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Toxicity, biotransformation, and mode of action of arsenic in two freshwater microalgae (Chlorella sp. and Monoraphidium arcuatum)

Jacqueline L. Levy jl53@uow.edu.au

Jennifer L. Stauber CSIRO, jenny.stauber@csiro.au

Merrin S. Adams msa344@uow.edu.au

William A. Maher University of Canberra

Jason K. Kirby

See next page for additional authors

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Toxicity, biotransformation, and mode of action of arsenic in two freshwater microalgae (Chlorella sp. and Monoraphidium arcuatum)

Abstract

The toxicity of As(V) and As(III) to two axenic tropical 15 freshwater microalgae. Chlorella sp. and Monoraphidium arcuatum, was determined using 72-h growth rate inhibition bioassays. Both organisms were tolerant to As(III) (72-h IC50, concentration to cause 50% inhibition of growth, of 25 and 15 mg As(III)/L, respectively). Chlorella sp. was also tolerant to As(V) with no effect on growth rate over 72 h at concentrations up to 0.8 mg/L (72-h IC50 of 25 mg As(V)/L). M. arcuatum was more sensitive to As(V) (72-h IC50 of 0.25 mg As(V)/L). An increase in phosphate in the growth medium (0.15 to 1.5 mg PO₄ $^{3-}$ /L) decreased toxicity, i.e. the 72-h IC50 value for M. arcuatum increased from 0.25 mg As(V)/L to 4.5 mg As(V)/L, while extracellular As and intracellular As decreased, indicating competition between arsenate and phosphate for cellular uptake. Both microalgae reduced As(V) to As(III) in the cell, with further biological transformation to methylated species (monomethyl arsonic acid and dimethyl arsinic acid) and phosphate arsenoriboside. Less than 0.01% of added As(V) was incorporated into algal cells, suggesting that bioaccumulation and subsequent methylation was not the primary mechanism of detoxification. When exposed to As(V) both species reduced As(V) to As(III), however only M. arcuatum excreted As(III) into solution. Intracellular arsenic reduction may be coupled to thiol oxidation in both species. Arsenic toxicity was most likely due to arsenite accumulation in the cell, when the ability to excrete and/or methylate arsenite was overwhelmed at high arsenic concentrations. Arsenite may bind to intracellular thiols, such as glutathione, potentially disrupting the ratio of reduced to oxidised glutathione and consequently inhibiting cell division.

Keywords

Toxicity, biotransformation, mode, action, arsenic, two, freshwater, microalgae, Chlorella, Monoraphidium, arcuatum, CMMB

Disciplines

Life Sciences | Physical Sciences and Mathematics | Social and Behavioral Sciences

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Authors

Jacqueline L. Levy, Jennifer L. Stauber, Merrin S. Adams, William A. Maher, Jason K. Kirby, and Dianne F. Jolley

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3	AND MONORAPHIDIUM ARCUATUM)						
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5	JACQUELINE L. LEVY,†‡ JENNIFER L. STAUBER,† MERRIN S. ADAMS,† WILLIAM A.						
6	Maher,§ Jason K. Kirby,§ Dianne F. Jolley‡*						
7	† Centre for Environmental Contaminants Research, CSIRO Energy Technology, Private						
8	Mail Bag 7, Bangor, NSW 2234, Australia						
9	‡ GEOQUEST, Department of Chemistry, University of Wollongong, NSW 2522, Australia						
10	§ Ecochemistry Laboratory, Applied Ecology Research Group, University of Canberra, ACT						
11	2601, Australia						
12							
13							
14	* To whom correspondence may be addressed Jacqueline Levy, via (djolley@uow.edu.au)						

15 Abstract- The toxicity of As(V) and As(III) to two axenic tropical freshwater microalgae, 16 Chlorella sp. and Monoraphidium arcuatum, was determined using 72-h growth rate 17 inhibition bioassays. Both organisms were tolerant to As(III) (72-h IC50, concentration to 18 cause 50% inhibition of growth, of 25 and 15 mg As(III)/L, respectively). Chlorella sp. was 19 also tolerant to As(V) with no effect on growth rate over 72 h at concentrations up to 0.8 mg/L (72-h IC50 of 25 mg As(V)/L). M. arcuatum was more sensitive to As(V) (72-h IC50 20 21 of 0.25 mg As(V)/L). An increase in phosphate in the growth medium (0.15 to 1.5 mg PO_4^{3-1} 22 /L) decreased toxicity, i.e. the 72-h IC50 value for *M. arcuatum* increased from 0.25 mg 23 As(V)/L to 4.5 mg As(V)/L, while extracellular As and intracellular As decreased, indicating 24 competition between arsenate and phosphate for cellular uptake. Both microalgae reduced 25 As(V) to As(III) in the cell, with further biological transformation to methylated species 26 (monomethyl arsonic acid and dimethyl arsinic acid) and phosphate arsenoriboside. Less than 27 0.01% of added As(V) was incorporated into algal cells, suggesting that bioaccumulation and 28 subsequent methylation was not the primary mechanism of detoxification. When exposed to 29 As(V) both species reduced As(V) to As(III), however only *M. arcuatum* excreted As(III) into 30 solution. Intracellular arsenic reduction may be coupled to thiol oxidation in both species. Arsenic toxicity was most likely due to arsenite accumulation in the cell, when the ability to 31 32 excrete and/or methylate arsenite was overwhelmed at high arsenic concentrations. Arsenite 33 may bind to intracellular thiols, such as glutathione, potentially disrupting the ratio of reduced 34 to oxidised glutathione and consequently inhibiting cell division.

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36 Keywords- Arsenic Algae Toxicity Biotransformation Phosphate

37

INTRODUCTION

38 Arsenic is a widespread contaminant in the environment. Anthropogenic sources, 39 together with natural sources, have led to extensive leaching of arsenic into surface, ground 40 and drinking waters [1]. Arsenic concentrations in freshwaters range from <1 to $10 \mu g/L$ [2], 41 with up to 5000 $\mu g/L$ reported in contaminated groundwaters [3].

42 Most studies investigating arsenic biotransformation have focussed on marine 43 environments [4], due to the formation of arsenoribosides and arsenobetaine in marine 44 invertebrates and macroalgae [5]. Arsenoribosides, believed to be the precursors of 45 arsenobetaine in marine invertebrates, have long been found in marine macroalgae [6], 46 however, their presence in freshwater microalgae has only recently been elucidated [7]. 47 Arsenic biotransformation and cycling in freshwater systems has thus far received 48 little attention, and little is known about the role of freshwater algae. Algae are an important 49 component of freshwater aquatic environments and could potentially remediate arsenic-50 contaminated waters in wetlands through adsorption and biotransformation of inorganic 51 arsenic. Microalgae in particular, have been shown to accumulate arsenic(V), with 52 bioconcentration factors ranging from 200 – 4000 [8,9]. However, more information about 53 the responses of freshwater algae to arsenic is required if they are to be used in remediation. 54 Literature data on the toxicity of arsenic to freshwater microalgae are limited to 55 chlorophytes and cyanophytes. Reported IC50 values (concentration to cause a 50%56 inhibition of growth) range over five orders of magnitude, from 0.048 to 202 mg/L 57 [10,11,12], and are generally well above environmental concentrations of arsenic. The wide variability in sensitivity to arsenic is likely due to biotic factors such as species type, differing 58 59 uptake/exclusion pathways, detoxification mechanisms and prior-exposure, as well as abiotic 60 factors such as arsenic species, phosphate concentrations, pH and exposure time.

3

61 While arsenic is toxic to microalgae at high concentrations, particularly at low 62 ambient phosphate concentrations, few studies have examined the mode of toxic action of 63 arsenic in freshwater microalgae. Most of our information on the mode of toxic action of 64 arsenic comes from studies with terrestrial plants or microorganisms such as bacteria and 65 yeasts [13,14]. A recent review of arsenic toxicity in terrestrial plants [13] showed that 66 arsenic toxicity to biota may be due to: (i) interference in phosphate metabolism, leading to 67 phosphate depletion or inhibition of adenosine triphosphate (ATP); (ii) oxidative stress due to 68 the generation of reactive oxygen species; and/or (iii) binding of arsenite to intracellular thiols 69 (sulfhydryl groups) of enzymes and tissue proteins, such as glutathione. 70 Aquatic and terrestrial biota have developed several strategies to detoxify metals and 71 metalloids such as arsenic. These include: (i) exclusion of arsenic from the cell [15]; (ii) 72 reduction of arsenate to arsenite followed by either excretion, or complexation with 73 glutathione and sequestration into vacuoles (e.g Saccharomyces cerevisiae, [14]); (iii) 74 production of other metal-binding proteins such as phytochelatins [16]; (iv) methylation to 75 less toxic organic forms, together with excretion [17]. Studies with microalgae have largely 76 focused on methylation as a potential detoxification process [17,18]. 77 The objective of this study was to investigate the toxicity, biotransformation and mode 78 of toxic action of arsenic in two axenic tropical freshwater microalgae, one arsenate-tolerant 79 species (*Chlorella* sp.) and one sensitive species (*Monoraphidium arcuatum*). The mode of 80 toxicity of arsenate and the detoxification processes in these two algae were compared. 81 Biotransformation of arsenic and arsenic speciation in cells was determined by microwave-82 assisted extraction and high performance liquid chromatography- inductively coupled plasma-83 mass spectrometry (HPLC-ICP-MS), enabling low detection limits for the quantitation of 84 arsenoribosides.

85

86

METHODS

87 Algal stock cultures

88 Chlorella sp. 12 and Monoraphidium arcuatum (Korš.) Hindák, originally isolated 89 from Lake Aesake, Papua New Guinea, were maintained axenically at CSIRO Energy 90 Technology, Sydney. Cultures were checked microscopically monthly and plated on agar 91 (2% bacto agent, 0.1% pepsin and 0.1% yeast extract) several times over 12 months to ensure 92 the absence of bacteria and other microorganisms. The algae were cultured in 1/5 strength 93 Jaworki's medium [19] and incubated at $27 \pm 1^{\circ}$ C on a 12:12 h light/dark cycle (Philips TL 94 40W cool white fluorescent lighting, 75 µmol photons/m²/s, Caringbah, NSW, Australia). 95 96 Growth rate inhibition bioassays 97 The toxicity of As(V) and As(III) to *Chlorella* sp. and *M. arcuatum* was determined 98 using 72-h growth rate inhibition bioassays. The test medium used in the bioassays was 99 synthetic softwater (96 mg/L NaHCO₃, 60 mg/L CaSO₄.2H₂O, 123 mg/L MgSO₄ and 4 mg/L 100 KCl) (Ajax and Asia Pacific Specialty Chemicals, Bacto Laboratories, Liverpool, NSW, 101 Australia) with a hardness of 80-90 mg CaCO₃/L, an alkalinity of 54 mg CaCO₃/L and a pH 102 of 7.6 \pm 0.1. The medium was vacuum filtered through an acid-washed 0.45 μ m cellulose 103 acetate/nitrate membrane filter (Millipore, Bedford, MA, USA) and stored at 4°C. 104 A batch method was used to conduct growth rate inhibition tests using 250-mL 105 borosilicate glass Erlenmeyer flasks, coated with Coatasil silanising solution (Ajax 106 Chemicals, Auburn, NSW, Australia) to prevent adsorption of arsenic to the glass. Test flasks 107 and sample storage vessels were soaked in 10% (v/v) nitric acid (BDH) overnight and rinsed 108 thoroughly with high purity Milli-Q deionised water (>18 M Ω /cm, Bedford, MA, USA). 109 On the initial day of each test arsenate stock solutions, 0.2 and 15 g/L As(V) using 110 Na₂AsO₄.7H₂O (May and Baker, Dagenham, England), or an arsenite stock solution, 15 g/L

111	As(III) using NaAsO ₂ (BDH, Poole, England), were prepared in test medium. Test media (70
112	mL) for control treatments or arsenic solutions were added to the flasks. Each test included
113	five arsenic concentrations and a control, each prepared in triplicate. For Chlorella sp.,
114	As(III) treatments ranged from 10 to 200 mg/L and As(V) treatments from 0.75 to 60 mg/L.
115	For <i>M. arcuatum</i> , As(III) treatments ranged from 5 to 50 mg/L and As(V) treatments from
116	0.025 to 0.4 mg/L. Nutrients KH ₂ PO ₄ (Ajax) and NaNO ₃ (Merck, Kilsyth, VIC, Australia)
117	were added to all flasks to give a final concentration of 0.15 mg PO_4^{3-}/L and 15 mg NO_3^{-}/L
118	(N:P molar ratio of 150:1). Additional toxicity tests with <i>M. arcuatum</i> , were carried out at a
119	higher phosphate concentration (15 mg NO_3^{-}/L ; 1.5 mg PO_4^{-3-}/L ; N:P molar ratio of 15:1) and
120	a lower nitrate concentration (1.5 mg NO ₃ ⁻ /L; 1.5 mg PO ₄ ³⁻ /L; N:P molar ratio of 150:1).
121	Prior to inoculation with algae, 20-mL subsamples were taken from each flask, pooled
122	for each treatment, for measurement of initial arsenic concentrations and stored at -18° C.
123	Measured concentrations, not nominal, were used to calculate toxicity test endpoints.
124	Cells in the exponential growth phase (5-6 days old) were used in bioassays after
125	centrifugation (2500 rpm, 7 minutes) and washing three times with Milli-Q water to remove
126	residual culture medium. Flasks were inoculated with $2-4 \times 10^4$ cells/mL, shaken by hand
127	and randomly placed in a growth cabinet ($27 \pm 1^{\circ}$ C, $12:12$ h light/dark cycle, Philips TL 40W
128	cool white fluorescent lighting, 140 μ mol photons/m ² /s). Test flasks were rotated, and shaken
129	twice daily to ensure sufficient gas exchange. The pH was recorded initially and after 72 h.
130	Cell density was determined daily using a Coulter Multisizer IIE Particle Analyser (70
131	μ m aperture; Coulter Electronics, Luton, UK), with a correction count of background
132	particles. The cell density data from 0 to 72 h were used to calculate the growth rate of
133	treatments by fitting a regression line to a plot of log_{10} (cell density) versus time (h). The
134	slope of the plot gave the cell division rate (μ) calculated as divisions per day. Growth rates
135	for treated flasks were expressed as a percentage of the control cell division or growth rate.

136

137 Statistical analysis

The 72-h IC50 was calculated using linear interpolation (ToxCalc Version 5.0.23C, Tidepool Software, San Francisco, CA, USA). The data were tested for normality and homogenous variance, and Dunnett's multiple comparison test was used to determine which treatments differed significantly from the controls (1 tailed, $P \le 0.05$) to estimate the NOEC (no observable effect concentration) and the LOEC (lowest observable effect concentration).

144 Intracellular and extracellular arsenic determination

145 The concentration of intracellular and extracellular arsenic after 72-h exposure to 146 As(V) at two different phosphate concentrations (both with 15 mg NO_3/L) was determined to 147 investigate the potential competitive uptake of phosphate and arsenate, using a modified 148 method of Franklin et al [20]. All manipulations were carried out in a Class-100 clean room. 149 M. arcuatum was incubated for 72-h with initial As(V) concentrations of 125, 250 and 150 1000 μ g/L for the low phosphate (0.15 mg/L) tests and concentrations of 250, 1000 and 3000 151 μ g As(V)/L for the high phosphate (1.5 mg/L) tests. Both tests included control treatments 152 (no arsenic). For each treatment, nine flasks were prepared, combining three after 72 h to 153 gain sufficient biomass for analysis, with three replicates per treatment. Each replicate was 154 mixed thoroughly and the cell density determined on a Coulter Multisizer IIE Particle 155 Analyser (70 µm aperture). Weighed sub-samples (145 mL) were filtered through a 25-mm 156 glass filtration unit (pre-acid-washed and rinsed with Milli-Q water) using a 0.45 µm GH-157 polypropylene filter (Pall, Ann Arbor, MI, USA). Approximately 50 mL of the filtrate was 158 collected and frozen until analysis (dissolved As fraction).

159 Cells were rinsed with 20 mL of arsenic-free, nutrient-free growth medium while in 160 the filtration unit to remove excess dissolved As(V) solution and to prevent overestimation of 161 arsenic bound to the outside of cells. This solution was retained for analysis (rinse fraction). 162 Following preliminary experiments, two 20-min washes with 0.1 M KH₂PO₄/K₂HPO₄ 163 buffer solution (pH 5.95) (Ajax, Merck)) were used for the optimum removal of extracellular 164 arsenic without obvious efflux of intracellular arsenic. Phase contrast microscopy showed that cells were healthy and intact after these treatments. Cells on the filter paper were 165 166 transferred to a Teflon tube, using 0.1 M phosphate buffer (final volume of 20 mL). This 167 mixture was shaken for 30 s, allowed to stand for 20 minutes, then re-filtered using a new 168 filter. The filtrate was retained for analysis. The process of cell-washing was repeated. 169 These samples were called the "extracellular" fraction. 170 Algal cells were returned to the Teflon tube using 25% (v/v) HNO₃ (Merck), made up 171 to a volume of 8 mL and left to digest for 30 minutes. The digest was microwaved at 90W for 172 5 min, diluted to 20 mL with Milli-Q water to give a final concentration of 10% (v/v) HNO₃, 173 and stored at 4°C ("intracellular" fraction). 174 The dissolved, rinse and extracellular arsenic fractions were analysed for total arsenic by hydride generation-atomic fluorescence spectrometry (HG-AFS). Because the 10% (v/v) 175 176 acid matrix interfered with the response from the AFS detector, the intracellular arsenic 177 fraction was analysed for total arsenic by ICP-MS (Perkin Elmer Elan-6000, Australia). 178 179 Arsenic speciation bioassays 180 To determine the inorganic and organic arsenic species in solution and in algal cells 181 following 72-h exposure to As(V), two speciation bioassays were conducted for both 182 Chlorella sp. and M. arcuatum. The first bioassay consisted of As(V) treatments of one 183 replicate each of 0, 10, 25 and 40 mg/L for *Chlorella* sp. and 0, 0.1, 0.2 and 0.3 mg/L for M.

arcuatum. The second speciation bioassay consisted of three replicates at one concentration
of arsenic only; 25 mg As(V)/L for *Chlorella* sp. and 0.2 mg As(V)/L for *M. arcuatum*(approximate 72-h IC50). This set-up was required logistically as each replicate required 15
flasks of algae to be combined to gain sufficient biomass for analysis.
The cell density of the pooled bioassay solution was determined and the solution was

filtered and both the filter paper and filtrate were collected for analysis. For the second
speciation bioassay, the cells collected on the filter paper were rinsed with 20 mL of Milli-Q
water to prevent overestimation of cellular arsenic due to carryover of dissolved solution.
This rinse solution was analysed for total arsenic by ICP-MS. The water samples and cellular
samples on the filter paper were frozen immediately following collection and were analysed
for arsenic speciation by microwave digestion and HPLC-ICP-MS.

195 For quality assurance purposes, three additional flasks were prepared for each 196 treatment concentration, and incubated for 72 h under the standard test conditions. Two of 197 these were blanks (no algae), used to determine arsenic speciation changes in solution due to 198 either chemical reduction, or the process of collecting the sample fractions. The third flask 199 (inoculated with algae) was used to check the overall arsenic mass balance, to account for all 200 the added arsenic as either in solution, on the cells or adsorbed to the flask. Adsorption to the 201 flask was determined by filling the empty flask with nitric acid (20 mL, 0.2% (v/v), Suprapur, 202 Merck). It was shaken, left to stand for 48 h and then analysed for total arsenic by ICP-MS. 203

204 Extracellular versus intracellular As(V) reduction

Reduction of As(V) to As(III) by *M. arcuatum* was further investigated to determine whether it occurred intracellularly or extracellularly. Control solutions (no arsenic) and As(V) treatments were inoculated with algae and incubated for 48 h. To test for nonbiological reduction of arsenic in solution, an additional control was prepared containing 209 As(V) but not algae. The solutions containing cells were centrifuged and the supernatant was 210 divided into 6×50 mL sub-samples, of which three were spiked with more As(V). Thus 211 there were three replicates of the following (cell free) solutions: (i) exposed to As(V) plus an 212 additional arsenic spike after 48 h; (ii) exposed to As(V) with no arsenic spike; (iii) not 213 exposed to As(V) plus an additional arsenic spike after 48 h and; iv)not exposed to As(V) 214 with no arsenic spike. All flasks were placed in the growth cabinet for a further 24 h. The 215 solutions were then stored at -15°C until analysis of inorganic arsenic species by HG-AFS. 216 This experiment was done three times, twice with algae exposed to and additional spikes of 217 100 μ g As(V)/L and once with algae exposed to 300 μ g As(V)/L and a spike of 300 218 $\mu g As(V)/L.$

219

220 *Effect of As(V) on cellular thiol groups in M. arcuatum and Chlorella sp.*

To determine whether As(V) reduction to As(III) was coupled to the oxidation of thiol groups such as glutathione in the cell, cellular sulfhydryl groups (-SH) were determined by spectrophotometry with 2-2'-dithiodipyridine using a modified method of Grassetti and Murray [21], adapted for algal cells by Stauber and Florence [22].

Test solutions were prepared in triplicate at three As(V) concentrations for both *M*. *arcuatum* (0 (control), 260 and 500 μ g As(V)/L) and *Chlorella* sp. (0 (control), 25 and 50 mg As(V)/L) using nutrient concentrations of 0.15 mg PO₄³⁻/L and 15 mg NO₃⁻/L. Solutions were inoculated with a high algal cell biomass (2-3 × 10⁵ cells/mL) and incubated for either 24 or 48 h under standard growth conditions.

After the exposure period, treatment flasks were combined and 30 mL (by mass) dispensed into four 50 mL polypropylene centrifuge tubes. Three replicates were exposed to 2-2'-dithiodipyridine, with the last replicate a blank. They were processed as per Stauber and Florence [22], and the absorbance of the samples at 341 nm and 233 nm was measured on a 234 UV-Visible spectrophotometer (Ultrospec IIE, LKB Biochrom, Cambridge, UK). Using a 235 calibration curve generated with freshly prepared 0.001 M reduced glutathione (GSH) 236 solution (Sigma) as a standard, the concentration of free thiols in controls and arsenic-treated 237 cells was calculated. Student t-tests were performed for pairs of As(V) concentrations to 238 determine if differences in the number of thiol groups were significant ($P \le 0.05$).

239

240 Arsenic analyses

HG-AFS, ICP-MS and HPLC-ICP-MS were all used to determine concentrations of total arsenic and arsenic species in solution and in algal cells. All calibration standards were prepared fresh on the day of analysis using matrix-matched solutions.

244 Total arsenic and inorganic arsenic speciation in solution were analysed by HG-AFS 245 using a PSA Excalibur system (PS Analytical, Kent, UK). Total arsenic was measured after 246 oxidative digestion of organics to As(V) in 1% K₂S₂O₈ in an autoclave for 30 minutes 247 (120°C). Quantitative reduction of As(V) to As(III) was achieved by standing for 30 minutes 248 with 32% HCl, 1.3% KI and 0.25% ascorbic acid. Online delivery of 33% (w/v) HCl and 249 1.5% (w/v) NaBH₄ (stabilised in NaOH) converted As(III) to AsH₃ for detection. Total 250 inorganic arsenic was determined by eliminating the persulphate digestion. For As(III) 251 determination, online delivery of 0.3M acetic acid-0.2M sodium acetate and 1.5% (w/v) 252 NaBH₄ (stabilised in NaOH) converted As(III) to AsH₃. Samples were in the same acetic 253 acid- sodium acetate matrix. Matrix-matched calibration curves using As(III) and As(V) 254 standards were generated and the total inorganic As and As(III) concentrations calculated 255 directly, and As(V) calculated by difference. 256 Samples requiring determination of total As in an acidic matrix (e.g. As adsorbed to

257 flask walls and intracellular As) were measured by ICP-MS (Perkin Elmer Elan-6000,

Australia) following a microwave digestion step [23].

259	Where low to trace concentrations of organic arsenic compounds were of interest in
260	cells, microwave-assisted extraction coupled with HPLC-ICP-MS was primarily used for
261	quantitative speciation analysis. Both anion- and cation-exchange HPLC-ICP-MS was used
262	according to the method thoroughly outlined and validated in prior work of Kirby and Maher
263	[24] and Kirby et al. [25]. Because microalgal cell masses were small (1-3 mg) following
264	freeze-drying (Labconco, Australia), they were extracted without subsampling. Calibration
265	curves were prepared using a mixed standard of sodium arsenite, sodium arsenate
266	heptahydrate, sodium dimethylarsenic (Alltech - Specialists, Australia) and disodium
267	monomethylarsenic (Alltech - Specialists, Australia) in Milli-Q water. Characterisation of
268	arsenosugars was done with standards previously isolated and purified as described in Kirby
269	and Maher, [24]. Standards were run at regular intervals throughout sample analysis.
270	
271	RESULTS
272	Toxicity of arsenic to microalgae
273	The effects of As(III) and As(V) on the growth rates of <i>Chlorella</i> sp. and <i>M. arcuatum</i>
274	are shown in Table 1. Growth rates of controls in the toxicity tests ranged from 1.2-1.8
275	doublings/day for <i>M. arcuatum</i> and 1.3-1.7 doublings/day for <i>Chlorella</i> sp., except for one
276	Chlorella sp. test where the growth rate was only 0.9 doublings/day, possibly due to late
277	inoculation at the end of the light cycle. Measured (initial) concentrations of As(III) and
278	As(V) ranged from 68-72% and 69-109% of nominal concentrations, respectively. The pH $$
279	increased by a maximum of 0.5 pH unit for all tests except for three individual M. arcuatum
280	treatments (0, 50, 100 μ g As(V)/L) that increased by up to 1.1 pH units.
281	Algal growth rate decreased as the concentration of arsenic increased. Slight
282	stimulation of algal growth (2-8%) occurred at the lowest arsenic concentrations in some
283	tests. Both species were insensitive to As(III) with 72-h IC50 values of 14.6 and 25.2 mg/L

for *M. arcuatum* and *Chlorella* sp., respectively. Complete growth inhibition (< 5% of
controls) was found at 50 mg As(III)/L for both species. The direct impact of adding As(III)
to these alga was not further investigated due to lack of inter-species sensitivity differences,
As(III) concentrations that caused toxicity were orders of magnitude above expected
environmental concentrations of total arsenic in freshwater and because As(V) is the

thermodynamically-favoured species in oxidised freshwaters [1, 2].

290 Data from the three individual As(V) toxicity tests for *Chlorella* sp. and *M. arcuatum* 291 were combined to determine a concentration-response curve for the toxicity of As(V) to each

alga (Fig. 1). Using non-linear regression, the 72-h IC50 for *Chlorella* sp. was 25.4 mg As/L,

with 95% confidence limits (CL) of 25.2 to 25.7 mg As/L. This alga showed similar

tolerance to both As(III) and As(V), however, both 72-h IC50 values were several orders of

295 magnitude above expected environmental arsenic concentrations. As(V) was about 100 times

more toxic to *M. arcuatum* than *Chlorella* sp., with a 72-h IC50 (95% CL) of 0.254 (0.253-

297 0.255) mg As/L. Significant effects of As(V) on the growth rate of *M. arcuatum* were found 298 at As(V) concentrations as low as 50 μ g/L in one test, however, the mean LOEC value from 299 three tests was 81 μ g/L, with a NOEC of 39 μ g As(V)/L.

The toxic mode of action of As(V) on *M. arcuatum* was of interest due to its greater
sensitivity to As(V) compared with *Chlorella* sp. Thus a number of more detailed
experiments were conducted, using *M. arcuatum*, to try and elucidate the mechanism of
toxicity (see below).

304

305 *Effect of phosphate on As(V) toxicity to M. arcuatum*

When the phosphate was increased to 1.5 mg/L, lowering the N:P ratio in solution
from 150:1 to 15:1, As(V) was much less toxic to *M. arcuatum*. The 72-h IC50 was 4.53
mg As(V)/L, compared to the standard bioassay with a 72-h IC50 of 0.254 mg/L (Table 1).

309 The NOEC and LOEC values also increased approximately 20-fold when the phosphate 310 concentration was increased. To establish that this was a result of changing the phosphate concentration and not changing the N:P ratio, a separate test with a lowered nitrate 311 concentration (1.5 mg NO₃⁻/L) and a molar N:P ratio of 15:1 was conducted (0.15 mg PO₄³⁻) 312 313 /L). In this test, the 72-h IC50 (0.183 mg As(V)/L) was only slightly (but significantly, 314 P<0.05) lower than the 72-h IC50 from the standard test using 15 mg NO₃⁻/L (0.254 mg As(V)/L) (Table 1). This suggests that the ameliorating effect on As(V) toxicity observed in 315 316 the high phosphate growth bioassay (N:P;15:1), was due to increasing phosphate 317 concentration alone.

318

319 *Effect of phosphate on the concentration of intracellular and extracellular arsenic*

The distribution of arsenic in the various algal fractions after 72 h are shown in **Table** 2 for low phosphate (0.15 mg PO_4^{3-}/L) and high phosphate (1.5 mg PO_4^{3-}/L) bioassays. Good recovery of arsenic was obtained (96-103% of the initial arsenic in solution), with most of the arsenic (> 99%) in the dissolved arsenic fraction. Arsenic concentrations in the cellular fractions were low. The concentration of arsenic in all fractions increased with increasing initial arsenic in the media.

Extracellular and intracellular concentrations of total arsenic on a per cell basis are shown in **Fig. 2**. Results are expressed this way to overcome substantially lower total cell numbers at higher arsenic concentrations, due to toxic effects on growth. Preliminary experiments showed that arsenic did not substantially alter the size of *M. arcuatum*, i.e. arsenic load did not change as a result of surface area or volume changes. Extracellular and intracellular concentrations of arsenic increased with increasing concentrations of arsenic added to the growth medium. The concentrations of intracellular and extracellular arsenic were significantly higher (P \leq 0.05) when the bioassay was carried out in low phosphate (0.15 mg PO₄³⁻/L) compared to high phosphate (1.5 mg PO₄³⁻/L) solutions (Table 2).

335

336 Speciation and distribution of arsenic in microalgae

337 The distribution of arsenic after 72-h exposure to As(V) is shown in Fig. 3 for both M. 338 arcuatum and Chlorella sp. Of the As recovered, > 94% was in solution, < 0.01% was 339 associated with the cells and < 1.3% was adsorbed to the flask walls. The amount of total As 340 adsorbed to the flask increased with increasing concentrations of As(V) used in the test 341 medium. Addition of a rinsing step resulted in up to 6.2% of total As being recovered in this 342 fraction, with cellular concentrations of As(V), As(III), DMA and MMA in Chlorella sp. 343 decreasing by 2-4 fold. This highlighted that carryover of dissolved arsenic, in the mg/L 344 range, results in overestimation of cellular arsenic.

345 As(III) was present in test media containing *M. arcuatum* after a 72-h exposure to 346 As(V) (Fig. 3). The percentage of initial As(V) reduced to As(III) decreased from 95% to 347 22% with increasing initial As(V) concentrations. However, no As(III) was detected in the 348 blanks (no algae), indicating that the presence of As(III) in M. arcuatum solutions was due to 349 biological reduction. In a separate As(V) exposure test, inorganic As concentrations were 350 measured at 24-h intervals throughout the 72-h bioassay with *M. arcuatum*. As(V) reduction 351 to As(III) was observed in the initial 24-h period and the reduction continued over time: 41% 352 and 65% of the 260 µg As(V)/L treatment was detected as As(III) at 24 and 48 h,

353 respectively; and 46% and 72% of the 500 μg As(V)/L treatment was detected as As(III) at 24

and 48 h. Between 48 and 72-h, As(V) reduction to As(III) was similar.

355 In contrast, As(III) was not detected in solution after 72 h in *Chlorella* sp. bioassays

356 (Fig. 3). There was no detectable MMA, DMA, phosphate arsenoriboside or other organic

357 species of arsenic in solution at 72 h for either algae.

358	Five arsenic species were detected in the cells; As(V), As(III), MMA, DMA and
359	phosphate arsenoriboside (P-sug) (Fig. 4). No other arsenic species were detected in any of
360	the tests. Cells contained predominantly As(V) followed by As(III). Although cellular
361	concentrations of arsenic species in Chlorella (after rinsing) were a maximum of 6-fold
362	higher than arsenic species accumulated by M. arcuatum, M. arcuatum had been treated with
363	concentrations of As(V) 100 times lower than Chlorella. Thus M. arcuatum accumulated
364	more arsenic from solution relative to Chlorella sp., and was consequently more sensitive to
365	As(V) than <i>Chlorella</i> sp.
366	For Chlorella sp., concentrations of cellular As(V) and As(III) generally increased
367	with increasing $As(V)$ concentrations in the growth medium, in contrast to <i>M. arcuatum</i>
368	where concentrations of As(V), DMA and phosphate arsenoriboside were at a maximum in
369	the 0.210 mg/L (rinsed cell) treatment (Fig. 4). Trace concentrations of phosphate
370	arsenoriboside were occasionally detected in Chlorella cells. Higher amounts of phosphate
371	arsenoriboside were detected in <i>M. arcuatum</i> with a mean value of $44.7 \pm 19.6 \times 10^{-18}$ g/cell

- in the 0.210 mg As/L treatment.
- 373

374 Cellular reduction of arsenic(V) to arsenic(III) by M. arcuatum

375 Preliminary experiments demonstrated that *M.arcuatum* did not inherently produce an376 exudate that could reduce arsenic.

377 *M. arcuatum* exposed to 0.1 mg/L and 0.3 mg/L As(V) for 48 h reduced the As(V) in

solution to As(III) by approximately 41% (0.04 mg As(III)/L) and 11% (0.03 mg As(III)/L),

379 respectively. There was no reduction of As(V) to As(III) in the growth medium in the

absence of *M. arcuatum* cells, indicating that arsenic reduction was biologically mediated.

381 After the initial 48-h exposure (control, 0.1 or 0.3 mg/L As(V)), the cells were

removed and the supernatant spiked with an additional 0.1 or 0.3 mg As(V)/L for a further 24

h. No further reduction of As(V) was observed in this period. This showed that the reduction of As(V) to As(III) only occurred in the presence of cells, i.e. the reduction was not due to an exudate released by the cells, generated in the presence of As(V).

386

387 *Effect of As(V) on cellular thiol groups in M. arcuatum and Chlorella sp.*

Oxidation of thiols such as glutathione has previously been shown to be a potential mechanism by which cell division is inhibited by metals in algal cells [19]. The hypothesis was that reduced glutathione (GSH) is oxidised (to GSSG) as As(V) is reduced to As(III). Decreased SH concentrations for As(V) treatments compared to controls (no As(V)) indicated

551 Decreased STI concentrations for Tis(t) inclusions compared to controls (no Tis(t)) indicated

392 that thiol groups were oxidised. Preliminary experiments with As(V) and glutathione in cell-

393 free solution, showed that As(V) did not oxidise GSH to GSSG in the absence of algal cells.

394 Arsenic toxicity and As(V) reduction to As(III) were similar when both high $(3 \times 10^5$

395 cells/mL) and low $(2-4 \times 10^4 \text{ cells/mL})$ initial cell densities were used.

396 In unexposed controls, *M. arcuatum* contained 8.2 ± 1.9 nmol SH 10⁻⁶ cells. Thiols

397 significantly decreased as the concentration of arsenic and time of exposure increased,

398 however, results were variable. After a 24-h exposure to $500 \ \mu g \ As(V)/L$, thiol

399 concentrations were significantly lower (P < 0.05) in two of the three tests (16% and 57% of

400 controls). After a 48-h exposure, thiol concentrations were 15 and 37% of controls in two of401 the three tests.

402 A similar pattern was observed for *Chlorella* sp., with a mean number of thiols of 7.9 403 \pm 3.3 nmol SH 10⁻⁶ cells in unexposed controls. After 48-h exposure to 25 mg As(V)/L (the 404 approximate IC50 for *Chlorella* sp.) thiol concentrations were significantly decreased (16-405 75% of controls). After a 72-h exposure, thiol concentrations were also decreased (55% of 406 control and 14% of control at 25 and 50 mg As(V)/L, respectively). When *Chlorella* sp. was 407 exposed to much lower arsenic concentrations, similar to that of *M. arcuatum* (i.e. 500 µg 408 As(V)/L) thiol concentrations were not significantly different after 24 h (92% of control) but 409 were significantly lower (P <0.05) after a 48- and 72-h exposure (61 and 68% of controls). 410 The variation observed is these results is likely due to the cells being damaged but not lysed 411 by the addition dithiodipyridine solution and vortexing. Thus, conversion of internal thiols 412 may differ between experiments. 413 414 DISCUSSION 415 Arsenic toxicity 416 The current results confirm that the toxicity of arsenic to freshwater microalgae 417 depends on the chemical species of arsenic, the algal species and the phosphate concentration 418 in the test medium. Arsenate and arsenite were approximately equally toxic to Chlorella sp., 419 with IC50s of 25.4 mg As(V)/L and 25.2 mg As(III)/L). M. arcuatum was more sensitive to 420 As(V) (IC50: 0.254 mg As(V)/L) than *Chlorella* sp. and more sensitive to As(V) than As(III) 421 (IC50: 14.6 mg As(III)/L). 422 The 12-14-d growth inhibition IC50 values for As(V) spanned five orders of 423 magnitude for Scenedesmus obliquus, Ankistrodesmus falcatus, Selenastrum capricornutum, 424 Scenedesmus quadricauda and Chlamydomonas reinhardtii (0.048, 0.256, 31, 61, 202 mg/L, 425 respectively) [10,11,12]. Although different test durations and conditions such as 426 photoperiods and phosphate concentrations make comparison difficult, this illustrates that 427 even in a single genus, there are large variations in sensitivity of microalgae to arsenic. 428 The toxicity of arsenic to freshwater microalgae is also dependent on the chemical 429 species of arsenic added. It has been reported that As(V) is more toxic than As(III) to 430 freshwater algae, while the reverse is true for marine algae and humans [1,26]. In 96-h 431 growth inhibition tests with the freshwater green alga Selenastrum capricornutum, IC50 432 values of 31 and 0.69 mg As/L were found for As(III) and As(V) respectively [27], while the

437 inhibition at As(III) concentrations > 40 mg/L [9].

438 Our results showed that a ten-fold increase in the phosphate concentration decreased 439 the toxicity of As(V) to *M. arcuatum* by approximately twenty-fold (Table 1). The reduced toxicity of As(V) was a result of higher phosphate concentrations, rather than simply due to 440 441 changing the N:P ratio (Table 1). The concentration of phosphate in solution significantly 442 affected the amount of arsenic adsorbed to the surface of M. arcuatum and the amount of 443 arsenic that was accumulated inside the cell (Table 2). At low phosphate concentrations, 444 intracellular and extracellular arsenic concentrations were high, corresponding to increased growth inhibition, compared to the bioassays carried out at high phosphate concentrations. 445 446 With an increase in phosphate concentration in the bioassay medium, less arsenic binds to the 447 algal cell, and less arsenic is taken up intracellularly (Fig. 2), supporting the hypothesis that 448 arsenate and phosphate compete for uptake in algal cells. This further supports the study by 449 Maeda et al [29] which showed that the toxic effect of 10 mg As(V)/L to Chlorella vulgaris decreased when the phosphate concentration increased from 14 to 14000 mg PO_4^{3-}/L . 450 451 However, these authors used high arsenic concentrations (1-1000 mg As(V)/L), high cell 452 densities, and an isolate from a contaminated environment. High cell densities decrease the 453 toxic load to cells [20], while there exists the potential for adaption and species succession in 454 polluted environments. Consequently Chlorella vulgaris was very tolerant to As(V) (52% growth inhibition at 5 g As(V)/L) when compared M. arcuatum in our study (IC50: 254 μ g 455 456 As(V)/L). Maeda et al. [30] also determined intracellular and extracellular As in Chlorella vulgaris using only water to remove extracellular arsenic. The amount of arsenic adsorbed to 457

458 and accumulated inside the cells increased 10-fold with each 10-fold increase in the 459 concentration of arsenic in the test medium. Similar trends were observed in *M. arcuatum* in 460 our current study, using a phosphate buffer to desorb As from the algal cell surface.

While phosphate has been shown to affect arsenate uptake into *M. arcuatum*, it is not known if arsenate reduces phosphate uptake into the alga, thereby contributing to inhibitory effects of arsenate on algal growth. However, because phosphate is an essential nutrient, competition between arsenate and phosphate for cellular uptake is likely to be one mode of toxic action in microalgae. Increases in arsenic have been shown to decrease phosphate uptake in five freshwater algae, *Anabaena variabilis, Chlamydomonas reinhardtii*,

467 *Cryptomonas erosa, Melosira granulata* and *Ochromonas vallesiaca* [31]. In contrast, it was

468 found that phosphate uptake in *Synechococcus leopoliensis*, a cyanophyte, was not affected by

arsenate even when the concentration of arsenate was fifty times that of phosphate, possibly

470 because this species had a highly specific phosphate transport system [32].

471

472 Arsenic accumulation and biotransformation

Accumulation of arsenic by freshwater microalgae typically increased with increasing
arsenate concentrations in the test medium (Fig. 4), similar to other studies [18,29,32].

475 Maeda et al. [9] also showed that accumulation of As only occurred in live *C. vulgaris* cells,

476 suggesting an active uptake mechanism.

In our studies, As(V) was the main arsenic species in cells, followed by 1-6% as
As(III). Maeda et al. [33] also found that >95% of arsenic was accumulated by freshwater
algae as inorganic species. They found that dimethylated arsenic was the major methylated
arsenic compound detected. However, while both *Chlorella* sp. and *M. arcuatum* in our study
methylated As(V) to MMA, DMA and phosphate arsenoriboside (Fig. 4), these products were
present only in low concentrations in the cells and were not detectable in solution.

483 In our study 0-12% of cellular arsenic occurred as the phosphate arsenoriboside. 484 Arsenoribosides have only recently been positively identified in one freshwater alga, 485 *Chlorella vulgaris*, with comparison to the retention time of arsenoriboside standards. 486 Glycerol, phosphate and sulfonate arsenoribosides were detected, with phosphate 487 arsenoriboside occurring in the highest concentration of 0.2-5% of accumulated arsenic [7]. 488 Agar plating was not carried out, but no significant differences occurred between cultures 489 treated with and without antibiotics. Arsenoribosides have also been identified in Nostoc 490 *flagelliforme*, a terrestrial cyanobacterium [34]. It is possible that the arsenoribosides 491 detected were produced by bacteria in the cultures rather than the microalgae themselves, but 492 our study with bacteria-free algal cultures confirms that microalgae exposed to low arsenic 493 concentrations can produce trace arsenoribosides, but it does not appear to be a major 494 detoxification pathway.

495 Bioassays with exponentially growing cells showed that when *M. arcuatum* (but not 496 *Chlorella* sp.) was exposed to As(V), As(III) was excreted into solution. Hellweger et al. [35] 497 found that As(III) excretion into solution was more likely during the exponential phase of 498 growth. Algae in the stationary phase of growth (phosphate limited) were more likely to 499 methylate arsenic to more complex organic arsenic compounds, which are then excreted. 500 Similarly, it was found that when Chlorella vulgaris accumulated inorganic, mono- and di-501 methylated arsenic over 20 days, it excreted inorganic As together with trimethylated arsenic 502 species after 4 days and dimethylated arsenic species after 14 days [18]. Trivalent 503 methylarsenic species have also been detected in the growth medium of the green alga 504 Closterium aciculare [36] but were not detected in our current study using HPLC-ICP-MS. It 505 is possible that, if methylation of arsenic occurs to a greater extent in stationary phase cells, 506 then larger concentrations of arsenoribosides may have been detected if stationary phase 507 rather than exponentially growing algae had been used in our experiments.

As(V) reduction to As(III) occurs intracellularly (or in the cell membrane) in *M*. *arcuatum* and the As(III) is then excreted into the test medium. Similar trends have also been found in bacteria and yeasts [14], with As(V) reduced to As(III) via an arsenate reductase and then removed from the cytosol by either a secondary carrier, using energy from an existing ion gradient, or in a complex with a second protein via an ATP-coupled pump.

513 Our study supports the biotransformation model of Cullen et al. [17,37] in which 514 arsenate is taken up by algal cells using a phosphate transport system, reduced to As(III) in 515 the cell by thiols and/or dithiols and then excreted into the growth medium, probably by an 516 active transport system. At longer exposure times, As(III) may be methylated to MMA, then 517 to DMA and trimethylated arsenic species, which then diffuses into the growth medium.

518 In the marine microalga Nitzchia closterium, toxicity of copper was shown to be a 519 cytosolic reaction between copper and GSH [22]. The cellular ratio of GSH:GSSG, critical to 520 mitotic cell division, was lowered. We hypothesised that in *M. arcuatum*, reduction of As(V) 521 to As(III) may be coupled with oxidation of GSH, ultimately resulting in inhibitory effects on 522 cell division. If this was the case, total thiol concentrations in the cells should be reduced in 523 the presence of arsenate, at concentrations that are inhibitory to algal growth. Thiol cell 524 concentrations were lower in *M. arcuatum* at high concentrations of As(V) (0.5 mg As(V)/L) 525 at 24 and 48-h compared to controls, but variability in the results suggest improvements must 526 be made to this technique for freshwater algae before strong conclusions can be made (as 527 using marine algae, e.g. [22], osmotic shock effectively lyses the cell). Excretion of As(III) 528 may not keep pace with arsenic reduction, leading to accumulation of As(III) in the cells. 529 As(III) is known to bind strongly to thiols in plants and animals [13]. As(III) appears to only 530 be toxic once accumulated inside cells, as As(III) in the medium was not toxic to either M. 531 arcuatum or Chlorella sp.

In *Chlorella* sp., thiol oxidation was also observed at As(V) concentrations that inhibit cell division (25 mg As(V)/L). This indicates that As(V) reduction may be coupled to thiol oxidation, but the alga lacks the arsenite transporter to excrete As(III) into the medium. It is possible that *Chlorella* sp. is able to detoxify arsenite inside the cell by sequestering it into subcellular compartments much like the yeast *Saccharomyces cerevisiae* complexes As(III) with glutathione, transferring the product from the cytosol into vacuoles via a specific transporter [14].

539 In freshwater environments, arsenic is unlikely to be toxic to *M. arcuatum*, except in 540 highly contaminated surface and groundwaters containing $>50 \mu g As/L$. In such 541 environments, it is likely that As(V) is taken up by algal cells due to its similarity to 542 phosphate, and is quickly reduced to As(III). Toxicity is most likely due to the presence of 543 As(III) in the cell, when the ability to excrete or sequester As(III) is overwhelmed and the 544 As(III) subsequently binds to intracellular thiols, inhibiting cell division. The disruption of 545 phosphate metabolism by incorporation of As(V) into phosphorylated compounds, vital to the 546 cycling of ATP, may also contribute to arsenic toxicity.

547

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	[NO ₃ ⁻]	[PO ₄ ³⁻]	N:P	72-h IC50 ^a (mg/L)	LOEC ^b (mg/L)
	(mg/L)	(mg/L)	(molar)		
As(III)					
Chlorella sp.	15	0.15	150:1	25.2 (23.3-29.2) ^c	_d
M. arcuatum	15	0.15	150:1	14.6 (11.7-17.7) ^c	3.75 ^c
As(V)					
Chlorella sp.	15	0.15	150:1	25.4 (25.2-25.7)	1.93
M. arcuatum	15	0.15	150:1	0.254 (0.253-0.255)	0.081
<i>M. arcuatum</i> - high PO_4^{3-}	15	1.5	15:1	4.53 (4.02-4.83) ^c	1.91 ^c
<i>M. arcuatum</i> - low NO ₃ ⁻	1.5	0.15	15:1	0.183 (0.170-0.192) ^c	0.054 ^c

Table 1. 72-h toxicity of As(III) and As(V) to *Chlorella* sp. and *Monoraphidium arcuatum*, under different nutrient conditions

^a IC50: concentration of As which inhibits growth rate by 50%, calculated from a

concentration response curve developed from 3 separate growth inhibition toxicity tests,

unless otherwise indicated. Brackets indicate 95% confidence limits

^b LOEC: lowest-observable-effect concentration, calculated as the geometric mean of three

LOECs from three separate tests, unless otherwise indicated

^c Results are calculated from a single growth inhibition toxicity test

^d LOEC > IC50 therefore not reported

[PO ₄ ³⁻]	Initial	Initial	%	Dissolved	As in	Extracellular As	Intracellular As
(mg/L)	nominal	measured	Recovery ^c	As (µg/L)	rinse ^d	$(\times 10^{-18} \text{ g/cell})^{\text{e,f}}$	$(\times 10^{-18} \text{ g/cell})^{\text{f}}$
	[As] (µg/L)	$[As] (\mu g/L)^b$			(µg/L)		
0.15	0 (control)	< 0.5	-	0 ± 0	0.0 ± 0.0	140 ± 15	ND ^g
	125	123	102 ± 1	125 ± 2	0.7 ± 0.0	1100 ± 270	1200 ± 390
	250	236	103 ± 0	244 ± 1	1.4 ± 0.2	1400 ± 120	2400 ± 380
	1000	1000	98 ± 2	985 ± 19	5.7 ± 0.3	3900 ± 410	2600 ± 110
1.5	0 (control)	< 0.5	-	0 ± 0	0.0 ± 0.0	15 ± 34	12 ± 21
	250	247	96 ± 1	237 ± 2	1.4 ± 0.4	40 ± 74	180 ± 22
	1000	935	101 ± 2	945 ± 22	7.5 ± 2.5	110 ± 50	400 ± 140
	3000	2880	99 ± 2	2840 ± 43	20 ± 1.5	1100 ± 100	1600 ± 330

Table 2. Mean arsenic distribution in *M. arcuatum* fractions after 72-h exposure to varying As(V) and PO_4^{3-} concentrations ^a

^a Mean calculated from 3 replicates, \pm one standard deviation (SD) from the mean

^b No SD indicated for initial measured arsenic as it was calculated from 3 pooled sub-samples

^c % Recovery = total As/measured initial As concentrations \times 100

^d Rinse was performed to prevent overestimation of extracellular As due to carryover from dissolved fraction.

^e Extracellular As is the combination of both phosphate washes

 $^{\rm f}$ Extracellular blank (phosphate buffer) was 0.6 (± 0.2) µg As/L and intracellular blank (acid matrix) was <0.1 µg As/L

^g ND not detected

Fig. 1. Effect of As(V) on 72-h growth rate of *Monoraphidium arcuatum* (\blacklozenge) and *Chlorella* sp. 12 (\Box). Concentration-response curves were based on combined data from three toxicity tests. Error bars represent one standard deviation of three replicates.

Fig. 2. Intracellular and extracellular arsenic concentrations in *M. arcuatum* when bioassays were carried out with varying As(V) concentrations (0-3000 μ g As(V)/L) at low and high phosphate concentrations (0.15 and 1.5 mg PO₄³⁻/L). Note that the legend is based on nominal concentrations of As, measured initial concentrations are given in Table 2 and vary slightly between low and high phosphate tests.

Fig. 3. Mass balance of arsenic species in solution after 72 hours of growth of (a) *Monoraphidium arcuatum* and (b) *Chlorella* sp. 12. As(V) and As(III) in solution, total arsenic (TAs) in the cells and adsorbed to the flask walls were measured for all test treatments. * This column in each figure is the average from 3 separate bioassays run with 0.210 and 26.4 mg As(V)/L for *M. arcuatum* and *Chlorella* sp. respectively; these tests incorporated a rinsing step of the algal cells to investigate the As carryover from solution to cells in the subsequent analysis, and thus a rinse fraction is shown only for these test treatments.

Fig. 4(a) Concentration of As species in *Chlorella* sp. 12 after 72-h exposure to 8.80-39.6 mg As(V)/L. Values indicated for 26.4 mg As(V)/L were the result of triplicate speciation bioassays which incorporated a rinsing step prior to analysing the cells. Values for control, 8.8 and 39.6 mg As(V)/L were the result of a single speciation bioassay and did not incorporate a rinsing step prior to analysing the algal cells. (b) Concentration of As species in Monoraphidium arcuatum after 72-h exposure to 0.103-0.298 mg As(V)/L. Values indicated for 0.210 mg As(V)/L were the result of triplicate speciation bioassays which incorporated a rinsing step prior to analysing the cells. Values for control, 0.103 and 0.298 mg As(V)/L were the result of a single speciation bioassay and did not incorporate a rinsing step prior to analysing the cells. Values for control, 0.103 and 0.298 mg As(V)/L were the result of a single speciation bioassay and did not incorporate a rinsing step prior to analysing the cells. Values for control, 0.103 and 0.298 mg As(V)/L were the result of a single speciation bioassay and did not incorporate a rinsing step prior to analysing the cells. WMA = monomethylarsonic acid; DMA = dimethylarsinic acid; P-sug = phosphate arsenoriboside.

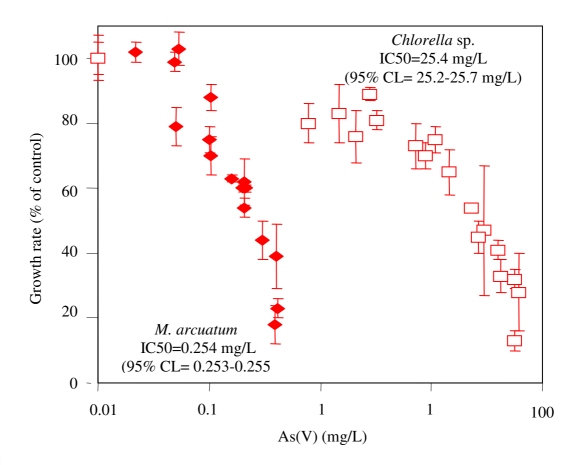


Fig. 1

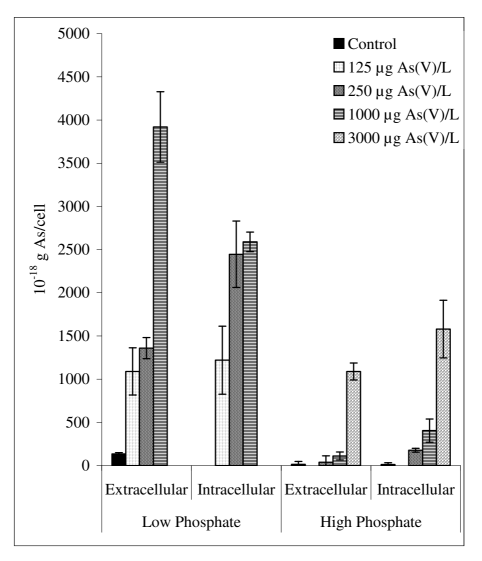
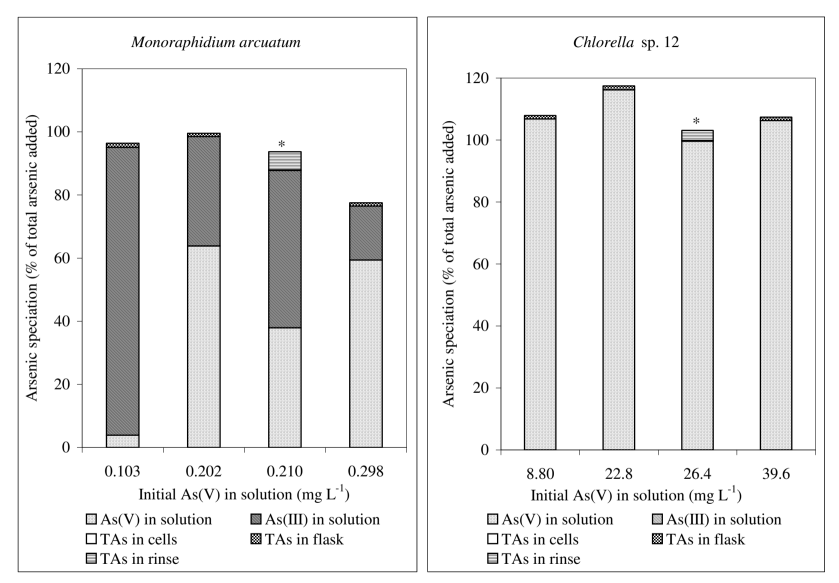


Fig. 2





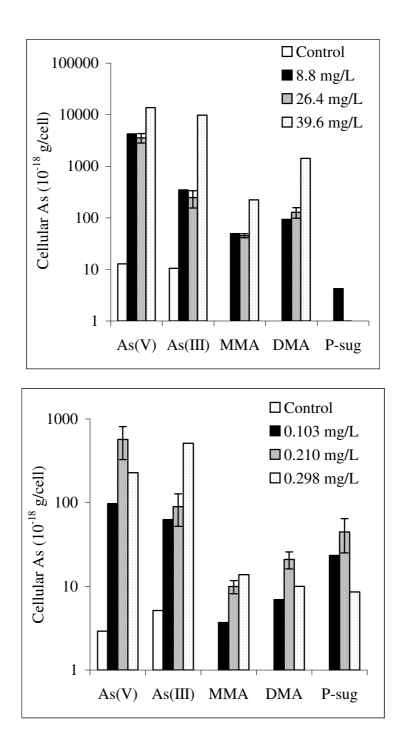


Fig. 4.