A Bacterium That Can Grow by Using **Arsenic Instead of Phosphorus**

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Life is mostly composed of the elements carbon, hydrogen, nitrogen, oxygen, sulfur, and phosphorus. Although these six elements make up nucleic acids, proteins, and lipids and thus the bulk of living matter, it is theoretically possible that some other elements in the periodic table could serve the same functions. Here, we describe a bacterium, strain GFA]-1 of the Halomonadaceae, isolated from Mono Lake, California, that is able to substitute arsenic for phosphorus to sustain its growth. Our data show evidence for arsenate in macromolecules that normally contain phosphate, most notably nucleic acids and proteins. Exchange of one of the major bio-elements may have profound evolutionary and geochemical importance.

iological dependence on the six major nutrient elements carbon, hydrogen, nitrogen, oxygen, sulfur, and phosphorus (P) is complemented by a selected array of other elements, usually metals or metalloids present in trace quantities that serve critical cellular functions, such as enzyme co-factors (1). There are many cases of these trace elements substituting for one another. A few examples include the substitution of tungsten for molybdenum and cadmium for zinc in some enzyme families (2, 3) and copper for iron as an oxygen-carrier in some arthropods and mollusks (4). In these examples and others, the trace elements that interchange share chemical similarities that facilitate the swap. However, there are no prior reports of substitutions for any of the six major elements essential for life. Here, we present evidence that arsenic can substitute for phosphorus in the biomolecules of a naturally occurring bacterium.

Arsenic (As) is a chemical analog of P, which lies directly below P on the periodic table. Arsenic possesses a similar atomic radius, as well as near identical electronegativity to P (5). The most common form of P in biology is phosphate (PO_4^{3-}) , which behaves similarly to arsenate (AsO₄³⁻) over the range of biologically relevant pH and redox gradients (6). The physicochemical similarity between AsO₄³⁻ and PO₄³⁻ contributes to the biological toxicity of AsO₄³⁻ because metabolic pathways intended for PO₄³⁻ cannot distinguish between the two molecules (7) and AsO₄³⁻ may

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be incorporated into some early steps in the pathways [(6)] and references therein]. However, it is thought that downstream metabolic processes are generally not compatible with As-incorporating molecules because of differences in the reactivities of P and As compounds (8). These down-

stream biochemical pathways may require the more chemically stable P-based metabolites; the lifetimes of more easily hydrolyzed As-bearing analogs are thought to be too short. However, given the similarities of As and P-and by analogy with trace element substitutions—we hypothesized that AsO₄³⁻ could specifically substitute for PO₄³⁻ in an organism possessing mechanisms to cope with the inherent instability of AsO₄³⁻ compounds (6). Here, we experimentally tested this hypothesis by using AsO₄³⁻, combined with no added PO_4^{3-} , to select for and isolate a microbe capable of accomplishing this substitution.

Geomicrobiology of GFAJ-1. Mono Lake, located in eastern California, is a hypersaline and alkaline water body with high dissolved arsenic concentrations [200 µM on average (9)]. We used lake sediments as inocula into an aerobic defined artificial medium at pH 9.8 (10, 11) containing 10 mM glucose, vitamins, and trace metals but no added PO₄³⁻ or any additional complex organic supplements (such as yeast extract or peptone), with a regimen of increasing AsO₄³⁻ additions initially spanning the range from 100 µM to 5 mM. These enrichments were taken through many decimal-dilution transfers, greatly reducing any potential carryover of autochthonous phosphorus

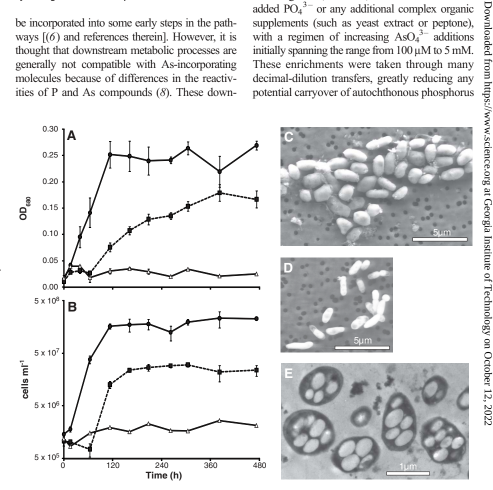


Fig. 1. Growth and electron microscopy of strain GFA]-1. (A and B) Growth curves of GFA]-1 grown on the defined synthetic medium amended with either 1.5 mM PO_4^{3-} (solid circles), 40 mM As O_4^{3-} (solid squares), or neither PO₄³⁻ nor AsO₄³⁻ (open triangles). Cell growth was monitored both by an increase in (A) optical density and (B) cell numbers of the cultures. Symbols represent the mean \pm SD of (A) n=6experimental and n=2 controls and (B) n=3 experimental and n=1 control. This was a single experiment with six replicates; however, material was conserved to extend the duration of the experiment to allow material for cell-counting samples. (C and D) Scanning electron micrographs of strain GFA]-1 under two conditions, (C) +As/-P and (D) -As/+P. (E) Transmission electron micrography of +As/-P GFA]-1 showed internal vacuole-like structures. Scale bars are as indicated in the figure (11).

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(11). The background PO_4^{3-} in the medium was $3.1 (\pm 0.3) \,\mu\text{M}$ on average, with or without added AsO₄³⁻, coming from trace impurities in the major salts (table S1) (11). The sixth transfer of the 5 mM AsO₄³⁻ (no added PO₄³⁻) condition was closely monitored and demonstrated an approximate growth rate (μ) of 0.1 day⁻¹. After 10^{-7} dilutions, we used the 5 mM AsO₄³⁻ enrichment to inoculate an agar plate that contained the same chemical composition as the artificial medium. An isolated colony was picked from the agar plates and reintroduced into an artificial liquid medium with no added PO₄³⁻, where we then progressively increased the AsO₄³⁻ concentration to determine the optimal level for growth. Currently, this isolate—strain GFAJ-1 identified by 16S ribosomal RNA sequence phylogeny as a member of the Halomonadaceae family of Gammaproteobacteria (fig. S1) (11)—is maintained aerobically with 40 mM AsO₄³⁻, 10 mM glucose, and no added PO₄³⁻ (+As/-P condition). Members of this family have been previously shown to accumulate intracellular As (12).

GFAJ-1 grew at an average μ_{max} of 0.53 day $^{-1}$ under +As/–P, increasing by over 20-fold in cell numbers after 6 days. It also grew faster and

more extensively with the addition of 1.5 mM $PO_4^{~3-}$ (–As/+P, μ_{max} of 0.86 day $^{-1}$) (Fig. 1, A and B). However, when neither AsO $_4^{~3-}$ nor $PO_4^{~3-}$ was added no growth was observed (Fig. 1, A and B). We include both optical density and direct cell counts to unambiguously demonstrate growth using two independent methods. Cells grown under +As/-P were oblong and approximately 2 by 1 µm when imaged by means of scanning electron microscopy (Fig. 1C) (11). When grown under +As/–P conditions, GFAJ-1 cells had more than 1.5-fold greater intracellular volume ($\approx 2.5 \pm 0.4 \,\mu\text{m}^3$) as compared with that of -As/+P ($\approx 1.5 \pm 0.5 \mu m^3$) (Fig. 1D) (11). Transmission electron microscopy revealed large vacuole-like regions in +As/-Pgrown cells that may account for this increase in size (Fig. 1E). These experiments demonstrated AsO₄³⁻-dependent growth, morphological differences in GFAJ-1 driven by AsO₄³⁻ in the growth medium, and that the level of PO₄³⁻ impurities in the medium was insufficient to elicit growth in the control (-As/-P).

Cellular stoichiometry and elemental distribution. To determine whether GFAJ-1 was taking up AsO₄³⁻ from the medium, we measured the

intracellular As content by means of inductively coupled plasma mass spectrometry (ICP-MS) (11). In +As/-P-grown cells, the mean intracellular As was $0.19 \pm 0.25\%$ by dry weight (Table 1), whereas the cells contained only 0.019 +/- 0.009 P by dry weight. This P was presumably scavenged from trace PO₄³⁻ impurities in the reagents and not likely due to carryover, given our enrichment and isolation strategy (11). Moreover, when grown +As/-P this intracellular P is 96.5% less than our measured P values for this microbe when grown -As/+P and far below the 1 to 3% P by dry weight required to support growth in a typical heterotrophic bacterium (13). In contrast, GFAJ-1 cells grown under -As/+P conditions had a mean P content of $0.54 \pm 0.21\%$ by dry weight. There was variation in the total As content of the +As/-P cells, which is possibly a result of collection during stationary phase and losses during the repeated centrifugations and washing cycles because of the potential instability of the cellular structures given their swollen state (Fig. 2, C and E). In contrast, the integrity of the -As/+P cells appeared robust (Fig. 2D), and thus intracellular P measured for these cells probably reflects their content. However, the low total intracellular P in +As/-P cells was consistently far below the quantity needed to support growth, suggesting that these low values are correct despite variation in data from the +As/-P cells. Low intracellular P in concert with high intracellular As was further confirmed through high-resolution secondary ion mass spectrometry and x-ray analyses, as discussed below.

We used radiolabeled ⁷³AsO₄ ³⁻ to obtain more specific information about the intracellular distribution of arsenic (11). We observed intracellular arsenic in protein, metabolite, lipid, and nucleic acid cellular fractions (Table 2). Stationary phase cells incorporated approximately 1/10 of the total intracellular ⁷³AsO₄³⁻ label into nucleic acids but more than three quarters of the 73 AsO₄³⁻ into the phenol extracted "protein" fraction, with a small fraction going into lipids. We caution that the large "protein" fraction is probably an overestimate because this extraction step likely contains numerous small, non-proteinaceous metabolites as well. To determine whether this distribution pattern reflected a use of AsO₄³⁻ in place of PO₄³⁻ in DNA, we estimated the average sequenced bacterial genome to be 3.8 megabase pairs (Mbp), which would contain approximately 7.5×10^6 atoms or 12.5×10^{-18} moles of P. Assuming one complete genome per cell, this would equal 0.39 fg of P in the genome. Using ICP-MS, we measured about 9.0 fg P per cell in the -As/+P condition, which implies that only ~4% of total intracellular P is associated with the genome. Because these cells were harvested in stationary phase (11), the fraction of P associated with RNA is likely small (14). Hence, roughly 96% of P is presumably distributed between the "lipid" and "protein" fractions. If AsO₄³⁻ is substituting for PO₄³⁻ in DNA, then we can assume that roughly the same fraction of the total intracellular

Table 1. Bulk intracellular elemental profile of strain GFA]-1. Cells were grown and prepared with trace metal clean techniques (11). Numbers in parentheses indicate replicate samples analyzed. As:P ratios were calculated based on all samples analyzed (11). Units are percent dry weight.

Condition	As	P	As:P
+As/-P (8)	$\textbf{0.19}\pm\textbf{0.25}$	0.019 ± 0.009	7.3
-As/+P (4)	0.001 ± 0.0005	$\textbf{0.54}\pm\textbf{0.21}$	0.002

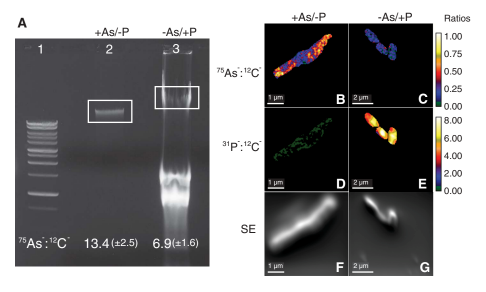


Fig. 2. NanoSIMS analyses of GFA]-1: extracted DNA and whole-cells elemental ratio maps. **(A)** Agarose gel loaded with DNA/RNA extracted from GFA]-1 grown (lane 2) +As/–P and (lane 3) –As/+P as compared with (lane 1) a DNA standard. Genomic bands were excised as indicated and analyzed with NanoSIMS. Ion ratios of ⁷⁵As^{-;12}C⁻ of excised gel bands are indicated below with 2σ error shown (all values multiplied by 10⁻⁶). **(B** to **G)** NanoSIMS images of whole GFA]-1 cells grown either [(B), (D), and (F)] +As/–P or [(C), (E), and (G)] –As/+P. Shown are the ion ratios of [(B) and (C)] ⁷⁵As^{-;12}C⁻, [(D) and (E)] ³¹P^{-;12}C⁻, and [(F) and (G)] secondary electron (SE). Ratios in (B) and (C) are multiplied by 10⁻⁴ and in (D) and (E) are multiplied by 10⁻³. The color bars indicate measured elemental ratios on a log scale as indicated. Length scale is as indicated on images; images contain equivalent pixel density (11).

 ${\rm AsO_4}^{3-}$ would reflect a similar distribution to our estimated ${\rm PO_4}^{3-}$ distribution. The distribution of intracellular $^{73}{\rm AsO_4}^{3-}$ in our experiments was consistent with these estimates. If ${\rm AsO_4}^{3-}$ is fulfilling the biological role of ${\rm PO_4}^{3-}$, then ${\rm AsO_4}^{3-}$ should act in many analogous biochemical roles, including DNA, protein phosphorylation, small molecular weight metabolites [such as arsenylated analogs of the reduced form of nicotinamide adenine dinucleotide phosphate (NADH), adenosine triphosphate (ATP), and intermediates like glucose and acetyl–coenzyme A (acetyl-CoA)], and phospholipids.

Our data suggested that As was present in a number of biomolecules, and in particular we sought to confirm the presence of As in the DNA fraction. Initially, we measured traces of As with ICP-MS analysis of extracted nucleic acid and protein/metabolite fractions from +As/-P-grown cells (table S1) (11). We then used high-resolution secondary ion mass spectrometry (NanoSIMS) to identify As in extracted, gel-isolated genomic DNA (Fig. 2A). These data showed that DNA from +As/-P cells had elevated As and low P relative to DNA from the -As/+P cells. NanoSIMS analysis of the DNA showed that the As:P ratio on an atom-per-atom basis was significantly higher in the +As/-P versus -As/+P grown cells (P =0.02) (Fig. 2A and table S2) (11). Whether expressed as an ion ratio relative to C (⁷⁵As⁻: ¹²C⁻), (Fig. 2A) or ³¹P⁻: ¹²C⁻ (table S2) (11), or normalized by relative ion yield and expressed as a concentration in parts per billion (table S2) (11), we saw a similarly consistent trend, with significantly higher As in the +As/-P DNA (P = 0.03) and higher P in the -As/+P DNA (P = 0.0003). In

Table 2. Intracellular radiolabeled 73 AsO₄ $^-$ arsenate distribution. All major cellular subfractions contained radiolabel after cell-washing procedures. Small molecular weight (s.m.w.) metabolites potentially include arsenylated analogs of ATP, NADH, acetyl-CoA, and others (11). SE is shown.

Cellular radiolabel recovered (percent of total)	
80.3 ± 1.7	
5.1 ± 4.1	
1.5 ± 0.8	
$\textbf{11.0}\pm\textbf{0.1}$	

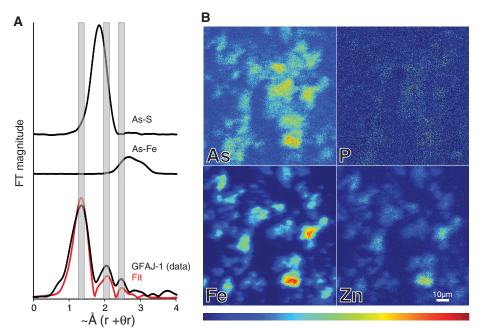


Fig. 3. X-ray analysis of GFAJ-1 +As/-P described similarity of As coordinated like P in DNA. (A) EXAFS comparisons of the Fourier transformed data for As in model environments and GFAJ-1 (washed and fixed, collected on whole cells). Identification of each spectrum is indicated on the figure and from top to bottom are As-S, As-Fe, and GFAJ-1 data (collected on whole cells) and fit to the GFAJ-1 data (red line). (B) XRF maps indicated the correlation between As, iron (Fe) and zinc (Zn) and not with P with some variability but consistent with the trend that these elements are often found together (fig. S3, element correlation plots). The scale bar in the Zn quadrant of the maps is as designated and applies to all parts of the figure. Given the spatial resolution of these images, the structures identified as containing high As, Fe, and Zn are aggregates of cells. Ranges as indicated in the color bar run from cold to hot, in units of μ g cm⁻², as follows: As, 0 to 1.6; P, 0 to 40; Fe, 0 to 32.1; and Zn, 0 to 2.8. Standards were used to calibrate signal and background (11).

both cases, the nonamended element concentration was equal or less than background levels. These measurements therefore specifically demonstrated that the purified DNA extracted from +As/–P cells contained As. Our NanoSIMS analyses, combined with the evidence for intracellular arsenic by ICP-MS and our radiolabeled ⁷³AsO₄³⁻ experiments, indicated that intracellular AsO₄³⁻ was incorporated into key biomolecules, specifically DNA.

Characterization of the intracellular arsenic chemical environment. We next used synchrotron x-ray studies to determine the speciation and chemical environment of the intracellular arsenic (11). Micro-x-ray absorption near-edge spectroscopy (µXANES) of +As/–P–grown cells exhibited an absorption edge characteristic of As(V) coordination with no evidence of As(III) observed. Best fits of the micro-extended x-ray absorption fine structure (uEXAFS) spectra are listed in Table 3 and shown in Fig. 3. The first neighbor shell around the arsenic in +As/–P cells consisted of four oxygen ligands (Table 3) but had a second shell that is inconsistent with our As-Fe and As-S models, free AsO₄³⁻ ions, or published spectra for organo-arsenicals (Fig. 3A) (15, 16). Although other arsenical compounds such as dimethylarsinate (DMA) also have As-O and As-C bonds, they have edge positions that are shifted to lower energy from the observed As(V) and have much shorter observed As-C bond distances (16). In contrast to the models, these As-O and As-C distances are consistent with that reported from the solved crystal structure of DNA for the analogous structural position of P relative to O and C atoms (Fig. 3A) (16, 17). Therefore, our x-ray data support the position of AsO₄³⁻ in a similar configuration to PO₄³⁻ in a DNA backbone or potentially other biomolecules as well. These data are also consistent with the presence of AsO₄³⁻ in small-molecular-weight metabolites (for example, arsenylated analogs of NADH, ATP, glucose, and acetyl-CoA) as well as arsenylated proteins in which AsO₄³⁻ would substitute for PO₄³⁻ at serine, tyrosine, and threonine residues (table S3) (1, 11). Micro-x-ray fluorescence data (uXRF) further confirmed our ICP-MS measurements and showed low background P, which contrasted with regions of high arsenic correlated with high iron and zinc (Fig. 3B and fig. S2) (11). These latter two elements are routinely used as proxies for the presence of cellular material (such as C, N, and O) in our experiments because these light elements could not be detected with x-ray fluorescence under our nonvacuum conditions. However, to further support the distribution of arsenic with cellular material we used NanoSIMS to map cellular ion ratios of ⁷⁵As⁻: ¹²C⁻ and ³¹P⁻: ¹²C⁻ (Fig. 2, B to G, and fig. S2) (11). These analyses confirmed, at a much finer resolution, that for the +As/-P condition the cells contain As with a low background of P (Fig. 2, B, D, and F). This is in contrast to the low As and higher P in -As/+P-grown cells (Fig. 2, C, E, and G). Because the x-ray absorption data

Table 3. Results of fitting arsenic K-edge EXAFS of GFA]-1. S02 = 1, global amplitude factor and E0 = 13.97, offset for calibration. Type, the coordination type; Number, the coordination number; R, interatomic distance; and σ^2 , the measure of the static disorder of the shell. See table S2 for comparison to P in P-containing biomolecules (11).

Туре	Number	R	σ^2
As-O	4.2 (0.6)	1.73 (2)	0.003 (2)
As-C	2.5 (0.5)	2.35 (4)	0.003 (2)
As-C	2.2 (0.5)	2.92 (6)	0.003 (2)

provided information about the average coordination of arsenic, our data identified a mixture of compounds in the cells. These results indicated that these compounds are dominated by arsenic(V)-oxygen-carbon-coordinated structures, and thus the bonding environment we described is consistent with our NanoSIMS data (Fig. 2A) and can be attributed to DNA. These data show that As is in the +5 redox state and bound to O and distal C atoms within acceptable covalent bond lengths, identifying AsO₄³⁻ assimilated into biomolecules within the cells in specifically relevant coordination.

Our data show arsenic-dependent growth by GFAJ-1 (Fig. 1). Growth was accompanied by AsO₄³ uptake and assimilation into biomolecules including nucleic acids, proteins, and metabolites (Figs. 2 and 3 and Tables 1 and 2). In some organisms, arsenic induces specific resistance genes to cope with its toxicity (7), whereas some dissimilatory arsenic-utilizing microbes can conserve energy for growth from the oxidation of reduced arsenic species, or "breathe" AsO₄³⁻, as a terminal electron acceptor (18). Our study differs because we used As as a selective agent and excluded P, a major requirement in all hitherto known organisms. However, GFAJ-1 is not an obligate arsenophile and grew considerably better when provided with P (Fig. 1, A and B). Although AsO₄³⁻ esters are predicted to be orders of magnitude less stable than PO₄³⁻ esters, at least for simple molecules (8), GFAJ-1 can cope with this instability. The vacuole-like regions observed in GFAJ-1 cells when growing under +As/-P conditions are potentially poly-β-hydroxybutyrate rich [as shown in other *Halomonas* species (19)], which may

stabilize As(V)-O-C type structures because nonaqueous environments appear to promote slower hydrolysis rates for related compounds (8). We propose that intracellular regions or mechanisms that exclude water may also promote this stability.

We report the discovery of an unusual microbe, strain GFAJ-1, that exceptionally can vary the elemental composition of its basic biomolecules by substituting As for P. How As insinuates itself into the structure of biomolecules is unclear, and the mechanisms by which such molecules operate are unknown.

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Supporting Online Material

www.sciencemag.org/cgi/content/full/science.1197258/DC1 Materials and Methods

Figs. S1 to S3 Tables S1 to S4 References

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