

1 Portable and rapid arsenic speciation in synthetic and
2 natural waters by an As(V)-selective chemisorbent,
3 validated against anodic stripping voltammetry

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18 Abstract

19

20 Inorganic arsenic speciation, i.e. the differentiation between arsenite and arsenate, is an important step
21 for any program aiming to address the global issue of arsenic contaminated groundwater, whether for
22 monitoring purposes or the development of new water treatment regimes. Reliable speciation by easy-
23 to-use, portable and cost-effective analytical techniques is still challenging for both synthetic and
24 natural waters. Here we demonstrate the first application of an As(V)-selective chemisorbent material
25 for simple and portable speciation of arsenic using handheld syringes, enabling high sample
26 throughput with minimal set-up costs. We first show that ImpAs efficiently removes As(V) from a
27 variety of synthetic groundwaters with a single treatment, whilst As(III) is not retained. We then
28 exemplify the potential of ImpAs for simple and fast speciation by determining rate constants for the
29 photooxidation of As(III) in presence of a TiO₂ photocatalyst. Finally, we successfully speciate
30 natural waters spiked with a mix of As(III) and As(V) in both Indian and UK groundwaters with less
31 than 5 mg L⁻¹ dissolved iron. Experimental results using ImpAs agreed with anodic stripping
32 voltammetry (ASV), a benchmark portable technique, with analysis conditions optimised here for the
33 groundwaters of South Asia. This new analytical tool is simple, portable and fast and should find
34 applications within the overall multi-disciplinary remediation effort that is taking place to tackle this
35 worldwide arsenic problem.

36

37 Keywords

38 Groundwater, water contamination, ion-exchange resin, electrochemistry, molecular recognition, solid
39 phase extraction

40

41 1. Introduction

42

43 Up to 200 million people worldwide are at risk from arsenic contaminated groundwater (Naujokas et
44 al., 2013). These waters are distributed globally with a variety of sources such as reducing aquifers in
45 the sediment basins of South Asia, oxidising aquifers in arid regions of Mexico and Chile, mining
46 activity in Burkina Faso, the USA and the UK, and geothermal waters (USA, New Zealand) (Smedley
47 2008). Arsenic contaminated waters thus display a wide range of properties, differing in pH, redox
48 conditions, presence of organic matter/bacteria, and the concentration of inorganic anions
49 (bicarbonate, phosphate, sulphate) and cations such as Fe(II) and Mn(II) which precipitate once
50 exposed to air. It is, therefore, not surprising that environmental arsenic presents itself as a range of
51 species, differing in redox state, protonation state, and incorporation of organic groups (Cullen and
52 Reimer, 1989).

53 Knowledge of arsenic speciation in groundwater systems is important for many reasons. Firstly, at pH
54 5-8, typical for most groundwaters, arsenic is present as inorganic H_3AsO_3 (“arsenite” or As(III)) and
55 $\text{H}_2\text{AsO}_4^-/\text{HAsO}_4^{2-}$ (“arsenate” or As(V)). Since As(III) is up to 60 times more toxic than As(V) in
56 humans, knowing the speciation is important when assessing public health risk (Shahid et al., 2018)
57 (Markley and Herbert, 2009). Secondly, since H_3AsO_3 is neutral, its removal is not efficiently
58 achieved through typical water treatment processes such as coagulation-flocculation or adsorption.
59 Instead, addition of a pre-oxidation step is needed (Hering et al., 2017). Thirdly, knowledge of the
60 speciation can help understand the cycling of arsenic within the environment (Shankar et al., 2014).

61 Speciation of inorganic arsenic in groundwater is thus needed but rarely achieved. For example, large
62 scale surveys of groundwater contamination including the 5 million tube wells tested in Bangladesh in
63 2000-2003 (Johnston and Sarker, 2007) have been limited to determination of total As. While arsenic
64 speciation is now routinely achieved using hyphenated techniques such as high-performance liquid
65 chromatography mass spectrometry (HPLC-ICP-MS), these remain expensive, time consuming, and
66 limited to specialist laboratories. For field samples, on-site speciation is preferable in order to avoid

67 problems of sample storage, primarily the progressive oxidation of As(III) on an hours-to-days
68 timescale (Gibbon-Walsh et al., 2011), however field speciation continues to be a specialist pursuit
69 and surveys of arsenic groundwater contamination around the world mostly continue to be limited to
70 determination of total arsenic (McGrory et al., 2017) (Rabbani et al., 2017) (Liang et al., 2018).

71 Reliable, low-cost and easy-to-use analytical techniques are also needed for the speciation of arsenic
72 in synthetic media. Research groups in industry and academia continue to investigate the chemistry of
73 arsenic including (i) chemical oxidation (e.g. Fenton's reagent) (Wang et al., 2013), (ii)
74 photooxidation (Chai et al., 2017), (iii) competitive adsorption between As(III) and As(V) (Qi and
75 Pichler, 2017), and (iv) oxidation over naturally occurring mineral surfaces (Ding et al., 2015), all
76 requiring reliable speciation techniques. Whilst previous studies often employ costly techniques
77 requiring specialist knowledge and time to set-up, such as hydride generation atomic fluorescence
78 spectrometry (HG-AFS) and HPLC-ICP-MS, many laboratories lack access to this specialised
79 analytical instrumentation.

80 Portable and low-cost arsenic speciation has been achieved by various techniques, including
81 colorimetric and electrochemical methods. The preferred molybdenum blue method for colorimetric
82 determination of arsenic, wherein a blue colour-complex is formed between molybdate and As(V),
83 suffers from a well-known phosphate interference. This is a major problem, particularly since
84 contaminated groundwaters in the Bengal Basin contain ten times more phosphate than arsenic
85 (McArthur et al., 2004) (Hug et al., 2008). Colorimetric speciation therefore necessitates a number of
86 chemical pre-treatments to remove phosphate and interconvert As(III) and As(V), complicating field
87 analysis (Okazaki et al., 2013). Speciation of arsenic by electrochemical means does not suffer from
88 phosphate interference and is best achieved using gold electrodes (Huang and Dasgupta, 1999), but
89 historically the perceived lack of "reliability, selectivity and sensitivity" has prevented wider use
90 (Brainina et al., 2000). More recently, several systems have been developed and successfully
91 deployed in environments including reducing groundwaters (Gibbon-Walsh et al., 2011), however
92 commercial uptake has still been limited (Borrill and Reily, 2019).

93 An alternative method is to separate arsenite and arsenate immediately at the time of sampling for
94 later detection under laboratory conditions (Chen et al., 2019) (O'Reilly et al., 2010) (Watts et al.,
95 2010). This can be achieved by passing the solution through a sorbent that is selective to As(V) but
96 leaves As(III) unaffected, for which chloride and acetate anion exchange resins have been used (solid
97 phase extraction, SPE) (Mihucz et al., 2017) (Bednar et al., 2004). By measuring total As before and
98 after treatment, the inorganic speciation can be determined. There are two potential drawbacks to ion-
99 exchange resins. Firstly, these resins are selective to a wide range of anions beside As(V). In
100 groundwaters such as the Bengal Basin, phosphate, sulphate and carbonate are 1, 2, and 4 orders of
101 magnitude more prevalent than arsenic (McArthur et al., 2004) (Ayers et al., 2016) and much
102 adsorption capacity may be lost to competing anions (particularly sulphate) (Pakzadeh and Batista,
103 2011), with resins reaching saturation more quickly than intended (Voice et al., 2011). Secondly is
104 the leaching of exchanged ions into the effluent, particularly chloride, where the polyatomic $^{40}\text{Ar}^{35}\text{Cl}^+$
105 ion has historically been an interference with the ^{75}As peak during ICP-MS analysis (Benramdane et
106 al., 1999), although modern instruments with collision cells now counter this problem (An et al.,
107 2015). Finally, SPE often requires additional elution steps to remove sorbed As(III) (Zhao et al.,
108 2019) (Montoro Leal et al., 2018) (Chen et al., 2016).

109 To date, no study has considered the application of arsenic-selective chemisorbent materials towards
110 the analytical speciation of arsenic. We recently reported an As(V)-selective receptor immobilised
111 onto polymeric beads yielding a chemisorbent material (ImpAs) with a very high affinity for arsenate
112 (patent number US 2017/0113949 A1) (Moffat et al., 2014). The active component of ImpAs is a
113 metal-organic complex containing two Zn(II) centres to which As(V) oxyanions coordinate. ImpAs
114 shows the greatest affinity towards As(V) at pH 6-8 and is thus suitable for most global ground waters
115 (Moffat et al., 2014). Chemisorbent resins offer an interesting new route towards arsenic speciation,
116 as their high selectivity reduces interference from other anions which may enable easy application in
117 challenging waters.

118 This work aims to assess the performance of ImpAs for the analytical speciation of arsenic in
119 synthetic and natural waters. We first tested the ability of ImpAs to separate As(V) from As(III) for

120 laboratory applications using simple buffer solutions and complex synthetic groundwaters. We
121 considered both Challenge Water (NSF Standard 53), a standard used to certify arsenic remediation
122 products in the USA, and a second synthetic groundwater, representative of arsenic-contaminated
123 groundwaters in West Bengal, India. We then applied ImpAs for the determination of As(III)
124 photooxidation rates (catalysed by TiO₂). Finally, we tested the ability of ImpAs to separate As(V)
125 from As(III) in natural waters, using samples from oxic waters in Cornwall, UK, and from anoxic
126 waters in West Bengal, India - regions both known for arsenic contamination. All experiments were
127 conducted in conjunction with anodic stripping voltammetry (ASV) used as the benchmark technique
128 for portable arsenic speciation, to validate the results of speciation (Figure 1).

129

130 2. Experimental

131

132 2.1. Materials and Reagents

133

134 All chemicals used were reagent or analytical grade (SI Table S1). Stock solutions (1000 mg L⁻¹, 13.3
135 mM) of As(V) and As(III) were prepared from Na₂HAsO₄·7H₂O and As₂O₃ respectively. Stock
136 solutions were calibrated against a standard solution of arsenic (Fluka, 1000 ± 4 mg L⁻¹, traceCERT)
137 and kept refrigerated. As(III) solutions were contained within opaque containers to prevent
138 photooxidation. Stock solutions of 24 mM aqueous hydrazine were prepared from hydrazine hydrate
139 or hydrazinium sulphate salt. Stock solutions of 0.5 M ethylenediaminetetraacetic acid (EDTA) were
140 prepared from EDTA tetrasodium salt, acidified to pH 8. A stock solution of 60 mM Fe(II) (pH 2.1)
141 was prepared from FeSO₄·7H₂O. A stock solution of 10 mM Mn(II) was prepared from MnCl₂·4H₂O.
142 A solution of 10 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) was prepared
143 from HEPES free acid and adjusted to pH 7.44 with small volumes of 1 M NaOH.

144 Synthetic groundwaters CW (pH 7.42), SGW-1 (pH 7.09) and SGW-2 (pH 8.32) were prepared to the
145 concentrations given in Table 1, following the procedure given in the Supplementary Information.

146 ImpAs was prepared as previously reported (Moffat et al., 2014), however improvements in the
147 synthesis increased the As(V) capacity to at least 16.0 ± 1.3 mg g⁻¹ (SI Figure S1).

148

149 2.2. ImpAs chemisorbent

150

151 2.2.1. Batch experiments

152

153 To verify that ImpAs does not remove As(III) from solution, batch experiments under equilibrium
154 conditions were carried out. Three background media were used: 10 mM HEPES, CW, and SGW-1.
155 Solutions of 1 mg L⁻¹ total As (13 μM) in ratios of 0:100, 25:75, 50:50, 75:25 and 100:0 As(III) to
156 As(V) were prepared in each media using 100 mg L⁻¹ (1.3 mM) As(III) and As(V) stock solutions. To
157 each solution 0, 1 or 20 g L⁻¹ ImpAs was added. Solutions were shaken overnight in an incubated
158 shaker at 25°C and filtered gravimetrically to remove ImpAs. Samples were kept wrapped in
159 aluminium foil to prevent photooxidation of As(III). As(III) and total As were determined by ASV
160 within 24 hours, and total As was determined by ICP-MS.

161

162 2.2.2. Syringe experiments

163

164 As a prototype portable device for high sample throughput, disposable plastic syringes (5 mL volume)
165 were loaded with ImpAs. To investigate the influence of mass loading, 0.2, 0.4 or 1 grams of ImpAs
166 were used, giving path lengths of 2.5, 5 and 12 mm respectively (Figure 2). Solutions of 1 mg L⁻¹
167 As(V) (13 μM) in 10 mM HEPES, CW or SGW-1 were prepared (total volume 50 mL). Each
168 treatment consisted of loading 5mL aliquots of solution into the top of the syringe by pipette, and
169 passing through the syringe until the total 50 mL had been processed. Sequential treatments were
170 performed, with 1 mL collected after each for determination of total As by ICP-MS. The flow rate
171 was controlled by hand, with solutions passed through the syringes dropwise (1 drop per second, *ca.* 1
172 minute per 5 mL aliquot). The effluent solution was collected directly, without discarding the first
173 portion. Experiments were performed with and without prior conditioning of ImpAs. Without
174 conditioning, ImpAs was used dry. When conditioned, the background media was passed through
175 ImpAs several times and care was taken to ensure that air space in the top of the syringe did not re-
176 aerate and thus dry out the resin. Conditioning took 30-60 seconds. A single syringe was prepared for
177 each mass of ImpAs, and these three syringes reused for all synthetic groundwater experiments. A
178 control experiment was performed in duplicate, with 1 mg L⁻¹ As(III) (10 mM HEPES) replacing the

179 1 mg L⁻¹ As(V) influent sample and [As(III) (aq)] determined by ASV to verify that As(III) was
180 neither removed nor oxidised during ImpAs treatment.

181

182 2.3. Electrochemical speciation

183

184 As(III) and total As were determined by ASV using a portable PDV6000 Ultra potentiostat (Modern
185 Water – UK) powered by 4x1.5 V AA batteries and equipped with the associated reference electrode
186 (Ag/AgCl//KCl(3M)) and auxiliary platinum electrode. The working electrode was a 30 µm diameter,
187 5 mm long gold microwire. Operating parameters are summarised in SI Table S2. Detection was made
188 by Linear Scan ASV (LSASV), scanning from -400 to +700 mV with a 4 or 8 V s⁻¹ scan rate and 1.6
189 mV step. The solution was stirred during the deposition step and then held for 10 seconds at -400 mV
190 before the stripping scan took place. All measurements were made in the presence of dissolved
191 oxygen.

192 As(III) was determined after addition of 10 mM HCl and 10-20 µM hydrazine to prevent oxidation
193 (Salaün et al., 2007) (Salaün et al., 2012), using a deposition potential of -0.7 V for 15s (SI Figure S2).

194 Total As was determined in 0.1 M HCl using a deposition potential of -1.3 V for 15s. In these
195 conditions of acidity and pH, immediate oxidation of As(III) to As(V) by the oxidant produced at the
196 auxiliary electrode ensures that all arsenic in the voltammetric is present as As(V) (Salaün et al.,
197 2007). The blank electrolyte prior to addition of the sample was used for background subtraction. The
198 As stripping peak was located at ca. +180 mV in 0.01M HCl and ca. +230 mV in 0.1M HCl. Peak
199 height was used for quantification. As(III) and total As were determined by the method of standard
200 additions, with a minimum of 2 additions of As(III) in each case (Gibbon-Walsh et al., 2011).

201 The working electrode was cleaned in 0.5 M H₂SO₄ by imposing a potential of -2.5 V for 30s
202 followed by a CV scan between -200 and +1500 mV at a scan rate of 1000 mV s⁻¹ (Salaün et al.,
203 2012) (SI Figure S3). Electrodes were cleaned at the start of each day, and sometimes between

204 measurements if electrode sensitivity appeared diminished (i.e. after analysis of very high iron content
205 synthetic waters).

206

207 2.4. Determination of photooxidation kinetics

208

209 Suspensions of 0.1 g L⁻¹ Degussa P25 (TiO₂) and 10 mg L⁻¹ As(III) (133 μM) were prepared in 10
210 mM HEPES (total volume 100 mL) and stirred overnight in the dark to achieve equilibrium
211 adsorption. Beginning at t=0, suspensions were irradiated by UV light (λ=368 nm) with a power of
212 13.7 mW cm⁻². Aliquots were taken at regular intervals, with TiO₂ removed using a 0.45 μm nylon
213 syringe filter (VWR 28145-489). Each aliquot was (a) analysed for total As by ICP-MS, (b) diluted
214 from 1 mL to 10mL with Milli-Q water and passed three times through a conditioned ImpAs syringe,
215 then analysed by ICP-MS for 'As(III)', and (c) speciated by ASV. A single ImpAs syringe was used
216 for all experiments. For each time series the value of k and its uncertainty were calculated as the slope
217 of the linear regression and its standard error. Experiments were performed in duplicate to give a final
218 average, and the final uncertainty determined as the difference between the maximum and average
219 values of k.

220

221 2.5. Sampling at field sites and arsenic spike experiment

222

223 Groundwater was collected from a natural spring sourced from granite bedrock, and surface water
224 from a stream feeding into the River Kennal, Cornwall, UK. Bottled mineral water from a French
225 volcanic spring was purchased. These waters were filtered using a 0.45 μm nylon membrane. West
226 Bengal groundwaters were sampled - well 3 (deep well, 400 feet), well 4 (shallow well) and STN13
227 (shallow well, 70 feet) in the village of Sahispur (23°04'16.9"N 88°36'33.4"E), and STN5 in the
228 village of Chakudanga (480 feet, 23°04'57.7"N 88°36'09.0"E), both in the Nadia district of West

229 Bengal, India. The sampling procedure was based on that of Gibbon-Walsh et al., 2011. Tube wells
230 were pumped continuously for 6-15 minutes before collecting samples in opaque plastic bottles to
231 prevent photooxidation of As(III). Samples were immediately spiked with either HCl (to pH 2) or 10
232 mM EDTA to prevent iron precipitation. Samples were filtered to remove solid particulates and
233 bacteria. Acidified samples were pH neutralised by addition of NaOH before conducting the arsenic
234 spiking experiment as ImpAs performs poorly under acidic conditions, owing to protonation of the
235 pyridine groups ($pK_a = 5.2$) disrupting the ImpAs metal-organic receptor (Moffat et al., 2014).

236 Natural samples were subdivided and spiked with arsenic stock solutions to give a wide range of
237 As(III):As(V) ratios. The first half of each subsample was collected without further treatment (for
238 determination of As(III) and total As by ASV, and total As by ICP-MS), whilst the second half was
239 passed through a syringe loaded with 1g ImpAs for determination of “As(III)”, with the results
240 labelled as *ImpAs/ICP-MS*. Using ASV, As(III) was determined within 24 hours of sampling and total
241 As within 48 hours. Samples were diluted at least 5 times with Milli-Q water during measurement by
242 ASV to minimise matrix effects and the most concentrated samples were diluted 60-100 times to
243 ensure analysis was made within the linear range of the method. Three ImpAs syringes were used to
244 treat all natural media, separated into (i) oxic waters, (ii) anoxic samples without EDTA, and (iii)
245 anoxic samples with EDTA. Absorbance at 254 nm was determined using a Jenway-7315 UV-vis
246 spectrophotometer with Milli-Q water used for background subtraction.

247

248 2.6. Determination of total As by ICP-MS

249

250 Samples were acidified to 2% (v/v) HNO_3 and analysed using the Agilent 7900 quadrupole ICP-MS.
251 Calibration curves of at least 5 points were obtained both with a single element arsenic standard
252 (Fluka, $1000 \pm 4 \text{ mg L}^{-1}$, traceCERT), and a multi-element standard (Inorganic Ventures, $10 \mu\text{g mL}^{-1}$,
253 NIST-traceable standard). Measurements were also normalised to internal standards (Yttrium) and the

254 uncertainty of each measurement was determined as the root-mean-square deviation (RSD) of each
255 measurement. The detection limit for arsenic was $0.41 \mu\text{g L}^{-1}$ (5.5 nM).

256 3. Results and Discussion

257

258 3.1. Demonstrating speciation by ImpAs chemisorbent

259

260 3.1.1. Speciation under equilibrium conditions

261

262 The first objective of this work was to investigate whether ImpAs can separate As(III) and As(V)
263 from synthetic waters. Batch reactions under equilibrium conditions were used to verify that As(V)-
264 selective ImpAs does not remove As(III) from solution, even after overnight shaking (in 10 mM
265 HEPES) (Figure 3a). Since separation of As(III) and As(V) fails when the sorbent is saturated, ImpAs
266 was kept in excess. The speciation, expressed as the fraction of As(III) in each sample, was
267 determined by Equation 1 and compared against the expected results (Figure 3b).

$$268 \quad \text{Speciation (\% As(III))} = 100 \times \frac{[\text{Total As (ImpAs treated)}]}{[\text{Total As (untreated)}]}$$

269

Equation 1

270 Using 1 g L⁻¹ ImpAs, the linear regression gave a slope of 0.952 ± 0.049 (within 5% of the expected
271 one-to-one calibration curve) with R² = 0.992. Equation 1 gave 2.9 ± 0.3% and 94.3 ± 2.2% As(III)
272 for mixing fractions 0% and 100% As(III) respectively. Similar results were obtained using 20 g L⁻¹
273 ImpAs: a calibration curve with a slope of 0.887 ± 0.022 with 3.5 and 92.6 % As(III) at mixing
274 fractions 0 and 100% As(III) respectively. These results indicate quantitative removal of As(V) and
275 separation from As(III) within ±7%.

276

277 3.1.2. Implementation as an easy-to-use, portable device

278

279 Having verified that ImpAs can speciate arsenic under equilibrium kinetics, ImpAs was loaded into
280 syringes to (i) provide a rapid and high throughput device for laboratory applications, and (ii) to
281 provide portability for on-site applications. Syringes were first tested for As(V) removal in 10 mM
282 HEPES with and without prior conditioning (i.e. resin wetting) (Figure 3c). We first aimed to identify
283 the appropriate amount of ImpAs for loading into 5 mL plastic syringes – balancing efficient removal
284 with low material consumption. Since As(V) sorption is a function of contact time between sorbent
285 and sample, syringes were prepared with 0.2 g (2.5 mm path length), 0.4 g (5 mm) and 1.0 g (12 mm)
286 ImpAs.

287 Without prior conditioning As(V) removal was limited and the 1 g ImpAs syringe only removed 57.9
288 $\pm 2.0\%$ As(V) in a single pass (Figure 3c). This is likely due to pore channelling between resin beads,
289 offering paths of least resistance through which influent contact with ImpAs is limited. However,
290 when ImpAs was conditioned first, it demonstrated very efficient removal of As(V) ($98.0 \pm 1.8\%$ with
291 a single pass, using 1g ImpAs syringe). When conditioned, increasing the mass of ImpAs also
292 increased As(V) removal due to increased contact (Figure 3c). In all cases ImpAs was unsaturated and
293 so incomplete As(V) removal was due to kinetic limitations rather than limited capacity. Cross-
294 contamination due to carry over was only identified for a single data point in Figure 3c. Control
295 experiments (N=2) demonstrated that As(III) was neither removed nor oxidised when passed through
296 the syringe (Figure 3c, grey diamonds).

297 Having established efficient As(V) removal using ImpAs syringes in HEPES buffer solutions, more
298 complex media were considered. Challenge Water (CW) is a synthetic groundwater used to test the
299 performance of arsenic removal units (The NSF Joint Committee on Drinking Water Treatment Units,
300 2016), whilst synthetic groundwaters SGW-1 and SGW-2 were formulated using published data on
301 West Bengal groundwater composition (McArthur et al., 2004) (Table 1). CW was high in sulphate,
302 which is known to interfere with anion exchange resins (Pakzadeh and Batista, 2011), whilst SGW-1
303 was high in phosphate, a known competitor of As(V) for ImpAs binding sites (Moffat et al., 2014).
304 ImpAs was effective in both CW and SGW-1 with 92.9 ± 1.1 (N = 2) and $96.9 \pm 0.7\%$ (N = 2) As(V)

305 removed in a single pass (Figure 3d), demonstrating that ImpAs-loaded syringes can speciate arsenic
306 in the presence of inorganic groundwater species.

307

308 3.2. Optimising electrochemical speciation for challenging waters

309

310 The objective of this section was to optimise our portable voltammetric system for arsenic speciation
311 in the challenging arsenic-contaminated waters of South Asia that often contain high levels of Fe(II)
312 and Mn(II) (Gupta and Singh, 2018). In West Bengal and Bangladesh, concentrations of 60 μM Fe(II)
313 (approximately 3.3 mg L^{-1}) and 20 μM Mn(II) (1.1 mg L^{-1}) are typical for arsenic contaminated
314 groundwaters (McArthur et al., 2004) (Hug et al., 2008) (Ayers et al., 2016). There are only few
315 papers dealing with the electrochemical speciation of arsenic in groundwaters of South Asia (Rasul et
316 al., 2002) (Paul et al., 2008) (Gibbon-Walsh et al., 2011) and they all differ in the voltammetric
317 methods and/or electrodes. Here, we tested the influence of high levels of Fe and Mn in synthetic
318 groundwater on the determination of As(III) and total As under acidic conditions, using fast linear
319 scan ASV (4-8 V s^{-1}).

320 Additions of high levels of Fe(II) generally resulted in a decrease of the As(III) signal (Figure 4a and
321 4b) with e.g. a loss of *ca.* 40% for addition of 120 μM Fe(II) (6.6 mg L^{-1}). Even at 300 μM Fe(II), the
322 arsenic peak is still well shaped and standard addition can still be achieved for reliable arsenic
323 determination. A peak at *ca.* +0.6 V whose intensity increased linearly with Fe additions is present on
324 the stripping scan, but its potential is significantly more positive than that of the arsenic. For the
325 determination of total As no significant decrease of the As peak was observed during addition of up to
326 300 μM Fe(II) (16.5 mg L^{-1}) (SI Figure S6). Similarly, addition of up to 100 μM Mn(II) (5.5 mg L^{-1})
327 affected neither the As(III) nor the As(V) peak (SI Figure S7).

328 Sensitivities and linear ranges were similar in synthetic groundwater (SGW-2) as in 10 mM HCl
329 (Figure 4c, 4d and SI Figure S9-10). Detection limits were higher in synthetic groundwater than in
330 blank HCl: for total As the LoD increased from 0.06 to 0.4 $\mu\text{g L}^{-1}$ (0.8 to 5 nM) and for As(III) the

331 LoD increased from 0.1 to 0.3 $\mu\text{g L}^{-1}$ (1 to 4 nM) (SI Table S2). Within the linear range, the average
332 uncertainty of measurement via the method of standard additions was $\pm 6\%$ (propagated from the
333 standard error of the slope in the linear regression). In all cases detection limits were $< 1 \mu\text{g L}^{-1}$
334 indicating that arsenic contaminated waters can be speciated. This showed that the inorganic ions
335 commonly encountered in South Asia, including natural levels of Fe(II) and Mn(II), do not
336 significantly impact arsenic analysis by the method of standard additions under this voltammetric
337 procedure.

338

339 3.3. Application 1: Determination of photooxidation rate constants in 340 synthetic solutions

341

342 Photooxidation of As(III) using TiO_2 as a catalyst was investigated as an example application with the
343 aim to highlight the advantages of using ImpAs as a simple and fast speciation tool. Remediation of
344 As(III) contaminated waters is challenging since this species adsorbs poorly to sorbent media such as
345 goethite (Qi and Pichler, 2017) and is hard to remove with membrane technology (Nicomel et al.,
346 2015). A number of research groups have thus investigated the use of photocatalysts such as TiO_2 to
347 oxidise As(III) to As(V) as a pre-treatment to reduce toxicity and improve adsorption (Fontana et al.,
348 2018).

349 In this work the oxidation of As(III) to As(V) was successfully monitored using the ImpAs/ICP-MS
350 procedure, with very similar results to those obtained by ASV (Figure 5a). The reaction was modelled
351 following pseudo-first order kinetics, with the rate equation

$$352 \quad -\frac{d[\text{As(III)}]}{dt} = k[\text{As(III)}]$$

353

Equation 2

354 where $d[\text{As(III)}]/dt$ is the rate of change in the concentration of As(III) with respect to time, and k is
355 the pseudo-first order rate constant (min^{-1}). The integrated solution to this equation is rearranged into
356 the linear form

$$357 \quad \ln\left(\frac{[\text{As(III)}]_t}{[\text{As(III)}]_0}\right) = -kt$$

358 *Equation 3*

359 where $[\text{As(III)}]_t$ and $[\text{As(III)}]_0$ are concentrations of As(III) at time 't' and 0 respectively. Plotting
360 Equation 3 in the linear form $y=mx+c$ indicated pseudo-first order kinetics, with $R^2=0.937\pm 0.006$
361 using ASV measurements, and $R^2=0.981$ using ImpAs/ICP-MS (Figure 5b). For each experiment, the
362 values of k as determined by the two different analytical procedures were within error of one another
363 (Figure 5c). The final average of the two repeats gave $k=0.0710\pm 0.0019 \text{ min}^{-1}$ using ASV, and
364 $k=0.0725\pm 0.0061 \text{ min}^{-1}$ using ImpAs/ICP-MS (Figure 5c). Again, the values of k obtained by the two
365 analytical methods were within error of one another, indicating that ImpAs successfully captured the
366 same reaction kinetics as ASV.

367

368 3.4. Application 2: Speciation of natural waters

369

370 3.4.1. Spike recovery experiment

371

372 These experiments aimed to demonstrate that ImpAs selectively removes As(V) from natural samples
373 thus enabling speciation to be determined by subsequent ICP-MS analysis. Commercial bottled
374 mineral water (France) and groundwater from a spring and surface water from a stream (both from
375 Cornwall, UK) were chosen as representative oxic waters, and four groundwater sources from tube
376 wells in West Bengal were sampled as anoxic reducing waters with high concentrations of dissolved
377 iron. After size filtration, clear groundwater samples from West Bengal needed immediate

378 acidification (to pH 2) with HCl or spiking with 10mM EDTA to prevent formation of yellow-brown
379 iron precipitates within a matter of minutes.

380 Results for three of these natural waters are shown in Figure 5. For untreated samples ICP-MS
381 showed that 100% of the As(V) spike added was recovered by total As determination, whilst ImpAs
382 treatment removed all of the As(V) spike added. Both determination of total As of untreated samples
383 by ICP-MS and As(III) determined by ImpAs/ICP-MS were consistent with the ASV determinations
384 of total As and As(III) respectively. Similar results were obtained for waters from both oxic and
385 anoxic environments. No colour change in the solution was observed during the course of the
386 experiment for these samples (though the resin became coated with a dark precipitate after treating
387 large volumes of the West Bengal groundwater samples – SI Figure S13).

388

389 3.4.2. Limitations

390

391 Samples preserved by acidification must be pH neutralised before treatment with ImpAs, to prevent
392 disruption of the metal-organic receptor. Unlike the other West Bengal groundwaters, for STN5 rapid
393 iron precipitation was visually observed both (i) immediately after sampling the water before
394 acidification and (ii) within minutes of pH neutralisation. This sample showed a very high iron
395 concentration (13 mg L^{-1} versus $\leq 5.0 \text{ mg L}^{-1}$ for all other samples), and 99% of all arsenic present
396 had been oxidised during storage. During the course of the STN5 spike experiment observed total As
397 failed to match the quantity of As(V) spike added (Figure 7a) owing to loss of arsenic during iron
398 precipitation. The concentration of As(III) was more stable, suggesting that losses of arsenic were
399 primarily due to adsorption and co-precipitation of As(V), in line with previous findings (Gibbon-
400 Walsh et al., 2011).

401 EDTA can be used instead of acidification to prevent iron precipitation. Whilst addition of EDTA
402 prevented loss of iron both in standing samples and during ImpAs treatment (SI Figure 14), EDTA

403 presented a new problem. By complexing Zn(II) present in the ImpAs resin, EDTA limited the ability
404 of ImpAs to retain As(V) (Figure 7b). A discussion of why ImpAs performance is limited in the
405 presence of EDTA, but not in the presence of weaker chelating agents such as dissolved organic
406 carbon (e.g. humic acids) is given in the Supplementary Information (Figures S15-S17).

407

408 3.5. Comparison of ImpAs/ICP-MS with ASV for rapid, reliable and 409 portable speciation of arsenic

410

411 Good agreement was found in the speciation of synthetic waters using ImpAs/ICP-MS and ASV,
412 generally within error of one another (Figure 8a). The cross-calibration curve gave $R^2 = 0.967$. In the
413 first case study, the difference between photooxidation rate constants determined by ASV and
414 ImpAs/ICP-MS was insignificant - the difference of 2.0% was within error bounds, suggesting good
415 accuracy. The precision of rate constants determined in duplicate by ImpAs/ICP-MS was slightly
416 lower than those obtained by ASV ($\pm 8.4\%$ versus $\pm 2.7\%$).

417 For natural waters the speciation determined by ImpAs/ICP-MS and ASV also agreed, but with lower
418 goodness of fit ($R^2 = 0.902$) (Figure 8b). Importantly, there was no significant difference in the cross-
419 calibration of ImpAs/ICP-MS and ASV speciation between oxic and anoxic water samples.

420 ImpAs/ICP-MS was much more time effective than ASV, requiring 30-60 seconds to condition the
421 resin and one minute to filter 5 mL of sample, plus 3 minutes analysis time by ICP-MS. ASV
422 meanwhile required approximately 20 minutes experimental work to measure one sample in duplicate
423 (35 seconds per scan, 15 scans to measure background, sample and 2 additions in triplicate). This
424 highlights how chemisorbents such as ImpAs might enable high throughput applications in the
425 laboratory. With a maximum As(V) capacity of 16 mg g^{-1} , a single syringe with 1.0 g ImpAs can
426 theoretically treat a maximum 16 L water contaminated with $1000 \mu\text{g L}^{-1}$ As(V). Owing to the high
427 sorption capacity of ImpAs, a single syringe can be reused to treat many samples - just three syringes
428 were used to treat nearly all samples in this work – separated into (i) all synthetic media samples (at

429 1.0 grams loading), (ii) all natural oxic waters, and (iii) all natural de-acidified West Bengal
430 groundwaters, with little trace of cross-contamination between samples.

431 Speciation using handheld syringes loaded with ImpAs is highly portable – especially as a single
432 syringe can treat many samples and unlike ASV, no hardware, batteries or additional reagents are
433 required. If coupled with a fast laboratory detection method (e.g. ICP-MS), high sample throughput
434 can be achieved. In contrast, although ASV has lower sample throughput and requires more manual
435 operation, ASV provides an immediate answer for on-site analysis and for near real-time monitoring
436 of oxidation experiments. ASV is more appropriate for natural samples with high iron levels ($> 5 \text{ mg}$
437 L^{-1}) where acidification or EDTA can be used to avoid precipitation.

438

439 4. Conclusions

440

441 This study aimed to develop the application of the As(V)-selective chemisorbent ImpAs as a fast,
442 portable and cost-effective method to separate As(III) and As(V) from aqueous solutions (synthetic
443 and natural) for applications in the design of arsenic treatment plants and monitoring campaigns in the
444 field.

445 The key finding is that ImpAs can speciate inorganic arsenic in synthetic and natural waters at pH 7;
446 this was true for both oxic waters and anoxic waters with $\leq 5 \text{ mg L}^{-1}$ dissolved iron. We have shown
447 here that chemisorbent materials offer an alternative and attractive approach towards the speciation of
448 arsenic for on-site and laboratory applications, being simple, portable, quick and easy-to-use with
449 little specialist knowledge required. In unstable waters where iron precipitation occurs within a few
450 minutes, ASV is presently recommended as a portable and relatively low-cost solution for on-site
451 arsenic speciation. Further work should consider the benefits and trade-offs between chemisorbents
452 and other portable techniques for speciation, particularly other solid phase extraction techniques such
453 as strong anion exchange resins.

454

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456

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461 6. References

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- 598

599 7. Tables

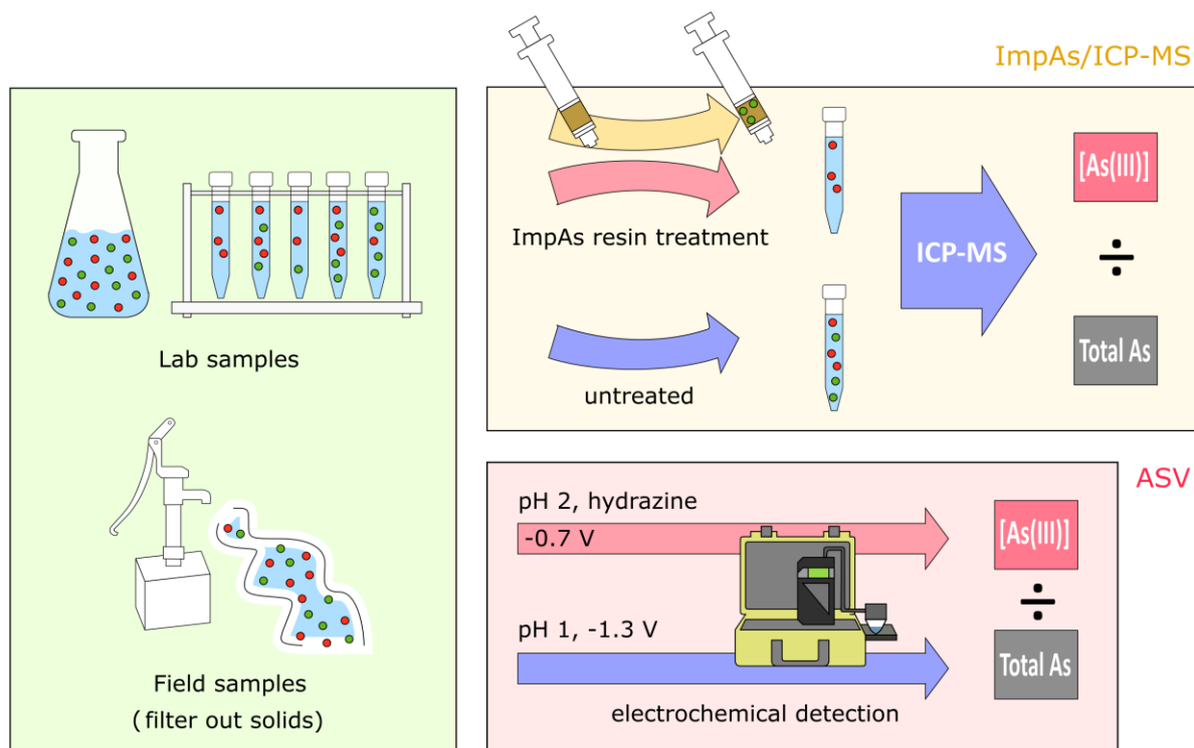
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Ion	Concentration (mg L ⁻¹)									
	Synthetic media			Natural media						
	SGW-1 (ImpAs experim ents)	SGW-2 (ASV experim ents)	CW	bottled mineral water	Cornish ground water	Cornish surface water	well 3 (West Bengal deep well)	well 4 (West Bengal shallow well)	STN13 (West Bengal shallow well)	STN5 (West Bengal deep well)
HCO ₃ ⁻	590	650	180	-	-	-	-	-	-	-
Ca ²⁺	34	1.5	47	2.5	2.5	2.1	16	14	18	19
Na ⁺	250	250	88	9.8	9.8	11	990	860	139	68
Mg ²⁺	37	39	13	9.0	3.9	3.1	29	19	17	16
Cl ⁻	160	160	82	-	-	-	-	-	-	-
NH ₄ ⁺	-	27	-	-	-	-	-	-	-	-
H ₄ SiO ₄	63	-	31	-	-	-	-	-	-	-
SO ₄ ²⁻	14	12	50	-	-	-	-	-	-	-
K ⁺	2.7	0.57	-	6.7	3.0	2.7	4.1	2.1	9.0	9.1
Fe ²⁺	-	0 or 3.4 added	-	<LoD	<LoD	<LoD	1.8	5.0	4.3	13
Mn ²⁺	0.97	-	-	0.0002	0.021	0.0002	0.13	0.14	0.3	0.3
HPO ₄ ²⁻	1.1	1.4	0.13	-	-	-	-	-	-	-
NO ₃ ⁻	0.62	4.5	8.8	-	-	-	-	-	-	-
Sr	-	-	-	0.057	0.060	0.047	0.49	0.27	0.37	0.35
Ba	-	-	-	<LoD	0.007	0.006	0.12	0.17	0.23	0.20
F ⁻	-	-	1	-	-	-	-	-	-	-
pH	7.09 ± 0.02	8.32 ± 0.02 ^b	7.42 ± 0.02	7.39 ± 0.02	6.52 ± 0.02	7.55 ± 0.02	7.3 ± 0.1	7.2 ± 0.1	8.2 ± 0.1	8.4 ± 0.1
absorbance at 254 nm (cm ⁻¹)	-	-	-	0.023	0.025	0.105	-	0.046	0.022	0.044

601 *Table 1: Composition of waters used in this study –. Synthetic groundwater compositions were determined from the mass of*602 *reagents used. The composition of natural waters was determined by ICP-MS using a multi-element standard. Blank cells*603 *indicate where a species was not added to solution or detected. '<LoD' refers to species that were under the detection limit.*604 *^a : SGW-1 and SGW-2 were both based on data from McArthur et al., 2004. SGW-1 was used for ImpAs experiments in*605 *London, whilst SGW-2 was used for electrochemistry experiments in Liverpool.*606 *^b : Since SGW-2 was used for optimisation of ASV conditions, this sample was always acidified to pH 1 or 2 during analysis.*607 *The uncertainty in pH is determined from the sensitivity of the two different pH meters used in the lab and in the field.*

608

609 8. Figures



610

611 *Figure 1: Concept schematic for arsenic speciation as performed by the two methods used in this study. As(III) is*
 612 *represented in red, and As(V) in green. Blue arrows denote determination of total As, whilst red arrows denote*
 613 *determination of As(III). Speciation is reported as the percentage of total As which is As(III).*

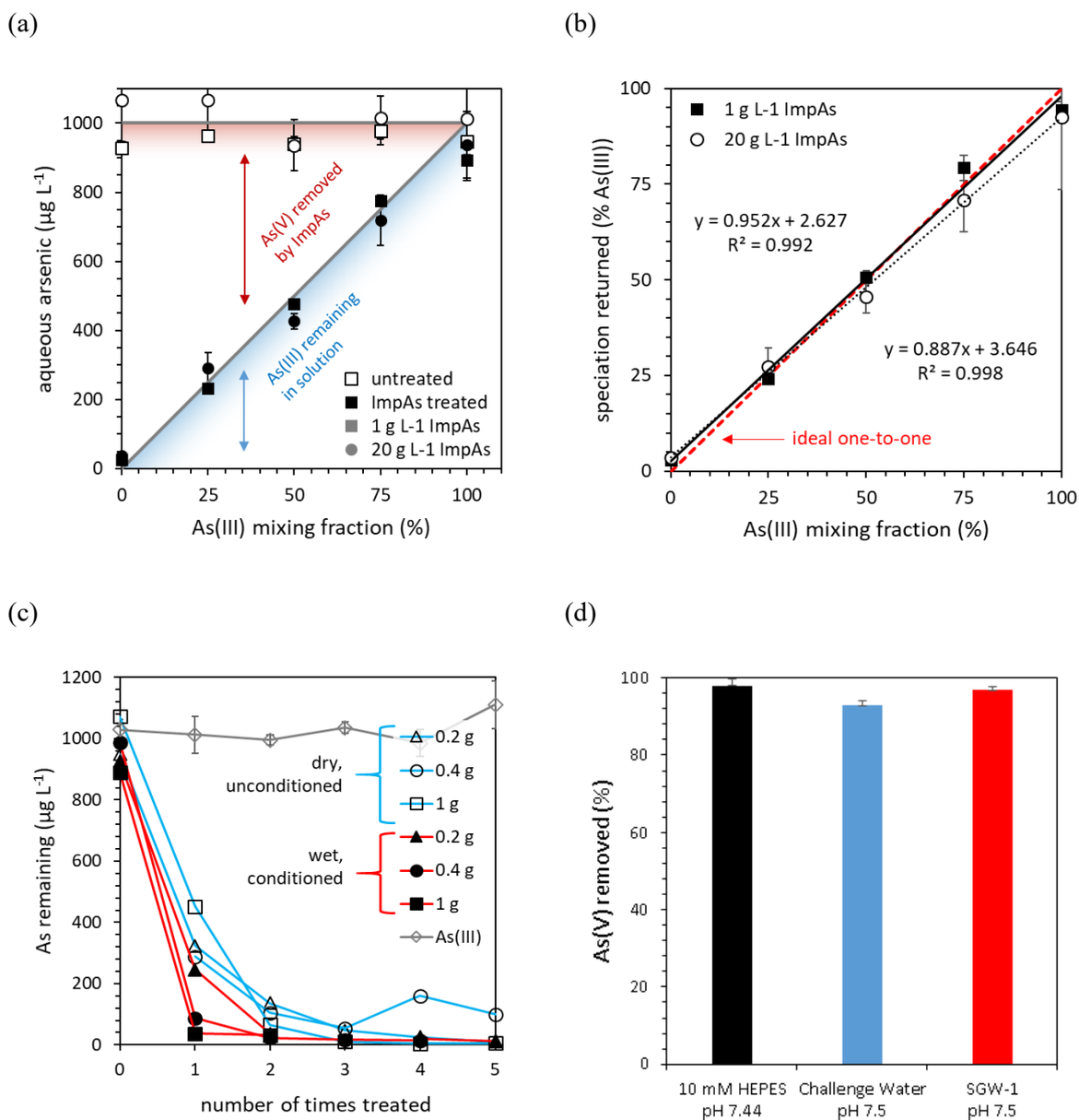
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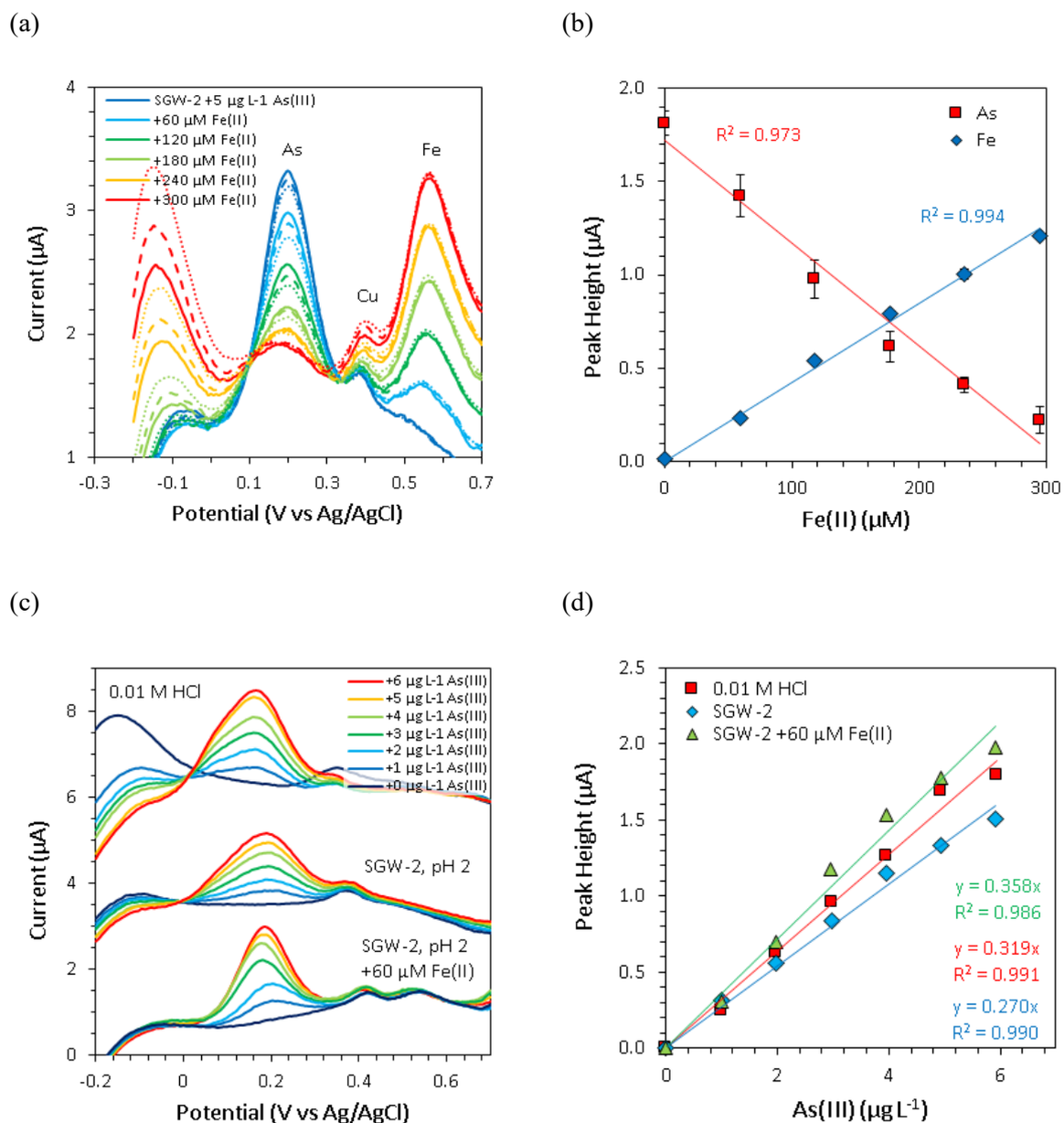
616 *Figure 2: Syringes loaded with 0.2, 0.4 and 1.0 gram ImpAs used in this work. Column path lengths were 2.5, 5 and 12 mm*
 617 *respectively.*

618



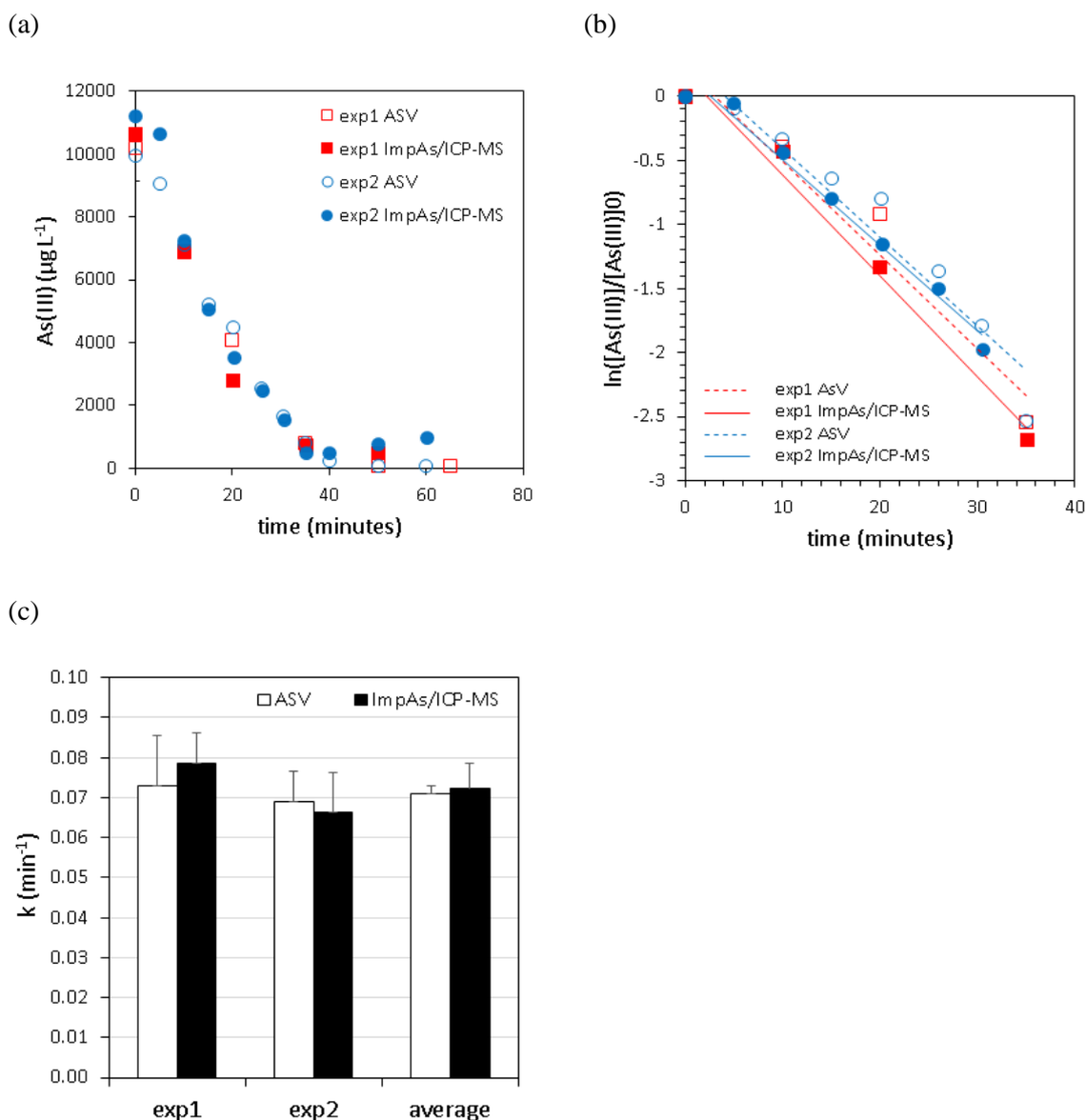
619 *Figure 3: ImpAs speciation method development. (a) The concentration of total As that remains in solution with and without*
 620 *addition of ImpAs as a function of As(III):As(V) mixing fraction, and (b) the resulting arsenic speciation calibration curves.*
 621 *(c) The removal of As(V) with sequential passes through syringes loaded with different masses of ImpAs. (d) As(V) removal*
 622 *efficiency in different media after a single pass through a syringe loaded with 1 g ImpAs (N=2).*

623



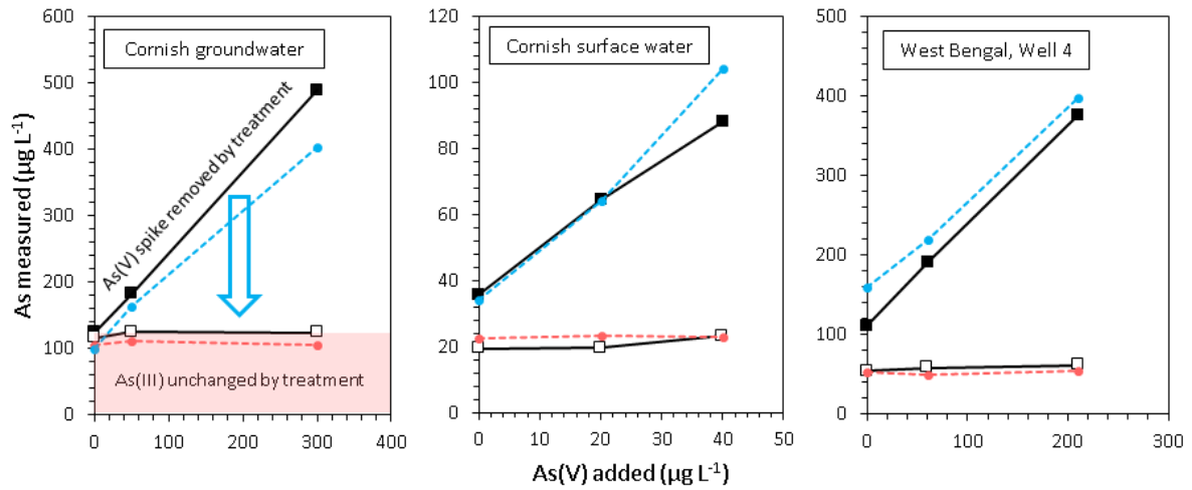
624 Figure 4: Electrochemical method development. (a) and (b) show the stability of As stripping peaks (5 $\mu\text{g L}^{-1}$ or 67 nM
 625 As(III) in synthetic groundwater with increasing additions of Fe(II). For each 60 μM addition of Fe(II), three scans were
 626 made (solid, dashed, and then dotted line) to check for peak stability. (c) and (d) show the linear range of As(III)
 627 determination by ASV in 0.01 M HCl and synthetic groundwater with and without Fe(II) as additions of 1 $\mu\text{g L}^{-1}$ As(III) (13
 628 nM) are made. Similar results for total As are shown in the Supplementary Information. LSASV conditions: 15s at -1.3 V
 629 (total As) or -0.7V (As(III)), 10s hold at -400 mV, stripping from -400mV to +700mV at 4 V s^{-1} . Detection was made at pH 2
 630 with 20 μM hydrazine.

631

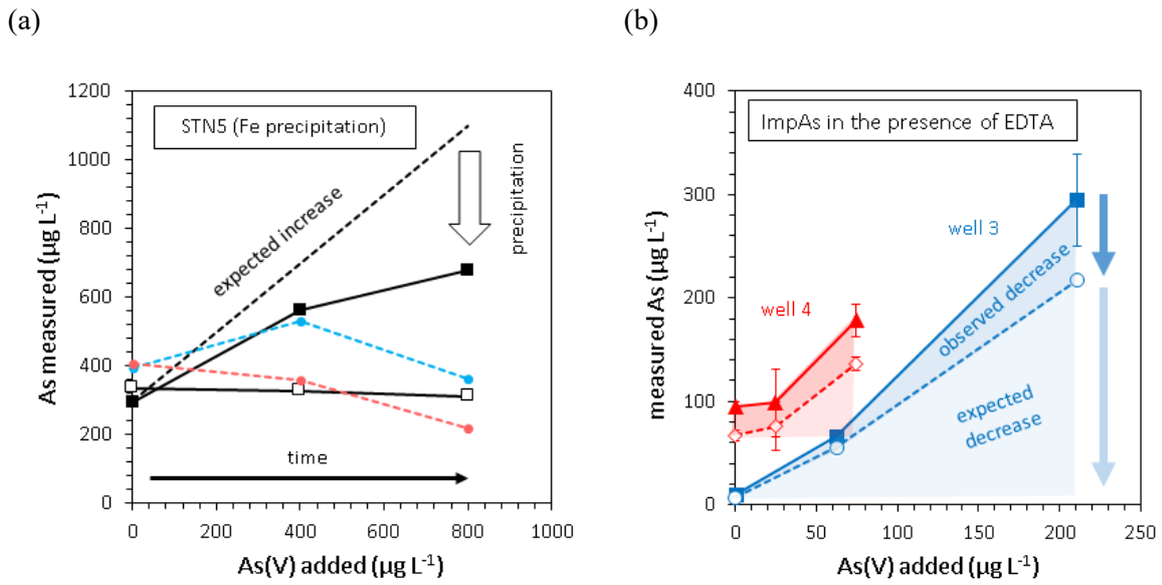


632 Figure 5: Case study 1 – kinetics for the photocatalytic oxidation of As(III) over TiO_2 , determined by ASV and ImpAs/ICP-
 633 MS. Conditions were 10 mg L^{-1} As(III), 0.1 g L^{-1} Degussa P25 TiO_2 powder, in 10 mM HEPES (pH 7.44). (a) Time series for
 634 photocatalytic oxidation of As(III) (two repeat experiments), (b) data fitted to a pseudo-first order kinetic model, and (c)
 635 comparison of rate constant, k , as determined by the two analytical methods. ASV measurements are shown as unfilled
 636 shapes, ImpAs/ICP-MS measurements are filled. Red squares indicate the first experimental repeat, and blue circles indicate
 637 the second experimental repeat. Solid lines and dashed lines indicate the linear best fit for ImpAs/ICP-MS and ASV results
 638 respectively.

639

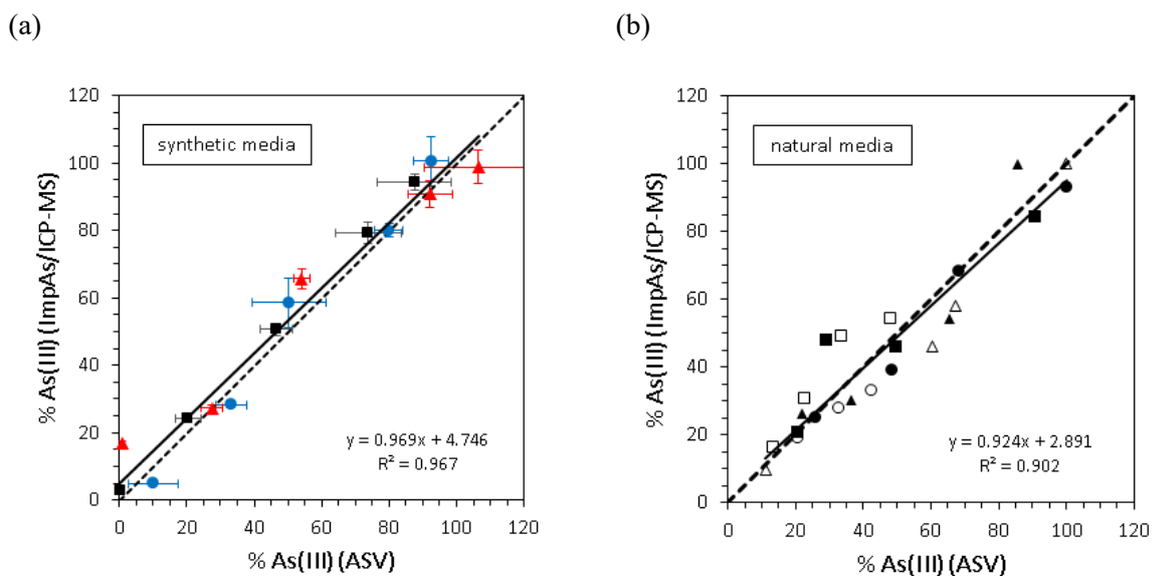


640 *Figure 6: Case study 2 – speciation in natural media. Samples were spiked with arsenic to obtain a mix of oxidation states*
 641 *and an ImpAs-loaded syringe used to remove As(V). By determining total As by ICP-MS before treatment with ImpAs (black*
 642 *filled squares) and after treatment (white squares) the speciation was determined. ASV determination is also given for*
 643 *comparison (dashed lines) with total As (blue) and As(III) (red).*



644 *Figure 7: Current limitations of speciation by chemisorbent. (a) Precipitation of iron during analysis of STN5 samples was*
 645 *associated with a loss of total As. (b) In the presence of 10 mM EDTA, ImpAs failed to remove all the As(V) spike, resulting*
 646 *in an overestimation of the proportion of As(III) present in the total As.*

647



648 *Figure 8: Comparison of arsenic speciation determined using the developed ImpAs/ICP-MS and ASV methods in (a)*
 649 *synthetic and (b) natural samples. (a) Cross-calibration of arsenic speciation in 10 mM HEPES (black squares), CW (blue*
 650 *circles) and SGW-1 (red triangles). Error bars indicate the propagated uncertainty of each measurement. (b) Cross-*
 651 *calibration of arsenic speciation in natural media. Data points correspond to bottled mineral water (black squares), Cornish*
 652 *groundwater (black circles) and surface water (black triangles), and West Bengal groundwater samples Well 4 (white*
 653 *squares), STN13 (white circles) and STN5 (white triangles). Dashed lines indicate the ideal one-to-one cross-calibration,*
 654 *and solid black lines indicate the linear best fit.*

655