nearest phylogenetic relatives [34-36].

Property	"Natranocalculus amylovorans" (3 isolates)	"Halalkalirubrum halophilum"	Halohasta (2 species)	Halonotius (4 species)
Cell morphology	flat pleomorphic nonmotile	pleomorphic, nonmotile	rods, motile	polymorphic rods, motility (V)
Di	red	red	red	red
Pigmentation PHA accumulation	red -	NR	NR	red
Anaerobic growth by:				
sugar fermentation	-	NR	NR	NR
sulfur respiration	-	NR	NR	NR
DMSO respiration	-	-	-	+(1 species)
Growth substrates				
carbohydrates:	starch-like alpha-	glucose, maltose,	glucose, sucrose;	glucose, arabinose,
	glucans, inulin,	fructose, sorbose,	mannose, galactose,	fructose, galactose,
	maltose, cellobiose,	lactose, xylose,	lactose, maltose (all	sucrose, maltose,
	trehalose, glycerol	mannitol, sorbitol	V)	raffinose, xylose, mannitol, sorbitol, glycerol, (all V)
organic acids:	none	acetate, pyruvate, lactate, fumarate, succinate, citrate	pyruvate, lactate, succinate, malate, fumarate, citrate (V);	pyruvate, citrate, tartrate (all V)
Amylase	+ (soluble starch)	- (soluble starch)	- (soluble starch)	- (soluble starch)
Esterase/lipase	- (tributyrin/olive oil)	- (Tweens)	- (Tween 80)	- (Tween 80)
Protease	- (gelatin, casein)	- (gelatin, casein)	- (gelatin, casein)	- (gelatin, casein)
Catalase/oxidase	+/+	+/+	+/V	V/V
Indole from tryptophane	+(w)	+	-	NR
Salinity range (opt.) M Na <sup>+</sup>	3-5 (4.0)	1.9-4.2 (2.5)	2.0-4.7 (2.5-3.0)	2.5-6.0**(3.0-4.0)
Mg <sup>2+</sup> demand	low	low	high	high
pH range (opt.)	7.2-9.3 (8.5-8.8)	7.0-10.5* (8.5-9.5)	5.5-9.0* (7.0-7.5)	5.0-9.0 (7.0-7.5)
Temperature max (°C)	48 (at pH 8.5)	42	45-50	45-50
Core lipids	C <sub>20</sub> -C <sub>20</sub> , C <sub>20</sub> -C <sub>25</sub> DGE	NR	NR	NR
Intact membrane polar lipids:				
phospholipids:	PG, PGP-Me	PG, PGP-Me	PG, PGP-Me, PA	PG, PGP-Me
glycolipids:	-	1, unidentified	S-DGD-1	S-DGD-1
sulfolipids:	-	PGS		PGS (V)
Respiratory lipoquinones	MK-8:8	NR	NR	NR
DNA G+C (% genomic)	51.5 (type strain)	58.4 (type strain)	58.8 (type species)	59.7-62.7 (4 species)
Type of hypersaline habitat	soda lakes	salt lake	solar saltern	solar saltern, saline soils

NR, not reported; (V), variable property in different species of the same genus; \* actual final pH values were not measured; \*\* reported for the type species, but not verified in any further research; PA, phosphatidic acid; PGS, phosphatidylglycerol sulfate; S-DGD-1, sulfated mannosyl glucosyl diether; other abbreviations (see **Table 2**).

<b>Table 4</b> . <i>Natranaeroarchaeum ae</i> Parameter	Species: Natranaeroarchaeum aerophilus sp. nov.
Author	Dimitry Y. Sorokin
Species name	aerophilus
Genus name	Natranaeroarchaeum
Specific epithet	aerophilus
Species status (SPST)	sp. nov.
Etymology	a.e.ro'phi.lus Gr. masc. n. aer, air; N.L. masc. adj. philus (from Gr. masc. adj. philos), friend, loving; N.L. masc. adj. aerophilus, air-loving.
Description of the new taxon	The cells are angular, flat, polymorphyc coccoids or rods, mostly nonmotile, variable in size from 1 to 3 µm. The cells lyze in hypotonic solutions below 1 M NaCl. produces red carotenoids. The core membrane diether lipids are composed of $C_{20}$ - $C_{20}$ DGE (extended archaeol). The polar lipid head groups include phosphatidylglycerolphosphate methyl ester (PGP-Me) and phosphatidylglycerol (PG). The dominant respiratory quinone is MK-8:8. Saccharolytic with limited substrate profile including several starch-like alpha-glucans, levan, maltose, trehalose and cellobiose. Capable of anaerobic sulfogenic growth with glucose and maltose as electron donors and sulfur as electron acceptor. Ammonium, urea and yeast extract serve as the N-source. Oxidase and catalase positive. Maximum growth temperature is 50°C. Extremely halophilic with a range of total Na <sup>+</sup> for growth from 3 to 5 M (optimum at 4 M) and moderately alkaliphilic, with a pH range for growth from 7.2 to 9.3 (optimum at 8.0-8.8). The G + C content of the DNA is 61.0% (genome of the type strain). Habitat - aerobic sediments and brines of hypersaline soda lakes. The type strain (AArc-St1-1 <sup>T</sup> =JCM 32519 <sup>T</sup> =UQM 41458 <sup>T</sup> ) was isolated aerobic sediments of hypersaline soda lakes in Kulunda Steppe (Altai, Russia). The species also includes other three closely related strains isolated from the same area. The draft genome of type strain is deposited in the GenBank under accession number JAKRVY0000000000.
Authors	Dimitry Y. Sorokin, Alexander G. Elcheninov, Tatjana V. Khizhniak, Michel Koenen, Nicole J. Bale, Jaap S. Sinninghe Damsté, Ilya V. Kublanov
Title	Natronocalculus amylovorans gen. nov., sp. nov., and Natranaeroarchaeum aerophilus sp. nov., dominant culturable amylolytic natronoarchaea from hypersaline soda lakes in southwestern Siberia.
Journal	Systematic and Applied Microbiology
Corresponding author	Dimitry Y. Sorokin
E-mail of corresponding author	soroc@inmi.ru; d.sorokin@tudelft
Designation of the type strain	AArc-St1-1
Strain collection numbers	JCM 32519 <sup>T</sup> =UQM 41458 <sup>T</sup>
16S rRNA gene accession numbers Genome accession numbers	MG584707- MG584709; ON003450 JAKRVY000000000 (type strain)
Genome accession numbers Genome status	Draft
G+C, %	61.0 (genome of type strain)
Country of origin	Russian Federation
Region of origin	Altai region
Date of isolation	2016
Source of isolation	Surface aerobic sediments from hypersaline soda lakes
Sampling dates	2015-07-07
Geographic location	S-W Siberia, Kulunda Steppe; southern Russia
Latitude	51°39' N; 49°10' N; 48°14' N
Longtitude	79°48' E; 46°39' E; 46°35' E
Depth	0-2 cm
Temperature of the sample	20°C
pH of the sample	10-11
Salinity of the sample	18-36%
Number of strains in study Source of isolation of non-type strains	Surface aerobic sediments and brines from hypersaline soda lakes, S-W Siberia, Kulunda Steppe; southern Russia
Growth medium, incubation conditions	4 M total Na <sup>+</sup> , pH 9; incubation - 37°C; starch as substrates; aerobic
Conditions of preservation	Deep freezing in 15% glycerol (v/v)
Gram stain	Negative
Cell shape	Pleomorphic flat coccoids
Cell size	0.8-2 µm in diameter
Motility (MOTY)	Mostly nonmotile
Sporulation	None
Colony morphology Temperature range for growth	Flat, compact, max. 2 mm, red
Lowest temperature for growth	nd nd
Highest temperature for growth	50
Optimal temperature for growth	37-40
Lowest pH for growth	7.2
Highest pH for growth	9.3
Optimum pH for growth	8-8.8
pH category	Moderately alkaliphilic
Lowest Na <sup>+</sup> concentration for growth	3.0 M
Highest Na <sup>+</sup> concentration for growth	5.0 M
Optimum salt concentration for growth	4.0 M total Na <sup>+</sup>
Other salts important for growth	KCl; Na-carbonates
Salinity category	Extremely halophilic
Relation to oxygen	Facultatively anaerobic
O <sub>2</sub> conditions for strain testing Carbon source used (class)	Fully aerobic Carbohydrates
Carbon source used (Class)	Carbonyurates

Specific compounds	Starch-like alpha-glucans, levan
Nitrogen source	Ammonium, urea, yeast extract
Terminal electron acceptor	$O_2$ and $S_8$
Energy metabolism	Chemoorganotrophic
Phospholipids	Core membrane lipids are C <sub>20</sub> -C <sub>20</sub> DGE (archaeol) and C <sub>20</sub> -C <sub>25</sub> DGE (extended archaeol). Polar head groups are phosphatidylglycerophosphate methylester (PGP-Me) and phosphatidylglycerol (PG)
Respiratory lipoquinones	MK-8:8
Glycolipids (GLYC)	-
Habitat (HABT)	Hypersaline soda lakes
Extraordinary feautures (EXTR)	Narrowly specialized amylolytics

# Table 5. Natronocalculus amylovorans: protologue

Parameter	Genus: Natronocalculus gen. nov.	Species: Natronocalculus amylovorans sp. nov.	
Parameter Author (AUTE)	Dimitry Y. Sorokin	Species. Ivaironocaicuius amyiovorans sp. nov.	
Species name (SPNA)	Diffility 1. Solokiii	amylovorans	
Genus name (GENA)	Natronocalculus	uniyiovoruns	
Specific epithet (SPEP)	-	amylovorans	
Species status (SPST)	-	sp. nov.	
Etymology (GETY/SPTY)	Na.tro.no.cal'cu.lus N.L. neut. n. natron, arbitrarily derived from the Arabic n. natrun or natron soda; L. masc. n. calculus, pebble, gravel; N.L. masc. n. Natronocalculus, soda loving pebble-shaped cells	a.my.lo.vo'rans. Gr. neut. n. <i>amylon</i> , starch; L. inf. v. <i>vorare</i> , to devour; N.L. part. adj. <i>amylovorans</i> , eating starch	
Type species of the genus (TYPE)	Natronocalculus amylovorans	yes	
Description of new taxon	Obligately aerobic and organotrophic member of <i>Halobacteriales</i> narrowly specialized on utilization of starch-like polymers. Core lipids are dominated by archaeol and extended archaeol with PGP-Me and PG polar groups. Glycolipids are not present. MK-8:8 is the dominant lipoquinone. Extremely halophilic and moderately alkaliphilic inhabitans of hypersaline soda lakes. The three-letter abbreviation is Ncl.	The cells are angular, flat, polymorphyc coccoids or rods, nonmotile, from 0.8 to 2.0 μm. The cells lyze in hypotonic solutions below 1 M NaCl. Colonies are orange-red. The core membrane diether lipids include of C <sub>20</sub> -C <sub>20</sub> DGE (archaeol) and C <sub>20</sub> -C <sub>25</sub> DGE (extended archaeol). The polar lipid head groups consists of hosphatidylglycerolphosphate methyl ester (PGP-Me) and phosphatidylglycerol (PG). The dominant respiratory menaquinone is MK-8:8. Obligately aerobic saccharolytic with limited substrate profile including several starch-like alpha-glucans, inulin, maltose, trehalose, cellobiose and glycerol. Ammonium and yeast extract serve as the N-source. Oxidase and catalase positive. Maximum growth temperature is 48°C. Extremely halophilic with a range of total Na* for growth from 3 to 5 M (optimum at 4 M) and moderately alkaliphilic, with a pH range for growth from 7.2 to 9.3 (optimum at 8.5-8.8). The G + C content of the DNA is 51.5% ( genome of the type strain). Habitat - aerobic sediments and brines of hypersaline soda lakes. The type strain (AArc-St2 <sup>T</sup> =JCM 32475 <sup>T</sup> =UQM 41459 <sup>T</sup> ) was isolated from aerobic sediments of hypersaline soda lakes in Kulunda Steppe (Altai, Russia). The species also includes 2 other closely related strains isolated from the same area. The draft genome of type strain is deposited in the GenBank under accession number JAKRVX0000000000.	
Authors (AUT)	Dimitry Y. Sorokin, Alexander G. Elcheninov, Tatjana V. Khizhniak, Michel Koenen, Nicole J. Bale, Jaap S. Sinninghe Damsté, Ilya V. Kublanov		
Title (TITL)		p. nov., and <i>Natranaeroarchaeum aerophilus</i> sp. nov., haea from hypersaline soda lakes in southwestern	
Journal (JOUR)	Systematic and Applied Microbiology		
Corresponding author (COAU)	Dimitry Y. Sorokin		
E-mail of corresponding author (EMAU)	d.sorokin@tudelft; soroc@inmi.ru		
Strain collection numbers (COLN)	-	JCM 32475; UQM 41459	
16S rRNA gene accession number (16 SR)	-	MG584710; ON000203; ON000205	
Genome accession numbers		JAKRVX000000000	
Genome status (GSTA)		Draft	
GC mol % (GGCM)	-	51.5 (genome type strain)	
Country of origin (COUN)	Russian Federation	Russian Federation	
Region of origin (REGI)	-	Altai region	
Date of isolation (DATI) Source of isolation (SOUR)	- Hypersaline soda lakes	2016 Surface sediments and brines of hypersaline soda lakes in southwestern Siberia	
Sampling dates (DATS)	2015	2015	
Geographic location (GEOL)	S-W Siberia	S-W Siberia	
Latitude (LATI)	-	51°39' N; 49°10' N; 48°14' N	
Longtitude (LONG)	-	79°48' E; 46°39' E; 46°35' E	
Depth (DEPT)		0-2 cm	
Temperature of the sample (TEMS)		20°C	
pH of the sample (PHSA)		10-11	
Salinity of the sample (SALS)		18-36%	
Number of strains in study (NSTR)	3	3	
Source of isolation of non-type strains (SAMP)	-	Surface sediments and brines of hypersaline soda lakes in southwestern Siberia	
Growth medium, incubation conditions		4 M total Na <sup>+</sup> , pH 9; incubation - 37°C; starch as	
(CULT)	B 6 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	substrates; aerobic	
Conditions of preservation (PRES)	Deep freezing in 15% glycerol (v/v)		
Gram stain (GRAM)	negative		
Cell shape (CSHA)	Pleomorphic, from flat irregular coccoids		
Cell size (CSZI)	-	0.8-2 μm in diameter	

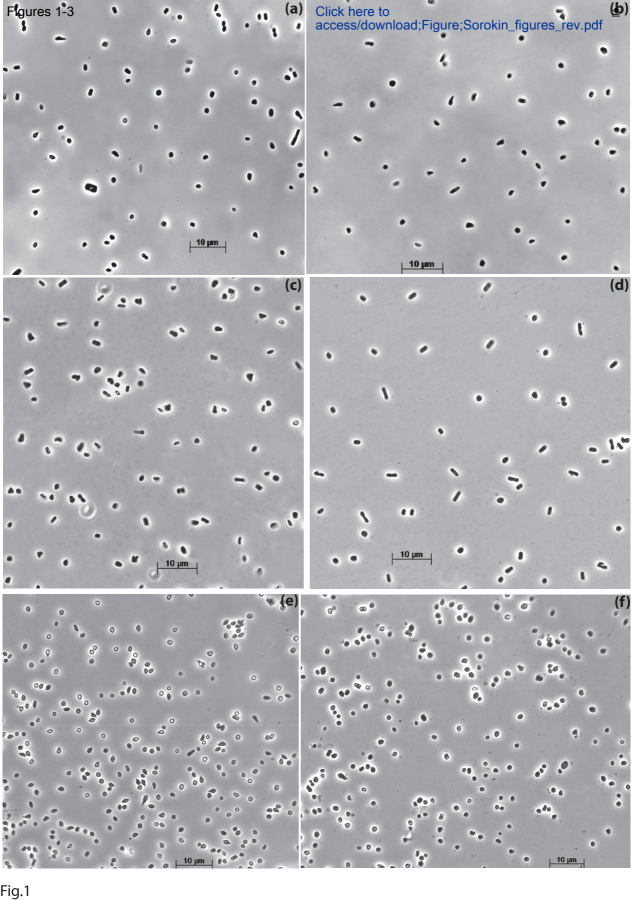
Motility (MOTY)	- nonmotile		
Sporulation (SPOR)	none		
Colony morphology (COLM)	Pink-orange, up to 2 mm, flat		
Temperature range for growth (TEMR)			
Lowest temperature for growth (TEML)			
Highest temperature for growth(TEMH)	48 (at pH 8.5)		
Optimal temperature for growth	37-40°C	-	
(TEMO)			
Lowest pH for growth (PHLO)	7.2		
Highest pH for growth (PHHI)	9.3		
Optimum pH for growth (PHOP)	8.5-8.8		
pH category (PHCA)	alkaliphile (optimum > 8.5)		
Lowest NaCl concentration for growth	3.0 M total Na <sup>+</sup>		
(SALL)			
Highest NaCl concentration for growth	5 M total Na <sup>+</sup>		
(SALH)			
Optimum salt concentration for growth	4.0 M total Na <sup>+</sup>		
(SALO)			
Other salts important for growth	Sodium carbonates		
Salinity category (SALC)	extremely halophilic		
Relation to oxygene (OREL)	aerobe		
O <sub>2</sub> conditions for strain testing (OCON)	aerobic		
Carbon source used (class) (CSUC)	carbohydrates		
Specific compounds (CSUC)	Starch-like alpha glucans and inulin		
Nitrogen source (NSOU)	Ammonium, yeast extract		
Terminal electron acceptor (ELAC)	02		
Energy metabolism (EMET)	chemoorganotrophic		
Phospholipids (PHOS)	Core membrane lipids are archaeol (C <sub>20</sub> -C <sub>20</sub> DGE) and extended archaeol (C <sub>20</sub> -C <sub>25</sub> DGE)		
	Polar lipids are phosphatidylglycerophosphate methyl ester (PGP-Me) and phosphatidylglycerol (PG)		
Glycolipids (GLYC)	((U)		
Respiratory lipoquinones	- MK8:8		
Habitat (HABT)	Hypersaline soda lakes		
nauliai (nad i)	rtypersamie soua takes		

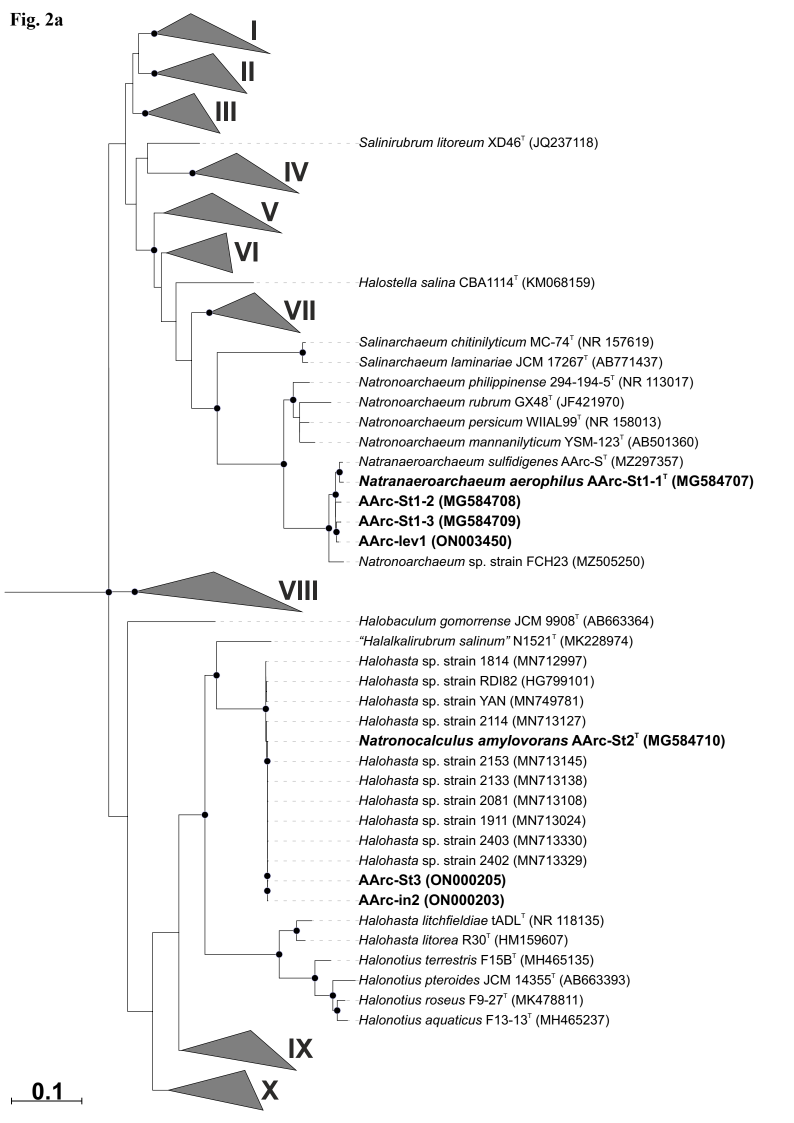
#### Legends to the figures

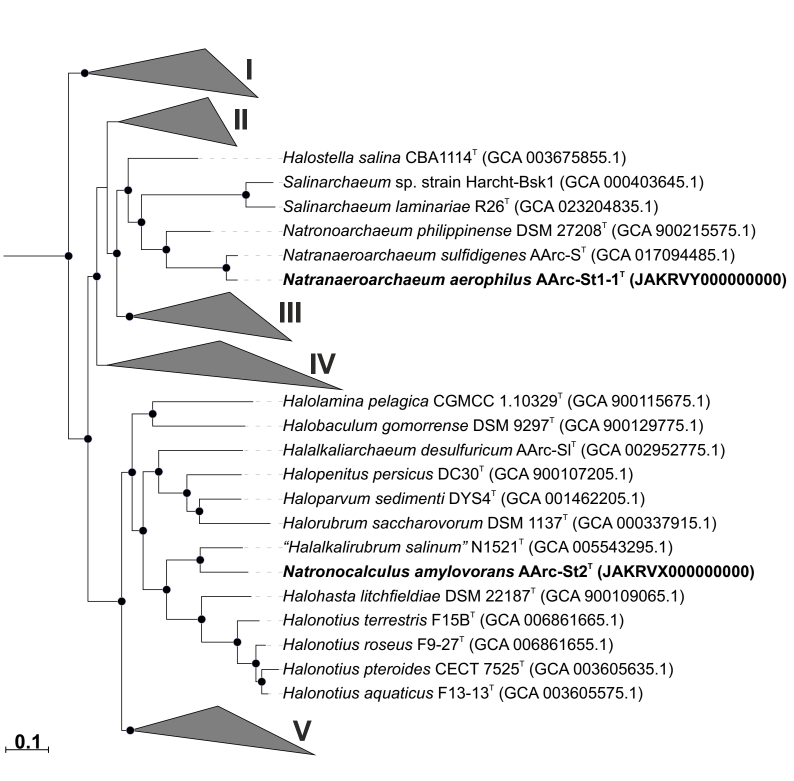
499 500

- 501 Fig. 1. Cell morphology (phase contrast microphotograps) of starch-utilizing natronoarchaea growing
- 502 aerobically at 4 M total Na<sup>+</sup>, pH 9 and 37°C. (a-d), group 1, including strains AArc-St1-1<sup>T</sup>, AArc-St1-
- 503 2, AArc-St1-3 and AArc-lev11, respectively. (e-f), group 2, including strains AArc-St2<sup>T</sup> and AArc-
- 504
- 505 Figure 2. Phylogeny of amylolytic natronoarchaea.
- 506 (a) Figure X. (a) 16S rRNA gene sequence-based maximum-likelihood phylogenetic tree, showing the position
- of AArc-St2<sup>T</sup> and AArc-St1-1<sup>T</sup> (in bold) within the *Halobacteria* class. The black circles at nodes indicate that 507
- 508 the percentage of corresponding support values was above 50. Archaeoglobus fulgidus VC-16<sup>T</sup>, Methanocella
- 509 paludicola SANAE<sup>T</sup>, Methanothermobacter thermautotrophicus Delta H<sup>T</sup> were used as an outgroup (not shown).
- 510 Species in clusters: I (Halodesulfurarchaeum, Halanaeroarchaeum, Halarchaeum, Halobacterium,
- 511 Salarchaeum, Halocalculus): II (Halomicroarcula, Haloarcula, Halorientalis, Halorhabdus, Halococcoides,
- 512 Halapricum, Salinirussus, Halovenus); III (Haloglomus, Natronomonas, Halosegnis, Salinirubellus,
- 513 Halomarina, Halocatena); IV (Halopelagius, Haloferax, Halogeometricum, Haloquadratum, Halobellus); V
- 514 (Halococcus, Halalkalicoccus, Haloarchaeobius, Halorubellus); VI (Halorussus, Halomicrococcus,
- 515 Haladaptatus); VII (Saliphagus, Natribaculum, Halovarius, Natronococcus, Halovivax, Natronobiforma, 516 Halostagnicola, Natronobacterium, Halopiger, Halobiforma, Natrarchaeobaculum, Natronolimnohabitans,
- 517
- Natronolimnobius, Natronorubrum, Natrinema, Haloterrigena, Natrialba, Natrarchaeobius, Salinadaptatus); 518
- VIII (Halosimplex, Salinibaculum, Halosiccatus, Halomicrobium); IX (Salinigranum, Haloplanus, Halobium,
- 519 Halegenticoccus, Halogranum, Haloprofundus, Halolamina); X (Halalkaliarchaeum, Halopenitus, Halorubrum,
- 520 Haloparvum).
- 521 (b) Maximum likelihood phylogenetic tree based on concatenated alignment of 122 conserved archaeal proteins 522
- and showing position of strains AArc-St2<sup>T</sup> and AArc-St1-1<sup>T</sup> (in bold) within the class *Halobacteria*. The branch 523 lengths correspond to the number of substitutions per site with corrections associated with the models. The black
- 524 circles at nodes indicate that the percentage of corresponding support values was above 50. Archaeoglobus 525 fulgidus VC-16<sup>T</sup>, Methanocella paludicola SANAE<sup>T</sup> and Methanothermobacter thermautotrophicus Delta H<sup>T</sup>
- 526 were used as an outgroup (not shown). Species in clusters: I (Halanaeroarchaeum, Halodesulfurarchaeum,
- 527 Halarchaeum, Halobacterium, Salarchaeum, Halocalculus); II (Halalkalicoccus, Halorussus, Halomicrococcus,
- 528
- Haladaptatus); III (Haloarchaeobius, Halorubellus, Halovivax, Saliphagus, Natronobiforma, Halostagnicola, 529
- Natrarchaeobius, Natrarchaeobaculum, Salinadaptatus, Halopiger, Natronolimnobius, Natronobacterium, 530
- Halobiforma, Natrialba, Natronococcus, Natrinema, Haloterrigena, Natronorubrum, Natronolimnohabitans); 531 IV (Halococcus, Halocatena, Halomarina, Natronomonas, Haloglomus, Halosegnis, Halorientalis, Halapricum,
- 532 Salinirussus, Salinibaculum, Halovenus, Halosimplex, Halococcoides, Halorhabdus, Halomicrobium,
- 533 Halosiccatus, Halomicroarcula, Haloarcula); V (Haloplanus, Salinirubrum, Haloprofundus, Halegenticoccus,
- 534 Halogranum, Salinigranum, Haloferax, Halopelagius, Halogeometricum, Halobellus, Haloquadratum).

- 537 Fig. 3. pH profiles for growth with soluble starch in amylolytic natronoarchaea at 4 M total Na<sup>+</sup> and
- 538 37°C. Actual final pH are shown. The results are average from two parallel incubations.







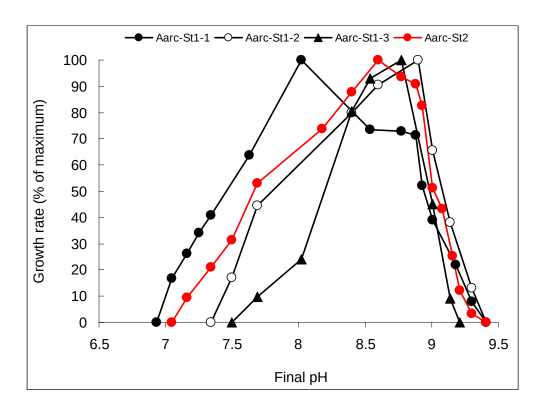


Fig. 3

e-Component

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