Accumulation and effects of cadmium on sulphate-reducing bacterial biofilms

Chris White and Geoffrey M. Gadd

Biofilms comprising a pure and a mixed culture of sulphate-reducing bacteria (SRB) were grown in continuous culture. When exposed to 20 or 200 μM Cd, both cultures accumulated Cd but the mixed culture accumulated more and continued to accumulate Cd during the experiment, whereas accumulation by the pure cultures ceased after 4-6 d. Unlike the pure culture, the mixed culture also accumulated both protein and carbohydrate throughout the experiment proportionally to Cd which showed that accumulation required the production of biofilm material. Electron microscopy showed the presence of polysaccharide and particulates in both pure and mixed cultures, irrespective of the presence of Cd. However, energy-dispersive X-ray analysis (EDXA) showed that accumulation of Cd in the form of CdS occurred in biofilms exposed to Cd while back-scattered electron imaging of sections indicated that the accumulation of Cd was localized in a superficial layer of the biofilm. The mechanism of uptake, therefore, appeared to be entrapment and/or precipitation of CdS at the biofilm surface. The relatively low Cd uptake by the pure culture biofilm was attributed to its less efficient growth and polysaccharide production. These results indicate that mixed SRB cultures are more effective than pure cultures for metal removal and underlines significant differences between the biology of pure and mixed cultures.

Keywords: Sulphate-reducing bacteria, biofilms, cadmium, toxic metal accumulation

INTRODUCTION

Sulphate-reducing bacteria (SRB) biofilms are ubiquitous in anaerobic environments as well as having a significant economic impact through biocorrosion of metal structures (Little et al., 1990; Hamilton, 1994; Lee et al., 1995). Biofilms are now regarded as a dominant state of growth in many habitats for bacteria, including SRB, due to the favourable conditions provided by this growth habit (Costerton et al., 1994, 1995) which include protection against stresses such as dehydration or chemical toxicity (Anwar et al., 1992; Yu & McFeters, 1994; Costerton et al., 1995). In addition, nutrients may be concentrated at surfaces by sorption (Costerton et al., 1995) and the close proximity of organisms with complementary metabolic capacities, e.g. for carbon substrates, may enhance consortium growth and enable metabolic processes to modify parameters such as pH in a manner that is favourable to the organisms (Wimpenny et al., 1993; Costerton et al., 1994).

There are a number of mechanisms by which metals potentially interact with SRB biofilms, although little or no direct experimental work has been carried out. Precipitation of metal sulphides is significant in free-living SRB (White & Gadd, 1996a, b, 1997). In addition, other significant interactions may include biosorption at cell surfaces (Gadd, 1992a, b; Gadd & White, 1993) and other biofilm components such as extracellular polymeric substances (EPS). EPS, produced during both suspended and biofilm growth of SRB, is comprised of a mixture of polysaccharides, mucopolysaccharides and proteins with the detailed composition varying with species and growth conditions (Zinkevich et al., 1996). It has been shown to bind significant amounts of potentially toxic metals (Beech & Cheung, 1993) and has been reported to protect SRB against toxicity during biocorrosion of copper alloys (Videla, 1994). Another potentially significant role for biofilm EPS is sorption and entrapment of fine particulates which could include...
Precipitation by biogenic sulphide is an efficient means of remediating toxic metal-polluted waters (Hammack & Edenborn, 1992; Barnes et al., 1994; White & Gadd, 1996a, b; 1997; White et al., 1997; Gadd, 1996) but processes currently in operation are of low intensity and require large working volumes (Gadd & White, 1993). Further development of bioprecipitation would benefit from reduction of the reactor volume, which could be achieved by the use of biofilm reactors. The current study was therefore undertaken to examine both the mechanisms involved in Cd uptake by sulphate-reducing biofilms and the effects of exposure to dissolved Cd on biofilm growth. Two biofilms were investigated which comprised a pure and a mixed culture, thus providing a direct comparison between the growth of pure and mixed culture biofilms, their interactions with Cd and potential efficacy for bioremediation of liquid effluents.

**METHODS**

**Organisms and culture.** The mixed sulphate-reducing culture was initially selected from natural sediment samples by chemostat culture (White & Gadd, 1996a). This culture contained fermentative organisms in addition to the sulphate-reducing component when grown on sugars (White & Gadd, 1996b) but these were not evident when lactate was used as substrate and methanogenesis was not detected on any substrate (White & Gadd, 1996a, b). The pure culture was isolated from this culture by inoculating 20 ml anaerobic roll-tube cultures of Postgates B solid medium (Postgate, 1984) with 0.1 ml of a 10⁸ dilution of chemostat culture and incubating at 20 °C for 96 h. Isolated colonies, identified by their black coloration, were picked and transferred to anaerobic liquid batch culture in 10 ml sterile SL10 medium (Widdel & Pfennig, 1981, 1982) in screw-top tubes under N₂. SL10 medium comprised a salts solution (g per 950 ml): Na₂SO₄, 4.0; KH₂PO₄, 0.2; NH₄Cl, 0.25; NaCl, 1.0; MgCl₂, 6H₂O, 0.4; KCl, 0.5; CaCl₂, 2H₂O, 0.15; FeSO₄, 0.02 which was sterilized by autoclaving (121 °C, 20 min) in 950 ml batches, cooled under N₂ and made up to 1 litre by means of the following additions. The two trace element solutions were sterilized separately by autoclaving (121 °C, 15 min). The first comprised (mg l⁻¹) MnCl₂, 4H₂O (100.0), CoCl₂, 6H₂O (190.0), ZnSO₄, 7H₂O (144.0), H₂BO₃ (6.0), NiCl₂, 6H₂O (24.0), CuCl₂, 2H₂O (20.0) and Na₂MoO₄, 2H₂O (3.0). The second comprised (mg l⁻¹) Na₂SeO₃, 5H₂O (6.0) and Na₂WO₄, 2H₂O (8.0). The vitamin solution comprised (mg l⁻¹) 4-aminobenzoic acid (4), vitamin A (1.0), nicotinic acid (10.0), Ca-(+)-pantothenate (5.0), pyridoxine/HCl (15.0) and thiamin/HCl (10.0) and was sterilized by filtration through a sterile 0.45 µm pore size cellulose nitrate membrane filter (Millipore). One millilitre of each trace element and vitamin solution was added aseptically under N₂ to the salts after cooling. Thirty millilitres NaHCO₃ (840 g l⁻¹) and 150 ml Na₂S, 9H₂O (50 g l⁻¹), pre-sterilized by autoclaving in sealed tubes (121 °C, 15 min) were also added and the medium pH was adjusted to 6.5 by addition of sterile 5 M HCl. Liquid cultures were incubated for 72 h and used to inoculate further roll-tube cultures repeating the above process until a pure culture was obtained.

Both pure and mixed cultures were selected for attached growth using the following procedure. Thin glass coupons were made by cutting 25 mm square microscopic coverslips into four parallel pieces, each of which was placed in a 12 ml screw-top tube which was sealed and sterilized by autoclaving (121 °C, 15 min). Sterile, N₂-sparged SL10 medium (10 ml) was added aseptically to each tube which was then inoculated with 0.5 ml of either culture and incubated at 20 °C until a visible biofilm developed on the coverslip (7–14 d). The biofilm growth was then preferentially subcultured by removing the glass coupon and biofilm, snapping it in two pieces and using each half to inoculate a further culture on a similar coupon, thus inoculating two new cultures from each old culture. Once established, the biofilm cultures were maintained by the same procedure, subculturing at 14 d intervals.

**Experimental cultures.** Chemostat culture experiments were carried out in 11 glass Quickfit reaction vessels (Merck) equipped with a side-arm for outflow. The medium was stirred by means of a Flatspin solid-state magnetic stirrer (Merck) with a 25 mm cross-head stirrer bar operating at a constant 500 r.p.m. This maintained circulation with minimal turbulence. The vessel top-plate comprised a 6-mm-thick polypropylene disc with 20 concentrically arranged 10 mm diameter openings (Fig. 1a). The openings were closed with 9.5–110 mm diameter silicone rubber bungs which could be pierced to allow access for feeds or used to support 0.3-mm-thick 140×9 mm polystyrene coupons which were inserted...
into 5-mm-deep slots cut into the narrow end of the bungs and held in place using polymethacrylate adhesive (Fig. 1b). SL10 salts (900 ml) were sterilized in the vessel, cooled in a stream of N₂ and trace elements and vitamins then added. The assembled coupons and bungs were sterilized separately by immersion in 50% (v/v) ethanol for 24 h and inserted into the top plate, maintaining a stream of sterile N₂. Inoculum for 11 continuous cultures was developed, also in continuous culture, in a 10 ml syringe barrel which was sealed at the plunger end with a silicone rubber stopper containing a 0.3 × 12 × 60 mm polystyrene Plastikard (Slater) coupon. The Luer fitting of the syringe acted as an inlet port and a needle inserted through the stopper acted as an outlet. The system was sterilized by recirculating 200 ml 50% ethanol for 24 h after which the ethanol was displaced by pumping in SL10 medium, allowing 10 changes (100 ml). A glass coupon from a 14-d-old batch culture was then added and colonization and growth was allowed to proceed without medium addition for 48 h, after which medium flow was started (2 ml h⁻¹) and continuous culture maintained at 20 °C for 14–21 d. The biomass was then resuspended by transferring two coupons to 10 ml sterile, anaerobic SL10 medium in a 14 mm i.d., 12 ml test-tube, containing 1 cm³ 0.5 mm diameter sterile glass beads and filling the head-space with N₂ before sealing and vortexing (3 min, 200 r.p.m.). The beads were allowed to settle and the resuspended biomass was used as inoculum for the experimental culture. After 48 h batch-growth, continuous culture was started and the biofilm was allowed to develop for 7 d in SL10 medium at 20 °C and a dilution rate of 0.2 h⁻¹ to produce a standardized mature biofilm prior to the start of the experimental run. Experimental runs were carried out under identical conditions except that an appropriate volume of test-tube, containing 1 cm³ of N₂ and trace elements and vitamins then added. The required final Cd concentration and Na₂S was omitted from both control and metal-containing cultures to avoid premature precipitation of the added Cd. CdSO₄ stocks were made up in growth was allowed to proceed without medium addition for 24 h. The positions of individual coupons were numbered and they were removed in a predetermined random sequence. Approximately 5 mm was removed from the distal end of each coupon prior to four 10 mm long subsamples being removed from the remainder of the coupon for electron microscopy and chemical assays of the biofilm Cd, protein and carbohydrate contents, respectively. This ensured that the samples were taken from areas that were submerged under a minimum of 10 mm of medium during growth.

**Sampling.** Samples were taken by removing entire coupons. Duplicate samples were taken during growth in addition to quadruplicate final samples.

**Analysis.** Cd was assayed by atomic absorption spectrophotometry (AAS) following digestion in 6 M HNO₃ (White & Gadd, 1995, 1996a). Protein was extracted by vortexing the coupon in 10 or 20 ml 0.5 M NaOH with the addition of approximately 0.5 cm³ of 0.5 mm diameter glass beads followed by extraction for 30 min and assaying by the Bradford method (White & Gadd, 1996a). Carbohydrate was assayed using the anthrone method (Herbert et al., 1971) after vortexing with glass beads in the same way and removing a suitable volume of suspended biofilm. Sulphide was assayed polarographically (White & Gadd, 1996a).

**Electron microscopy.** Coupon samples for scanning electron microscopy and energy-dispersive X-ray analysis (EDXA) were fixed by immersion in 40% ethanol which strengthened the biofilm by partially dehydrating the extracellular polymeric material of the biofilm. They were then cut into 5 mm segments suitable for further preparation, dehydrated in successive 10% increments for ethanol concentration with three changes of absolute ethanol and critical-point-dried after which specimens for photography were sputter-coated with gold and those for EDXA analysis were carbon-coated. Dehydrated specimens were prepared for back-scattered electron imaging (BEI) by resin embedding (Wierzchos & Ascaso, 1993). The specimen was infiltrated with LR White resin for 48 h and cured for 24 h at 60 °C. A section through the specimen was then cut and the surface polished using a microtome. Microscopy, BEI and EDXA were carried out using a JEOL JSM-35 scanning electron microscope and EDXA spectra were analysed using an Apple computer.

![Fig. 2. Scanning electron micrographs of (a) pure and (b) mixed culture biofilms grown in unamended SL10 medium. A typical chain of cells including swollen, sporulating cells (s) is visible in (a). (b) shows a greater variety of cell-types present (although chains of cells are visible). Both micrographs show the accumulation of dehydrated extracellular polymers and granular material, although neither was grown in the presence of Cd. An area of this material is marked 'g' in (a) while it is generally distributed in (b). Some dead material, such as fragments of cell wall, is also present in both cultures. Bars, 10 μm.](image-url)
equipped with a Link Interface P1445 and software (Link Systems).

RESULTS AND DISCUSSION

Pure culture identification

The pure culture was a sulphate-reducing strain which utilized lactate and ethanol but grew poorly or not at all on acetate as carbon/energy source. The organism was pleomorphic, growing in suspended culture as a slightly curved rod during early growth but also producing swollen cells and endospores in batch culture. When grown as an attached (biofilm) culture, this strain typically produced chains of cells in which those furthest from the point of attachment were medium-length rods becoming sequentially elongated, swollen and drop-shaped, sporulating closer to the substratum (Fig. 2a). The chains were either attached basally producing streamers or attached throughout their length. The strain was provisionally identified on the basis of these characteristics as being a Desulfotomaculum species (Holt et al., 1994). Morphologically similar organisms formed a large proportion of cells visible in the mixed culture and appeared to perform a structural role in the biofilm (Fig. 2b).

Metal uptake

Biofilms of both pure and mixed cultures accumulated Cd when exposed to both 20 and 200 μM Cd. Both pure and mixed cultures took up significantly more Cd at 200 μM than at 20 μM but the amount taken up by the pure culture was significantly less than that by the mixed culture at both concentrations (Table 1). This appeared to result primarily from differences between the time course of metal uptake by pure and mixed culture biofilms in that, while the initial uptake rate of Cd by both cultures was similar, the pure culture apparently ceased to accumulate Cd after 4–6 d at either concentration while the mixed culture continued to accumulate the metal at both 20 and 200 μM Cd throughout the experiment (Fig. 3).

Effects of Cd on biofilm growth

Carbohydrate content, derived primarily from the EPS component of the biofilm, can be used as an estimate of biofilm growth (Pacepavicius et al., 1997). However, possible responses by the biofilm culture to Cd include inhibitory effects on growth and the secretion of extracellular metal-binding proteins (Fortin et al., 1994) or (muco)polysaccharides (Beech & Cheung, 1995) which could result in differential effects on the biofilm protein and polysaccharide production. Since protein is a more significant component of cells than EPS and the converse is true of polysaccharide, protein and polysaccharide content can together provide independent indices of cell growth and EPS development and a fuller picture of the effects of Cd on biofilm growth than either can provide singly. Both 20 and 200 μM Cd are potentially toxic to heterotrophic organisms in culture (Gadd, 1992a) and free ion concentrations of Zn and Ni in the same range have been shown to be toxic to sulphate-reducing bacteria (Poulson et al., 1997). However, there was no significant reduction in the protein content of either pure or mixed culture biofilms as a result of growth in the presence of either 20 or 200 μM Cd (Table 1, Fig. 4a, b). This suggested that the bioavailability of Cd to the biofilm was reduced by factors such as precipitation as CdS or binding to biofilm components. The protein content of the mixed culture was approximately tenfold greater than that of the pure culture, irrespective of the Cd concentration and, while Cd had no effect on the protein content of the pure culture at either concentration, the presence of 200 μM Cd in mixed cultures resulted in a small but significant increase in protein content over the control (Table 1). When the protein content of biofilms over the whole course of the experiment is considered, it is apparent that there was no overall accumulation or loss of protein.

Table 1. Cd, EPS and protein accumulation by mixed and pure sulphate-reducing biofilm cultures exposed to varying Cd concentrations

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Mixed culture</th>
<th>Pure culture</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cd (nmol cm⁻²)</td>
<td>Protein (mg cm⁻²)</td>
<td>EPS (μmol cm⁻²)</td>
</tr>
<tr>
<td>Control (0 μM Cd)</td>
<td>5.12</td>
<td>0.99</td>
</tr>
<tr>
<td>20 μM Cd</td>
<td>244.22</td>
<td>1.03</td>
</tr>
<tr>
<td>200 μM Cd</td>
<td>780.25</td>
<td>1.18</td>
</tr>
<tr>
<td>LSD (P &lt; 0.01)</td>
<td>92.08</td>
<td>0.08</td>
</tr>
</tbody>
</table>

Each value is the mean of eight determinations with the least significant difference (LSD) being calculated by variance analysis and comprising the residual SE of the whole data set for the three treatments multiplied by an appropriate t value for the required significance level at 21 degrees of freedom (Sokal & Rohlf, 1981). Mean values differing by more than this value can be considered as distinct at the significance level noted. Cd, protein and EPS content of pure and mixed culture biofilms subjected to 0 (control), 20 and 200 μM Cd were compared using Student’s t test (Sokal & Rohlf, 1981) and the cultures differed significantly in all parameters at P < 0.01.
Effects of Cd on sulphate-reducing biofilms

Fig. 3. Accumulation of Cd by (a) pure and (b) mixed culture biofilms grown in continuous culture with no added Cd (control) (C), 20 (●) and 200 µM Cd (□) over a period of 14 d following 7 d cultivation with no Cd present in all cases. Each point is the mean of at least two separate samples and the bars indicate the SEM.

by the pure culture (Fig. 4a), but that the mixed culture accumulated protein, the greatest accumulation occurring at 200 µM Cd (Fig. 4b). The polysaccharide and mucopolysaccharide components of SRB extracellular polymers are complex (Zinkevich et al., 1996) but, while not providing an absolute measurement of total EPS, the anthrone assay gives a useful index of carbohydrate content and EPS development (Kogel-Knabner, 1995). The carbohydrate content of the pure culture biofilm remained constant throughout the experiment (Fig. 5a) and there was also no significant difference between control cultures and those exposed to 20 or 200 µM Cd (Table 1). The control mixed culture, with no Cd present, contained approximately three times the amount of carbohydrate as the pure culture but also exhibited no net accumulation over time. In contrast, the mixed culture biofilms exposed to both 20 and 200 µM Cd both appeared to accumulate carbohydrate during the course of the experiment, with the greatest accumulation occurring at 200 µM Cd (Fig. 5b). Both the protein and carbohydrate content of mixed biofilms showed a significant correlation with the Cd content of the biofilms yielding $R^2$ values of 0.65 and 0.60, respectively ($P<0.01$ at 53 degrees of freedom in both cases) (Sokal & Rohlf, 1981). This accumulation of EPS and protein could have resulted from the production of metal-binding polymers or, alternatively, it may have been due to the interaction between the biofilm and solid amorphous CdS particles precipitated in the biofilm or its immediate environment. Numerous sites in bacterial

Fig. 4. Biomass growth in SRB biofilms grown in continuous culture, indicated by accumulation of protein, in (a) pure and (b) mixed culture biofilms grown with no added Cd (control) (C), 20 (●) and 200 µM Cd (□) over a period of 14 d following 7 d cultivation with no Cd present in all cases. Each point is the mean of at least two separate samples and the bars indicate the SEM.

Fig. 5. Development of extracellular polymeric material by SRB biofilms in continuous culture, indicated by accumulation of polysaccharide, in (a) pure and (b) mixed culture biofilms grown with no added Cd (control) (C), 20 (●) and 200 µM Cd (□) over a period of 14 d following 7 d cultivation with no Cd present in all cases. Each point is the mean of at least two separate samples and the bars indicate the SEM.
EPS are able to interact with such particulate material (Walker et al., 1989; Flemming et al., 1990; Flemming, 1995) and clay particles are incorporated into bacterial biofilms where they produce an apparent stimulation of biofilm growth by mechanical stabilization (Vieira & Melo, 1995).

Electron microscopy

Scanning electron micrographs of pure and mixed culture biofilms confirmed that the morphological diversity of the pure culture was less than that of the mixed culture. Because of pleomorphism, the pure culture contained a range of cell morphologies but some morphological types, such as coccoidal forms, were absent (Fig. 2a). Both in situ inspection with the naked eye and electron microscopy showed that pure culture biofilms also covered a lower proportion of the surface than mixed culture biofilms. The depth of pure culture biofilms was less than that of the mixed culture, although 2–4 mm streamers, formed from chains of cells, were only present in the pure culture. Both pure and mixed cultures contained dehydrated EPS with granular, presumably precipitated, material adhering to it and to cell surfaces. This was apparently present in similar amounts in both control cultures and those exposed to Cd [little of this material is visible in Fig. 2(a) as this micrograph was selected to illustrate the cell-types occurring in the culture]. However, EDXA analysis of the surface of the biofilm surface showed the presence of Cd in biofilms which had been grown in the presence of 20 or 200 μM Cd which was absent in controls (Fig. 6). Although EDXA analysis cannot quantify uptake, the peak representing sulphur was also greatly elevated in spectra from specimens exposed to Cd, indicating that a major component of the deposited Cd was CdS. This was especially striking in the spectrum obtained from the mixed culture exposed to 200 μM Cd where the CdS deposit was sufficiently extensive and continuous to leave only Cd and S peaks detectable and effectively mask the presence of the other elements detected in control and 20 μM EDXA spectra. This confirmed that precipitation and/or binding of CdS was the major mechanism of Cd accumulation by the biofilms although the existence of other mechanisms of lesser importance cannot be discounted. Metallic elements, such as Fe, Co, Ni and Zn, were supplied as micronutrients in the growth medium and therefore were potentially able to

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**Fig. 6.** EDXA spectra obtained from (a) pure and (b) mixed culture biofilms following 21 d continuous culture. The top spectrum in each set was obtained from control cultures grown in the absence of Cd and the middle and lower spectra were obtained from cultures grown in 20 and 200 μM Cd, respectively, for 14 d following 7 d development in the absence of Cd. Clear Cd peaks are visible in the lower spectra which are absent in the controls and the presence of Cd and S peaks alone can be seen in the bottom right (mixed culture, 200 μM Cd) spectrum. The Ti peak visible in the pure culture spectra derives from titanium white filler and was the only metallic element present in the coupon. The total emission count for each spectrum was $3.1 \times 10^5$–$3.5 \times 10^5$. 

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Effects of Cd on sulphate-reducing biofilms compete with Cd for sulphide. However, except for Fe (71.9 μM), these were largely present in concentrations less than 1 μM. Under the pH conditions occurring in the system, the solubility product constant for FeS (log $K_c = -18.43$) is higher than that of CdS (log $K_c = -28.3$) (Sillen, 1964) but would allow significant competition at the lower Cd concentration of 20 μM. However, the presence of a large excess (1-0.4-5 mM) of sulphide in both pure and mixed culture systems would allow precipitation of both metals, with the relative abundance governed by their concentrations and the formation constants of their sulphides. This would therefore result in a precipitate comprising almost entirely of CdS when Cd was present at 200 μM. Quantitative chemical analysis for metals other than Cd was not carried out. However, peaks representing Fe and less significant Cu and Zn peaks were visible in EDXA spectra of both pure and mixed biofilms from control and 20 μM Cd cultures and it can be assumed that this represented deposition of FeS. There was no distinct peak for Fe in either biofilm at 200 μM Cd. It is not feasible to quantify metals from EDXA spectra as they are affected by a number of factors, e.g. penetration of the target, which would be strongly affected by the presence of precipitates, but the spectra are consistent with the metal sulphide chemistry outlined above. However, the experimental system employed was not well suited to obtaining quantitative mass-balance and kinetic data and this is the subject of further experimental work in progress. BEI also showed an accumulation of electron-dense material in the biofilm exposed to 200 μM Cd which was not present in the control (Fig. 7a, b). This was accumulated in a superficial layer of the biofilm and the thickness of the control biofilm was similar to the lower layer of the exposed biofilm, which contained no Cd. Although the biofilms were dehydrated, and therefore shrunken to an unknown degree, this distribution suggested that the accumulation of Cd occurred in the biofilm surface, whether by precipitation or entrapment of particles, and co-accumulation of Cd(S) and biofilm material led to the greater thickness of the biofilm.

Conclusions

Both pure and mixed culture biofilms accumulated Cd by mechanisms which appeared to be primarily the result of entrapment or nucleation of insoluble sulphides. In comparing the interactions of the pure and mixed cultures with Cd, a number of differences between the responses of the two cultures emerged. The growth of neither culture was inhibited by Cd but, even in the absence of Cd, the pure culture showed significantly less growth than the mixed culture, containing approximately 13% of the protein and 36% of the EPS content of the mixed culture. The pure culture used was a preferentially lactate-utilizing organism while the mixed culture contained both lactate- and acetate-utilizing organisms which would allow a higher energy yield from the substrate. The possibility also exists that other complementary metabolic capacities occurred within the mixed culture. Similarly, the pure culture biofilm accumulated only 25–30% of the amount of Cd accumulated by the mixed culture. Since accumulation of Cd within the mixed biofilm was accompanied by proportional accumulation of both protein and EPS, which did not occur in the pure culture biofilm, we can suggest that Cd accumulation by the pure culture was limited by its lower capacity to produce Cd-binding extracellular components. Although it is generally believed that mixed cultures have several advantages over pure cultures for environmental biotechnology, there have been few studies in which a direct comparison has been made. This study shows that the mixed culture biofilm differed from the pure culture considerably in its growth responses to Cd, although this was composed of a major component species purified from the same mixed culture, and confirms that pure culture studies may be a poor guide to the behaviour of more complex mixed culture systems. However, while the long-term per-

Fig. 7. Back-scattered electron images obtained from polished sections of mixed culture SRB biofilms embedded in LR White resin following growth in continuous culture for (a) 21 d with no Cd added and (b) 7 d with no Cd followed by 14 d with 200 μM Cd. The layer of dark, electron-dense granules in (b) indicates the position of precipitated CdS in the biofilm. The dark band is the 'Plastikard' coupon which contained titanium white filler.
formance of the mixed culture was better than that of the pure culture, both were essentially similar in their direct interactions with Cd and biofilms of both cultures were capable of precipitating and immobilizing considerable quantities of Cd as entrapped CdS, demonstrating the potential of sulphate-reducing biofilms for bioremediation.

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