

An arsenic metallochaperone for an arsenic detoxification pump

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Environmental arsenic is a world-wide health issue, making it imperative for us to understand mechanisms of metalloid uptake and detoxification. The predominant intracellular form is the highly nephritic arsenite, which is detoxified by removal from cytosol. What prevents arsenite toxicity as it diffuses through cytosol to efflux systems? Although intracellular copper is regulated by metallochaperones, no chaperones involved in conferring resistance to other metals have been identified. In this article, we report identification of an arsenic chaperone, ArsD, encoded by the *arsRDABC* operon of *Escherichia coli*. ArsD transfers trivalent metalloids to ArsA, the catalytic subunit of an As(III)/Sb(III) efflux pump. Interaction with ArsD increases the affinity of ArsA for arsenite, thus increasing its ATPase activity at lower concentrations of arsenite and enhancing the rate of arsenite extrusion. Cells are consequently resistant to environmental concentrations of arsenic. This report of an arsenic chaperone suggests that cells regulate the intracellular concentration of arsenite to prevent toxicity.

antimony | resistance pump | metalloid | ArsD | ArsA

All cells possess regulatory mechanisms that control the intracellular concentration of metals, including essential metals, which become toxic in excess (copper and zinc), toxic metals (cadmium and lead), and toxic metalloids (arsenite and antimonite) (1–3). For example, metallothioneins are small proteins found in both eukaryotes and prokaryotes that bind transition metal ions such as copper and zinc for storage and heavy metals such as cadmium for detoxification (4). In plants and some fungi phytochelatins are cysteine-rich peptides that bind heavy metal ions for detoxification (5). In contrast to these types of metal-binding peptides and polypeptides are metallochaperones, which sequester metals in the cytoplasm and deliver them to protein targets (1, 6, 7). In some cases the targets are metalloenzymes such as superoxide dismutase, where the *Saccharomyces cerevisiae* metallochaperone Lys-7 (CCS in humans) is required for insertion of copper ions (8). Although the best characterized metallochaperones are for copper, UreE is a nickel chaperone that delivers Ni²⁺ to urease (9), and frataxin is a chaperone that delivers iron to proteins for assembly of iron–sulfur clusters and heme (10). Other targets are membrane transporters, predominantly transport ATPases, that participate in metal ion homeostasis and detoxification such as *S. cerevisiae* Atx1 (Hah1 in humans), the metallochaperone for the yeast Ccc2p (ATP7A and ATP7B in humans) Cu(I)-translocating P-type ATPase (11). To date, no metallochaperones involved in detoxification of metals other than copper have been identified.

Arsenic is the most prevalent environmental toxic metal and is first on the Superfund list of hazardous substances (www.atsdr.cdc.gov/cercla/05list.html). Health effects associated with exposure to low levels of arsenite include cardiovascular and peripheral vascular disease, neurological disorders, diabetes mellitus, and various forms of cancer (12, 13). Chronic exposure to the low levels of arsenic that exist naturally in both water and soil (14) provides competitive pressure to maintain arsenic tolerance or detoxification systems in most, if not all, living organisms, including humans (15). To date, nearly every pro-

karyote with a sequenced genome has an *ars* operon, as do a number of archaea, and some budding and filamentous fungi have clusters of arsenic detoxification genes. *ars* operons are diverse but most often contain three genes for an arsenic-responsive transcriptional repressor (*arsR*), an arsenite efflux membrane protein (*arsB*), and an arsenate reductase (*arsC*) (16). Some have two additional genes, *arsD* and *arsA*, first identified in the *Escherichia coli* plasmid R773 *ars* operon (17, 18). The 26-kDa homodimeric ArsD (120 aa residues per monomer) is a cytosolic protein with three vicinal cysteine pairs per monomer (Cys-12-Cys-13, Cys-112-Cys-113, and Cys-119-Cys-120) (see Fig. 5, which is published as supporting information on the PNAS web site). ArsD has weak As(III)-responsive transcriptional regulatory activity (18). ArsA is an ATPase that associates with ArsB to form an As(III)-translocating ATPase that extrudes arsenite from cells, conferring an additional level of resistance over cells with only ArsB (19). It is striking that the *arsD* and *arsA* genes always are found together in bacterial and archaeal arsenic resistance operons or gene clusters (see Table 1, which is published as supporting information on the PNAS web site), which led us to propose that five-gene *arsRDABC* operons evolved from three-gene *arsRBC* operons by insertion of the *arsD* and *arsA* genes as a unit (16). The linkage of ArsD and ArsA further suggests that they have associated functions in arsenic detoxification. In this report, we demonstrate that the ArsD serves as a chaperone for intracellular arsenite, transferring metalloid to the pump.

Results

ArsD Confers a Competitive Advantage to Cells Growing in Subtoxic Concentrations of Arsenite. When arsenite resistance was compared between cells expressing the *arsAB* genes or the *arsDAB* genes, the latter showed a modest increase in resistance, with the greatest differences observed at toxic (millimolar) concentrations of arsenite (see Fig. 6, which is published as supporting information on the PNAS web site). Does ArsD confer a competitive advantage for growth in subtoxic concentrations of arsenite? A molecular competition experiment was devised to examine this question. We used cells of *E. coli* strain AW3110, in which the chromosomal *ars* operon had been deleted, bearing either a plasmid with either *arsAB* or *arsDAB* in the same vector. Note that the *E. coli* chromosome has neither *arsD* nor *arsA* genes. Equal amounts of the two types of cells were mixed

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Abbreviations: bBBR, dibromobimane; MBP, maltose-binding protein; LB, Luria-Bertani.

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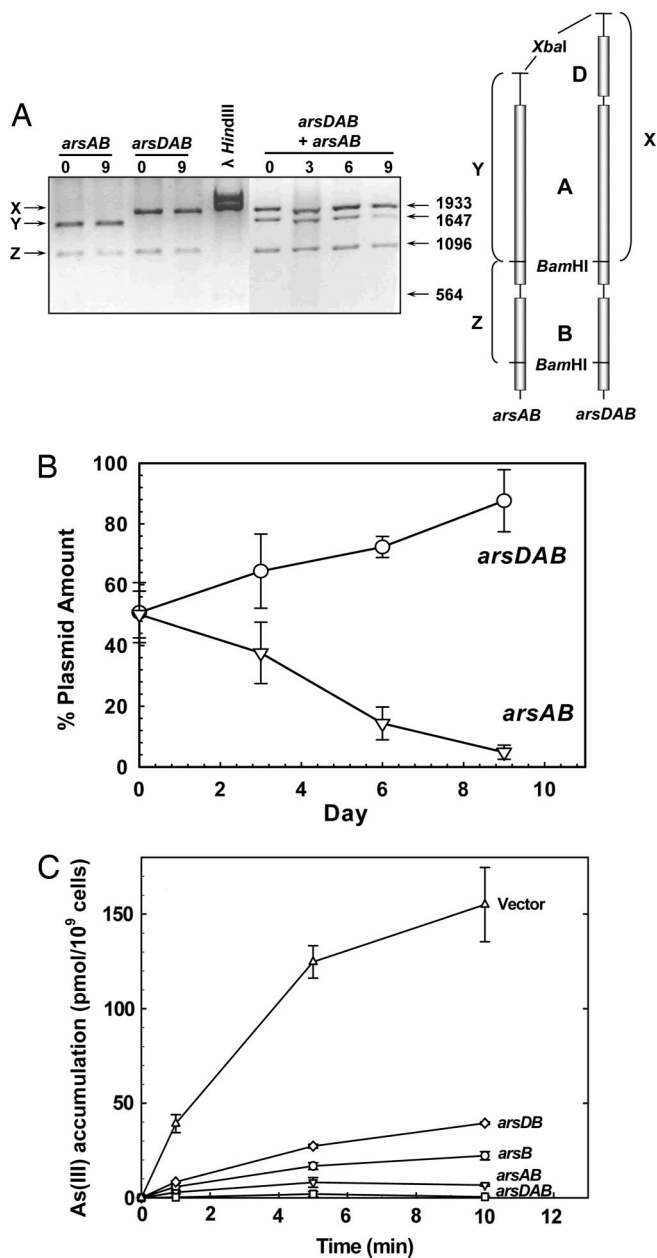


Fig. 1. ArsD enhances the activity of the ArsAB pump *in vivo*. (A) Molecular competition between cells with *arsDAB* and *arsAB*. Cells of *E. coli* strain AW3110 bearing either pSE-AB or pSE-DAB were grown separately or in mixed culture of each for 9 days with daily dilutions into fresh LB medium containing 10 μ M sodium arsenite, as described in *Materials and Methods*. The plasmids from equal amounts of cells were extracted and digested with restriction enzymes XbaI and BamHI, and equal amounts of DNA containing the restriction fragments were separated by electrophoresis on a 1% agarose gel. Both plasmids produced a large fragment (data not shown) of vector DNA, and both produced a small *arsB*-containing BamHI fragment of 1.1 kb (fragment Z). Digestion of pSE-AB also produced a 1.6-kb fragment containing most of *arsA* (fragment Y), and digestion of pSE-DAB yielded the 1.9-kb fragment X containing *arsD* and most of *arsA*. The restriction fragments were from cells with only pSE-AB before growth (lane 1) or after 9 days (lane 2), only pSE-DAB before growth (lane 3) or after 9 days (lane 4), or from a mixed culture of cells with either pSE-AB or pSE-DAB before growth (lane 6) or after 3 days (lane 7), 6 days (lane 8), or 9 days (lane 9). Lane 5 contains a standard of λ HindIII-digested DNA. (B) Cells with only *arsAB* are lost from the population. The fraction of each plasmid was calculated by quantifying bands X, Y, and Z by densitometry. The percentage of the cells with each plasmid was calculated as follows: *arsDAB*: $X/((\text{vector} + Z)/2)$; *arsAB*: $Y/((\text{vector} + Z)/2)$. The data are the mean of values from five separate gels representing three independent

together and grown in the presence of subtoxic arsenite (10 μ M), which is in the range of what is found in the environment (14). Although expression was under the control of the *tac* promoter in this experiment, when under the control of the *ars* promoter, the *ars* operon is induced fully by 1 μ M arsenate or arsenite (20). Each day the mixed culture was diluted 1,000-fold, and the relative amounts of the *arsDAB* and *arsAB* plasmids were determined by quantitative densitometry of the bands produced by restriction enzyme digestion with XbaI and BamHI (Fig. 1A). Digestion of both plasmids produced a 1.1-kbp BamHI fragment. The size of the XbaI–BamHI fragment in the *arsAB* plasmid was 1.6 kbp. The presence of *arsD* in *arsDAB* plasmid increased the size of this fragment to 1.9 kbp. After 9 days, cells with *arsDAB* had largely replaced those with only *arsAB* (Fig. 1B), indicating that ArsD provides a competitive advantage for growth in the low concentrations of arsenite that are ubiquitous in soil or surface waters.

The effect of the *ars* genes on arsenite accumulation was examined. Higher rates of extrusion result in lower accumulation (19). Cells of the arsenite-hypersensitive strain AW3110 with no *ars* genes (vector plasmids pSE380 and pACBAD) accumulated ≈ 150 pmol As(III) per 10^9 cells per 10 min (Fig. 1C). Cells expressing only *arsB* accumulated ≈ 22 pmol As(III) per 10^9 cells per 10 min. Expression of *arsAB* resulted in decreased accumulation to ≈ 7 pmol As(III) per 10^9 cells per 10 min, reflecting more efficient arsenite extrusion by the ArsAB pump than by ArsB alone (19). Cells coexpressing *arsD* and *arsAB* accumulated substantially less arsenite (≈ 1 pmol As(III) per 10^9 cells per 10 min) than did those with *arsAB*. Cells expressing *arsD* with only *arsB* accumulated arsenite to approximately the same level as cells expressing only *arsB*, indicating that ArsD does not affect expression or activity of ArsB. Immunoblotting showed that *arsD* does not affect the levels of ArsA produced (see Fig. 7, which is published as supporting information on the PNAS web site). These results clearly show that ArsD enhances the ability of the ArsAB pump to reduce intracellular arsenite.

Interaction of ArsD with ArsA. Yeast two-hybrid analyses were used to show that ArsD and ArsA physically interact (Fig. 2A). ArsA interacted with ArsD but not with ArsR or ArsC. ArsD interacted with ArsA and with itself, as expected because ArsD is a homodimer (21), but not with ArsR or ArsC. Homodimeric ArsR also interacts with itself but not with ArsD or ArsA. These results indicate specific interaction of ArsD and ArsA.

Because both ArsD and ArsA have metalloid binding sites composed of cysteine residues (19), direct physical interaction at those sites was examined by cross-linking with dibromobimane (bBBr), a thiol reagent that becomes fluorescent when it cross-links cysteine residues within 3–6 Å of each other (22). When ArsD was treated with bBBr, dimers and higher-order species were observed by SDS/PAGE (Fig. 2B Middle). All ArsD bands became fluorescent, showing formation of both intra- and intermolecular cross-links (data not shown). When equimolar ArsD and ArsA were reacted with bBBr, a cross-linked species was detected that reacted with both anti-ArsA and anti-ArsD antibodies (indicated by an asterisk in Fig. 2B Top and Middle, lane 2). This species migrated as a band with an apparent mass of ≈ 90 kDa. Although we cannot definitively rule out other combinations of ArsD and ArsA, the mass of this band corresponds to the predicted mass of an ArsD dimer cross-linked to an ArsA monomer. Note that the intensity of the bands from one

experiments. The error bars represent the standard deviation of the mean calculated by using SigmaPlot 9.0. (C) Cells expressing *arsDAB* maintain lower intracellular arsenite than do cells expressing *arsAB*. Transport of As(III) was assayed with AW3110 bearing vector plasmids pSE380 and pACBAD (Δ) or plasmids with *arsB* (\circ), *arsAB* (∇), *arsD* and *arsAB* (\square), or *arsD* and *arsB* (\diamond).

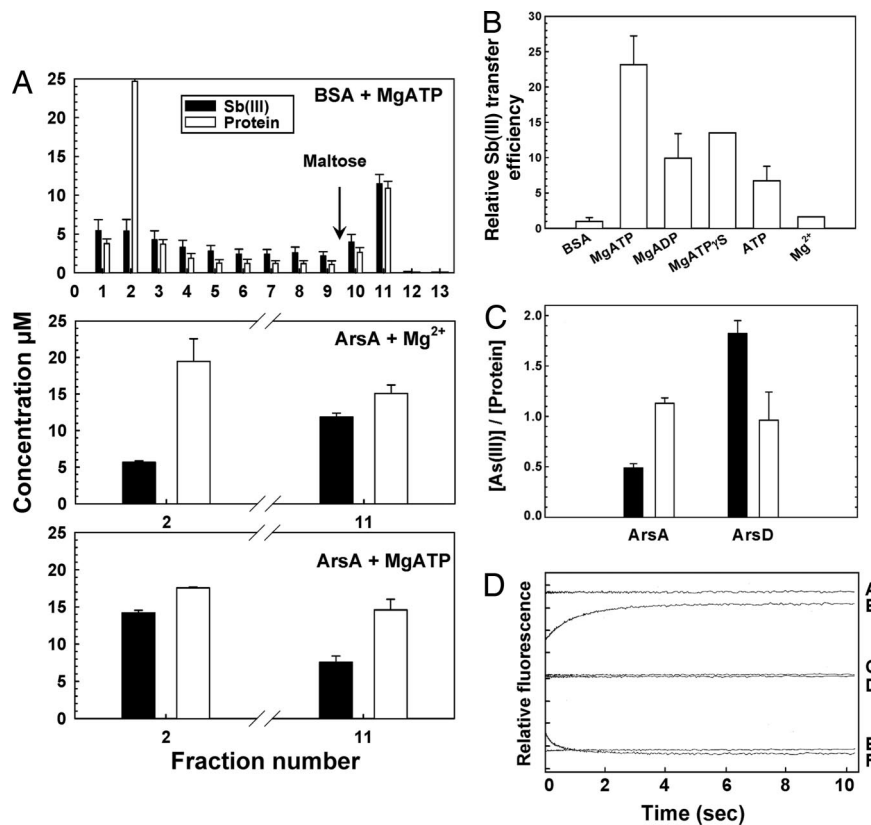


Fig. 3. Arsd transfers metalloid to and activates the ArsA ATPase. (A) Arsd releases Sb(III) from Arsd. Sb(III)–MBP–Arsd was bound to a 2-ml amylose column. One milliliter of either 20 μM Arsa or BSA preincubated with 1 mM MgCl₂ or MgATP was applied to the column. The column was washed with 8 ml of column buffer, and MBP–Arsd was eluted with 4 ml of 10 mM maltose. The protein in each fraction was identified by SDS/PAGE. BSA or Arsa eluted primarily in fraction 2, and most of the MBP–Arsd eluted in fraction 11. The molar concentration of each protein in the fractions was estimated from the absorption at 280 nm (white bars), and amount of Sb(III) was determined by inductively coupled plasma mass spectroscopy (black bars). (B) A nucleotide enhances the Arsa-induced release of Sb(III) from Arsd. Sb(III) release from Arsd was assayed in the presence of the indicated nucleotides and is expressed relative to the values with BSA. The values are the average of two independent assays. (C) Arsd transfers As(III) to Arsa. The molar ratio of As(III) to either Arsa or Arsd monomer was measured with protein either alone (black bars) or in the presence of the partner protein (white bars). The values are the mean of three independent assays. (D) The transfer of Sb(III) from Arsd to Arsa time-resolved by stopped-flow fluorescence spectroscopy. Equal volumes of the following reagents were mixed in a stopped-flow device, and the changes in protein fluorescence (excitation = 285 nm; emission >340 nm) monitored. Curve A, 2 μM Arsd + 10 μM Arsa; curve B, 2 μM Arsd/10 μM Sb(III) + 10 μM Arsa; curve C, 10 μM Arsa + buffer; curve D, 10 μM Arsa + 10 μM Sb(III); curve E, 2 μM Arsd + 10 μM Sb(III); curve F, 2 μM Arsd/10 μM Sb(III) + buffer. Each division on the y axis represents a fluorescence change of 5% relative to that of 2 μM Arsd/10 μM Arsa.

conferred by efflux pumps, including transport ATPases and resistance–nodulation–cell division (RND) transporters (26, 27). However, to date, the only other metallochaperones identified appear to be involved in delivery to metal-containing proteins. For example, a nickel chaperone delivers Ni²⁺ to the enzyme urease (9), and the chaperone frataxin delivers iron to partners for assembly of iron–sulfur clusters and heme (10). In this article, we report the existence of a chaperone for the toxic metalloid arsenic that enhances resistance by delivering the metalloid to the ArsAB efflux pump. Although there are superficial similarities with copper chaperones, Arsd is unrelated to any other chaperones in sequence or mechanism. Our yeast two-hybrid data suggest that Arsd interacts with Arsa in the absence of metalloid. We propose that Arsd interacts with Arsa with low affinity in the absence of metalloid and high affinity when metalloid is bound. Arsd has been shown to bind metalloid with higher affinity than Arsa does. For transfer from a high-affinity binding site on one protein to a relatively lower-affinity site on another protein, interaction of the two proteins must produce conformational changes in Arsd and/or Arsa. Indeed, binding of Sb(III) to Arsd is so tight that dissociation takes hours in the absence of Arsa, but the release rate is stimulated by more than four orders of magnitude by Arsa (Fig.

3D). We further propose that cysteine residues on Arsd are involved in the interaction with Arsa and that they are brought into close proximity to the metalloid binding sites of Arsa, allowing transfer of metalloid directly from the binding site on Arsd to the binding site on Arsa. It is plausible that this interaction destabilizes the metalloid binding sites on Arsd, reducing its affinity, while stabilizing binding to Arsa by occluding metalloid within the complex, thus facilitating transfer to Arsa. In this manner, the thermodynamically unfavorable process of transferring metalloid from a high- to a low-affinity site is overcome.

Our stopped-flow studies of the transfer of Sb(III) from Arsd to Arsa clearly are consistent with a mechanism in which this metallochaperone accelerates the rate of transfer to its partner. Because Arsd has higher affinity for metalloid than Arsa, it can scavenge the cytosol for free metalloid for delivery to Arsa, allowing the ArsAB pump to confer resistance at significantly lower concentrations of As(III). The *K_m* of Arsb as a secondary carrier is 140 μM, but natural waters range in concentration of total inorganic arsenic from 7 nM to 70 μM, and concentrations of arsenic in drinking water in the worst arsenic-contaminated wells in West Bengal and Bangladesh are ≈40 μM (14). The *ars* operon is induced fully by 1 μM arsenate or arsenite (20), so

buffer containing 62.5 mM Tris·HCl (pH 6.8), 0.1 M 2-mercaptoethanol, and 2% SDS at 50°C for 30 min.

Metalloid Transfer Assays. Cells of *E. coli* BL21(DE) expressing MBP–ArsD were lysed in the presence of 1 mM potassium antimonyl tartrate in buffer A lacking DTT. The cytosol containing the MBP–ArsD–Sb(III) complex was applied to a 2-ml amylose column, which was washed with 20 ml of the same buffer at room temperature. BSA or purified ArsA (1 ml of 20 μM) was applied to the columns with 1 mM ATP, ADP, ATPγS, and/or MgCl₂, as indicated. The column was washed with 8 ml of buffer, and MBP–ArsD was eluted with 4 ml of 10 mM maltose. As shown from SDS/PAGE, fraction 2 contained nearly all of the BSA or ArsA with little MBP–ArsD, and fraction 11 contained nearly all of the MBP–ArsD with little or no BSA or ArsA. Antimony was quantified by inductively coupled plasma mass spectroscopy. Transfer activity was calculated as ([Sb(III)_{ArsA}]/[ArsA])/([Sb(III)_{ArsD}]/[ArsD]).

To further demonstrate that ArsA accepts As(III) from ArsD, the amount of arsenic on each protein was determined after

interaction. To determine the amount of As(III) bound to ArsA, a mixture of 3 μM his-tagged ArsA, 25 μM sodium arsenite, 2.5 mM MgCl₂, and 2 mM ATPγS with or without 9 μM MBP–ArsD was incubated at 37°C for 10 min. To isolate the As(III)–ArsA complex, MBP–ArsD and free As(III) were removed with a microspin gel-filtration column with a 0.3-ml layer of amylose resin applied at the top of the spin column. ArsA and bound arsenic concentrations were quantified. The amount of As(III) bound to ArsD was determined similarly by adding excess ArsA (9 μM) to ArsD (3 μM). His-tagged ArsA and free As(III) were removed by using a gel-filtration spin column with a 0.3-ml layer of Ni-nitrilotriacetic acid resin applied at the top of the spin column, and the amounts of ArsD and As(III) were determined as described above. The details of fluorescence measurements and ATPase assays with these proteins are described in *Supporting Materials and Methods*.

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