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Abstract A microcosm study was undertaken to examine the effects of dissolved cadmium at various concentrations (0, 10, and 100  $\mu\text{g}\cdot\text{L}^{-1}$ ) on biofilm accumulation and diatom assemblages. A natural biofilm sampled from the Riou-Mort River (Southwest France) was inoculated into three experimental systems, where biofilm settled on glass slides. Samples collected after 1, 2, 4, and 6 weeks of colonization were analyzed for metal accumulation (total metal content and intracellular metal content in the biofilm), biomass (as measured through dry weight and ash-free dry matter), and quantitative as well as qualitative analysis of diatom assemblages. There was a positive correlation between cadmium accumulation and dissolved cadmium concentrations and duration of exposure: a linear relationship was found between concentration factors (CFs) of growing biofilms and time (CFs/day = 0.25 and 0.38 under contaminations of 10 and 100  $\mu\text{gCd}\cdot\text{L}^{-1}$ , respectively). Biofilm settlement, more than photosynthetic activity, was affected by high cadmium concentrations: we observed for all stages of settlement a drastic and significant ( $p < 0.05$ ) reduction in biofilm biomass and in diatom densities in the highest cadmium contamination, compared to control and low cadmium concentration units.

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Keywords (separated by '-') Cadmium toxicity - Bioaccumulation - Biofilms - Diatom densities

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Footnote Information

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## 4 Cadmium Toxicity and Bioaccumulation in Freshwater Biofilms

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33 **Keywords** Cadmium toxicity · Bioaccumulation ·  
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### Introduction

Because of their specific ecological preferences (Lange-  
Bertalot 1979; Steinberg and Schiefele 1988; van Dam  
et al. 1994), benthic diatoms are commonly used to assess  
water quality and a great number of methods based on the  
use of diatoms have been proposed and are applied for the  
evaluation of eutrophication and organic pollution in rivers  
(a review of the major indices used in Europe is given in  
Prygiel et al. 1999). These methods are determined with  
consideration to nutrients and eutrophic conditions, how-  
ever, interactions between nutrients and toxicants often  
occur and are not evidenced through current indices. With  
the implementation of the European Water Framework  
Directive (2000/60/EC), there is a need to take into account  
priority substances such as heavy metals; studies are nec-  
essary for the improvement of diatom monitoring of these  
pollutions.

Aquatic primary producers from polluted sites are gen-  
erally considered to be passive absorbers of the toxicants  
present in waters, in which all their vital functions  
happen: nutrition, respiration, reproduction, excretion, etc.  
Although a number of studies have assessed the accumu-  
lation of cadmium and the biological effects in higher  
organisms in rivers of Southwest France (Andres et al.  
2000; Baudrimont et al. 2003, 2005), cadmium toxicity is  
often observable from the beginning of the food chain.  
Previous in situ (Ivorra et al. 1999, 2000; Gomez and  
Licursi 2003; Morin et al. 2007) and experimental studies  
(Interlandi 2002; Ivorra et al. 2002; Gold et al. 2003a, b)  
have underlined the impact of combined nutrients and  
metals on diatom community structures but interactions  
between contaminants are likely to bias the sensitivity of  
species to single factors (Lozano and Pratt 1994; Guasch  
et al. 1998). To avoid this bias, we propose to set up an

69 in vitro experiment to describe biofilm development under  
70 metal contamination. Microcosm studies stray from field  
71 conditions but allow the control of physicochemical  
72 parameters; in order to limit the differences with environ-  
73 mental conditions we worked on natural assemblages of  
74 diatom species. In this study, biofilm growth was quantified  
75 through dry weight, ash-free dry mass and chlorophyll *a*  
76 measurements in order to point out the effects of cadmium  
77 on structural as well as functional characteristics of the  
78 periphyton.

79 Therefore, these studies linked shifts in diatom consortia  
80 to water metal concentrations and did not reflect the real  
81 exposure in the organic matrix. Indeed, periphytic biofilms  
82 accumulate metals following three main mechanisms  
83 (Holding et al. 2003): (i) absorption in extracellular poly-  
84 meric substances (EPS), (ii) cell surface adsorption, and  
85 (iii) intracellular uptake. Here we propose to discriminate  
86 between metal adsorbed to abiotic or biotic materials and  
87 intracellular (nonexchangeable) metal by assaying the  
88 metal concentrations in the periphyton after no treatment or  
89 EDTA washing of periphyton samples (Behra et al. 2002).  
90 Moreover, many authors have reported metal-induced  
91 deformities of the frustule in polluted streams (McFarland  
92 et al. 1997; Shehata et al. 1999; Gomez and Licursi 2003;  
93 Nunes et al. 2003), yet correlations between exposure and  
94 occurrence of abnormal valves still lack experimental  
95 validation. The use of the frequency of abnormal frustules,  
96 if pointing to a metal stress, would be worthy for routine  
97 biomonitoring of metal contaminations.

98 The present microcosm study is aimed at characterizing  
99 biofilm growth (using measurements of dry weight, ash-  
100 free dry mass, and diatom density) under various metal  
101 levels and determining the accumulation kinetics of dis-  
102 solved cadmium in a natural freshwater biofilm, as well as  
103 metal toxicity to diatom assemblages and individuals.

## 104 Materials and Methods

### 105 Field Sampling

106 The Riou-Mort stream, a small tributary of the river Lot  
107 located in the industrial basin of Decazeville (Southwest  
108 France; 44°N / 2°E), exhibits polymetallic pollution from  
109 its confluence with the Riou-Viou, a stream carrying  
110 seepage from a former zinc factory, presenting high levels  
111 of dissolved cadmium (Say 1978). In the year 2000, Audry  
112 et al. (2004) measured concentrations of 16 mg·L<sup>-1</sup> dis-  
113 solved cadmium (average value).

114 Diatom communities were sampled from the field, at a  
115 site on the Riou-Mort located upstream of the contamina-  
116 tion source, in March 2006. Microbenthic biofilms were  
117 grown on 20 glass slides (300 cm<sup>2</sup> per slide) immersed in

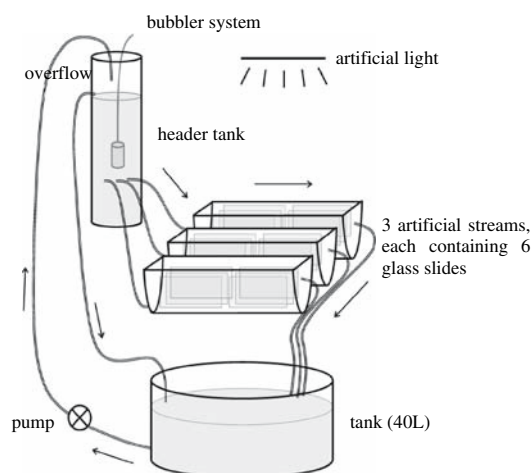
the stream in plastic racks for 5 weeks before collection,  
118 scraping, suspension in stream water, and transport to the  
119 laboratory (Morin et al. 2007).  
120

### Experimental Design

121  
122 In the laboratory, biofilm suspensions were separated into  
123 four aliquots; each aliquot was inoculated in the water  
124 column of three independent experimental units (EUs).  
125 Each system runs in a closed circuit and is composed of  
126 replicate artificial streams (60 cm long, 6-cm radius)  
127 equipped with six glass substrates, connected in parallel to  
128 a 40-L tank (Fig. 1). Using an external pump, water was  
129 circulated from the tank to a header tank providing a steady  
130 supply of water to the streams.

131 EUs were placed in an air-conditioned room, at a light  
132 intensity of approximately 70 μmol·s<sup>-1</sup>·m<sup>-2</sup> (10:14  
133 light:dark regime) and under continuous water movement  
134 at a velocity of approximately 0.4 cm·s<sup>-1</sup>. The reservoirs  
135 were filled with 40 L of modified Woods Hole culture  
136 medium (without EDTA and supplemented with silica)  
137 diluted fourfold (Gold et al. 2003a). The levels of nitrates  
138 and orthophosphates in the systems were typical nutrient  
139 concentrations found in the Riou-Mort river in 2004 and  
140 2005. Before the start of the experiment, the microcosms  
141 were equilibrated overnight with the culture media, to  
142 which the experimental concentrations of cadmium had  
143 been added.

144 During the course of the experiment (6 weeks), physical  
145 and chemical variables of the water (temperature, pH,  
146 electric conductivity, dissolved oxygen concentration and  
147 saturation) (WTW, Weilheim, Germany) were determined  
148 daily at the end of the light cycle. Nutrient (orthophos-  
149 phate, nitrate) concentrations were analyzed weekly at the



**Fig. 1** Schematic representation of one experimental unit. Arrows indicate direction of flow

150 laboratory from 1L water samples, according to French and  
151 international standards (NF T90-023 and NF EN ISO  
152 13395, respectively). Depending on the results of the  
153 analyses, culture medium was added as required to com-  
154 pensate for the decrease of nutrient concentrations due to  
155 algal uptake.

## 156 Water Column Contamination Protocol

157 Each individual experimental unit represented a different  
158 treatment: control (EU1), low contamination (EU2;  
159  $10 \mu\text{Cd}\cdot\text{L}^{-1}$ , in accordance with concentrations found in  
160 the polluted river Riou-Mort), high contamination (EU3;  
161  $100 \mu\text{Cd}\cdot\text{L}^{-1}$ ), which approximately corresponds to  
162 extreme values recorded in a tributary of the river Riou-  
163 Mort, the Riou-Viou, by the GEMA team, University  
164 Bordeaux 1 (e.g., more than  $60 \mu\text{Cd}\cdot\text{L}^{-1}$  in October  
165 2004).

166 The EUs were contaminated with a cadmium chloride  
167 solution ( $\text{CdCl}_2$ ; Merck, Darmstadt, Germany) to reach  
168 nominal test concentrations of 10 and  $100 \mu\text{g}\cdot\text{L}^{-1}$ . Cad-  
169 mium concentrations were measured (and, when necessary,  
170 corrected) daily during the first week and twice per week  
171 during the rest of the experiment. Water samples were  
172 filtered and acidified with  $\text{HNO}_3$ , to determine cadmium  
173 concentrations by atomic absorption spectrophotometry  
174 (Varian AA400—Zeeman correction, Victoria, Australia).  
175 The detection limit was  $0.1 \mu\text{gCd}\cdot\text{L}^{-1}$ . The accuracy of the  
176 analytical methods was checked periodically using two  
177 certified biological reference materials (Tort-2, lobster  
178 hepatopancreas; and Dolt-2, dogfish liver; NRCC-CNRC,  
179 Ottawa, Canada). Values were consistently within the  
180 certified ranges (data not shown).

## 181 Biofilm Sampling and Analyses

182 After a 1- and 2-week colonization, two glass slides were  
183 removed at random from each artificial stream; one slide  
184 was sufficient for biofilm collection at weeks 4 and 6. The  
185 streams were then reset with new glass slides, to preserve  
186 identical flow conditions.

187 Both faces of the glass substrates were carefully scraped  
188 with a cutter blade and the biofilm was suspended in a  
189 standard volume of 100 mL to form a single sample per  
190 stream (i.e., three replicate periphyton samples per exper-  
191 imental unit) per sampling date and per treatment.

192 Each suspended biofilm sample was separated into  
193 aliquots assigned to various analyses. Twenty milliliters  
194 ~~was~~ used for particulate matter analysis: biofilm dry  
195 weights (DW) were determined following the European  
196 standard NF EN 872; after drying and weighing, samples

were ashed at  $500^\circ\text{C}$  for 1 h in a muffle furnace (Solax  
Technology Ltd., China) and the results were reported as  
ash-free dry mass (AFDM). Growth rates inferred from  
AFDM measurement data were calculated for the expo-  
nential phase (Biggs 1990) and were expressed as  
micrograms of AFDM per unit area of glass substrate per  
day.

Forty milliliters ~~was~~ put into Teflon jars to assay metal  
concentrations, following two protocols (Behra et al.  
2002): (i) 20 mL was used for determination of total  
metal in the biofilm, and (ii) 20 mL was washed for 10  
min with 4 nM EDTA at pH 8 to determine the intra-  
cellular metal content of the periphyton. After filtration  
and mineralization of each sample (after washing with  
EDTA or not), cadmium concentrations were measured by  
atomic absorption spectrophotometry (Varian AA400—  
Zeeman correction, Victoria, Australia) and by autosam-  
pler, with a  $0.1 \mu\text{g}\cdot\text{L}^{-1}$  detection limit. Finally, ~~5 mL was~~  
preserved in a formaldehyde solution for countings and  
diatom identifications to the species level (see Quantita-  
tive and Qualitative Analysis of Diatom Assemblages,  
below).

## Quantitative and Qualitative Analysis of Diatom Assemblages

Enumeration was done in each formalin-preserved sample  
(100  $\mu\text{L}$ ) using a Nageotte counting chamber: the total  
number of cells counted in 10 fields (1.25  $\mu\text{L}$  each, 0.5 mm  
deep) using light microscopy at  $400\times$  magnification (pho-  
tomicroscope Leica DMRB, Wetzlar, Germany) was then  
recorded as cells per unit area of sampled substrate  
(number of diatom cells $\cdot\text{cm}^{-2}$ ).

The sample fractions devoted to taxonomic analysis of  
diatom assemblages were prepared according to ANSP  
protocols (Charles et al. 2002), i.e., digestion in boiling  
hydrogen peroxide (30%  $\text{H}_2\text{O}_2$ ) and hydrochloric acid  
(35%) followed by three cycles of centrifugation of the  
sample and pellet rinsing with distilled water. After the  
last treatment, the pellet was once again resuspended in  
distilled water and pipetted onto coverslips, which were  
mounted onto slides after air-drying, using the high  
refractive index medium Naphrax (Brunel Microscopes  
Ltd., UK; RI=1.74). Diatom identifications were con-  
ducted at a magnification of  $\times 1000$ ; individual microscope  
fields were scanned until a minimum of 400 valves had  
been identified using specialized literature (Krammer and  
Lange-Bertalot 1986–1991). Individual deformities (twis-  
ted cells and/or diatoms with deformed valve wall  
ornamentation) were observed and their frequency deter-  
mined. Note that the valves of initial cells, which  
typically have a morphology different from that of

**Table 1** Physical and chemical parameters and nutrient concentrations measured during the 6-week experimental period in the water column of the three experimental systems

		EU1	EU2	EU3
pH	Week 0	7.1 ± 0.0	7.0 ± 0.0	7.0 ± 0.0
	Week 1	7.4 ± 0.1	7.5 ± 0.1	7.4 ± 0.1
	Week 2	7.3 ± 0.0	7.5 ± 0.0	7.2 ± 0.0
	Week 4	7.7 ± 0.0	8 ± 0.0	7.8 ± 0.0
	Week 6	7.9 ± 0.0	7.9 ± 0.0	7.7 ± 0.0
Temperature (°C)	Week 0	18.0 ± 0.2	18.3 ± 0.2	17.8 ± 0.2
	Week 1	18.8 ± 0.1	17.6 ± 0.1	18.0 ± 0.1
	Week 2	18.9 ± 0.1	17.9 ± 0.2	18.2 ± 0.1
	Week 4	18.8 ± 0.1	16.5 ± 0.2	17.4 ± 0.2
	Week 6	18.7 ± 0.1	17.6 ± 0.2	18.1 ± 0.1
Dissolved O <sub>2</sub> (mg·L <sup>-1</sup> )	Week 0	8.5 ± 0.2	8.9 ± 0.2	8.3 ± 0.2
	Week 1	8.7 ± 0.1	8.4 ± 0.1	8.3 ± 0.1
	Week 2	8.5 ± 0.1	8.2 ± 0.1	8.2 ± 0.0
	Week 4	9.2 ± 0.1	8.9 ± 0.1	9.2 ± 0.1
	Week 6	8.8 ± 0.0	8.3 ± 0.1	8.2 ± 0.1
Oxygen saturation (%)	Week 0	89.0 ± 1.1	89.1 ± 1.1	87.1 ± 1.1
	Week 1	93.5 ± 0.9	87.6 ± 0.9	86.5 ± 0.9
	Week 2	91.7 ± 0.5	87.9 ± 1.5	85.4 ± 0.9
	Week 4	97.3 ± 0.5	86.9 ± 0.9	97.5 ± 0.7
	Week 6	94.2 ± 0.6	86.3 ± 1.0	84.9 ± 1.4
Conductivity (μS·cm <sup>-1</sup> )	Week 0	143 ± 6.2	152 ± 6.2	135 ± 6.2
	Week 1	153 ± 1.2	180 ± 2.5	139 ± 1.0
	Week 2	163 ± 1.5	201 ± 3.5	155 ± 2.5
	Week 4	183 ± 1.1	254 ± 2.5	181 ± 2.1
	Week 6	202 ± 2.1	320 ± 5.6	220 ± 3.5
Cadmium (μg·L <sup>-1</sup> )	Week 0	<d.l.	5.34 ± 0.22	70.38 ± 3.74
	Week 1	<d.l.	8.26 ± 0.59	106.59 ± 4.46
	Week 2	<d.l.	7.87 ± 0.41	92.52 ± 2.48
	Week 4	<d.l.	12.90 ± 1.19	101.65 ± 4.03
	Week 6	<d.l.	9.72 ± 0.14	110.50 ± 2.84
Nitrates (mg·L <sup>-1</sup> )	Week 0	33.5	33.3	32.6
	Week 1	27.3	24.4	27.6
	Week 2	25.7	22.5	28.0
	Week 4	22.2	23.6	31.6
	Week 6	18.9	29.6	37.2
Orthophosphates (mg·L <sup>-1</sup> )	Week 0	2.9	2.9	2.9
	Week 1	2.5	2.1	2.4
	Week 2	2.3	1.9	2.2
	Week 4	1.7	1.9	1.9
	Week 6	0.9	1.9	2.2

Note. Data are means ± standard deviation. d.l.: detection limit, 0.1 μg Cd·L<sup>-1</sup>. The water was changed at the end of weeks 3 and 5

vegetative valves (larger and usually presenting distorted costae) (Round et al. 1990), were not taken into account in diatom counts.

## Data Analyses

Data were evaluated statistically in two ways. First, variations in DW, AFDM, density, and abnormal form frequencies with treatments and duration of exposure were examined by means of a linear mixed-effects (LME) model for repeated measurements, with the treatments and dates as fixed effects and the subsamples taken from each artificial channel as random effects. In this case, averaging is only an approximate analysis, and a specific statistical approach adapted to the subsampling design used is needed to avoid inflated test statistics, deflated *p*-values, or incorrect rejection of the null hypothesis. These statistical analyses were computed with the statistical modeling environment R (Ihaka and Gentleman 1996). Second, PC-ORD software (McCune and Mefford 1999) was used for conducting Hierarchical Cluster Analysis. A dendrogram based on relative abundances of the species occurring most (the 80 species that had the highest cumulative relative abundances) was drawn using Ward's method.

Values are mean ± standard deviation.

## Results

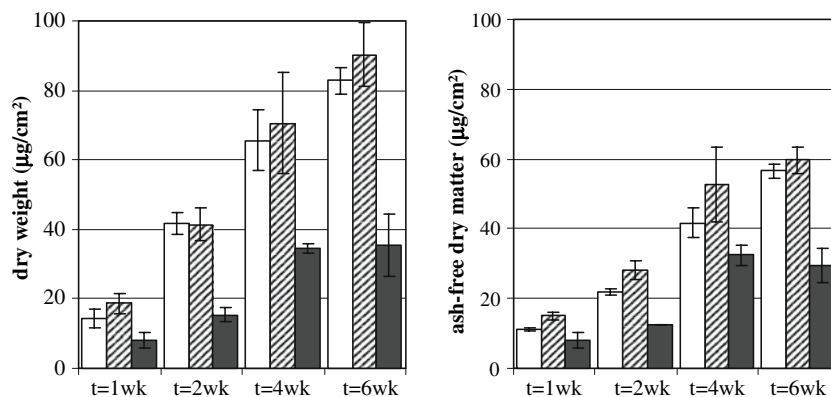
### Physical and Chemical Characteristics of Microcosm Water (See Table 1)

The pH of the water did not differ significantly (*p* = 0.77) among experimental units. A slight increase was observed, ranging from 7 at the beginning of the experiment to 8 on the last dates. Temperature and oxygenation were relatively constant throughout the experimental period and similar between treatments. Temperatures averaged 17.9°C (±0.8°C). Dissolved oxygen concentrations were stable, with a mean value of 8.7 mg·L<sup>-1</sup> (±0.8 mg·L<sup>-1</sup>), and oxygen saturation was quite high, at 92% (±4%). The mean conductivity increased regularly in all streams, from 140 μS·cm<sup>-1</sup> to reach 200 μS·cm<sup>-1</sup> in EU1, 220 μS·cm<sup>-1</sup> in EU3, and 320 μS·cm<sup>-1</sup> in EU2.

Nitrate and phosphate concentrations were around 33 mg·NO<sub>3</sub>·L<sup>-1</sup> and 2.9 mg·PO<sub>4</sub>·L<sup>-1</sup> in all systems at the beginning of the experiment (week 0). Concentrations decreased gradually in all the EUs, until additions of culture medium at the end of weeks 3 and 5 restored nutrient to comparable levels in all systems, followed by stabilization. In EU1, however, a continuous decrease in nitrate and phosphate concentrations occurred.

Cadmium concentrations in the EUs matched the nominal concentrations well. Actual concentrations were below

**Fig. 2** Composition of the periphytic layer in the experimental units: white bars—EU1, 0  $\mu\text{g Cd}\cdot\text{L}^{-1}$ ; hatched bars—EU2, 10  $\mu\text{g Cd}\cdot\text{L}^{-1}$ ; gray bars—EU3, 100  $\mu\text{g Cd}\cdot\text{L}^{-1}$ . Error bars: standard deviation of three replicates



296 0.1  $\mu\text{g}\cdot\text{L}^{-1}$  in EU1,  $9.07 \pm 2.55 \mu\text{g}\cdot\text{L}^{-1}$  in EU2, and  
297  $95.93 \pm 14.57 \mu\text{g}\cdot\text{L}^{-1}$  in EU3 during the experiment.

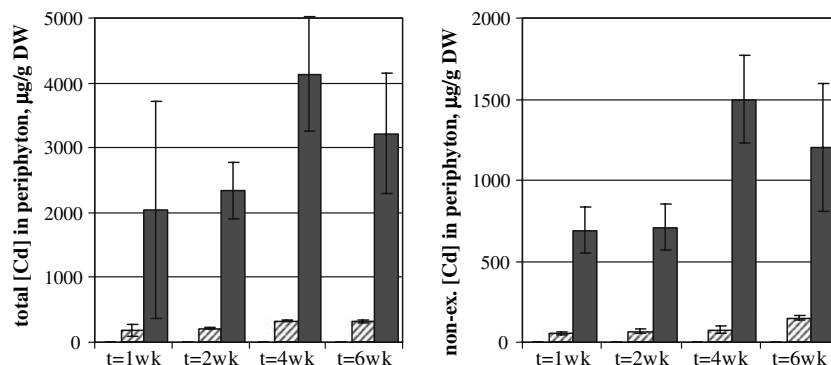
### 298 Biofilm Biomass and Metal Content

299 The mean DW and AFDM of the biofilms increased  
300 throughout the 6-week exposure (Fig. 2). In EU3, however,  
301 DW and AFDM measurements tended to stabilize between  
302 week 4 and week 6. The LME model performed on the  
303 dataset confirmed this date effect (mean DW and AFDM  
304 ranged from 13.6 and 11.4  $\mu\text{g}\cdot\text{cm}^{-2}$ , respectively, at week  
305 1 to 69.5 and 48.5  $\mu\text{g}\cdot\text{cm}^{-2}$  at week 6;  $p < 0.05$  in both  
306 cases) and also underlined a treatment effect. EU1 and EU2  
307 exhibited similar DW values during the whole course of the  
308 experiment, whereas EU3 values were half those of EU1  
309 and EU2 for almost all dates. Indeed, the tests did not  
310 discriminate EU1 and EU2 but expressed strong differ-  
311 ences from EU3 (with  $p < 0.01$  for the treatment  $\times$  date  
312 effect at weeks 2 and 6). A slight but statistically signifi-  
313 cant increase in AFDM ( $p = 0.08$  at week 4) was  
314 observed between EU1 (mean AFDM =  $32.8 \mu\text{g}\cdot\text{cm}^{-2}$ )  
315 and EU2 (mean AFDM =  $38.9 \mu\text{g}\cdot\text{cm}^{-2}$ ). AFDM in EU3  
316 was significantly reduced for all stages of settlement  
317 ( $p < 0.01$ ), with an average value of  $20.6 \mu\text{g}\cdot\text{cm}^{-2}$ .  
318 Growth rates in the exponential phase were compar-  
319 able for EU1 ( $1.4 \mu\text{g}\cdot\text{cm}^{-2}\cdot\text{day}^{-1}$ ;  $R^2 = 0.99$ ) and EU2

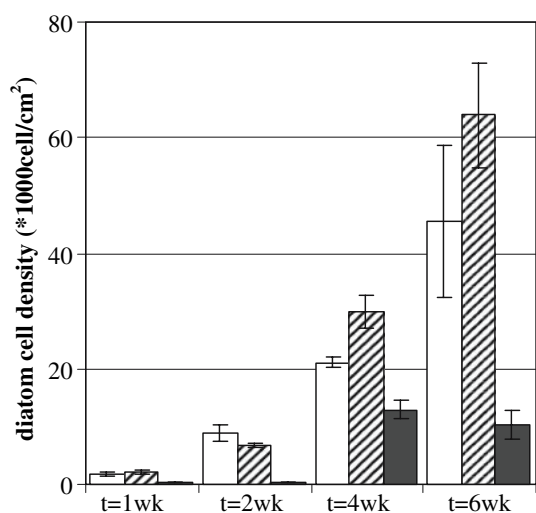
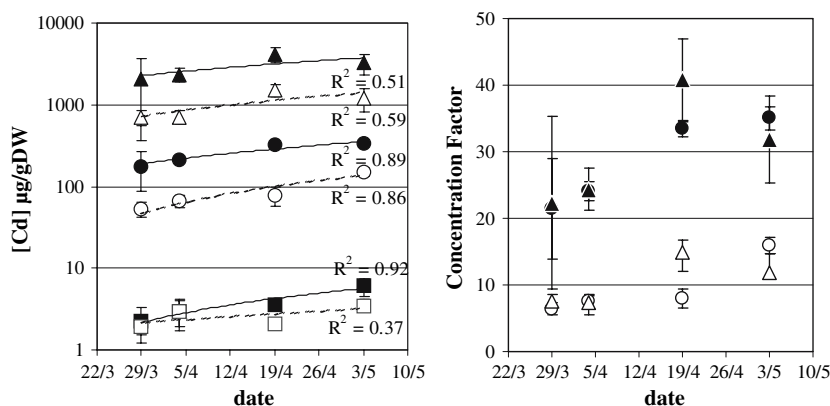
( $1.6 \mu\text{g}\cdot\text{cm}^{-2}\cdot\text{day}^{-1}$ ;  $R^2 = 0.87$ ). A cadmium overload of  
320  $100 \mu\text{g}\cdot\text{L}^{-1}$  in EU3 induced a slight decrease in growth  
321 rate ( $0.9 \mu\text{g}\cdot\text{cm}^{-2}\cdot\text{day}^{-1}$ ;  $R^2 = 0.73$ ).  
322

323 Metal contents in biofilms as measured with both proto-  
324 cols (without and after an EDTA wash) reflected their  
325 exposure history. Biofilms from the cadmium-enriched  
326 systems (EU2 and EU3) sequestered increasing concentra-  
327 tions per DW unit as the biofilm settled. We observed a  
328 logarithmic accumulation of nonexchangeable and total  
329 cadmium in periphyton (Figs. 3 and 4). Cadmium contents  
330 increased from week 1 to week 4, and then stabilized at  
331 around 3 and 5  $\mu\text{g}\cdot\text{g}^{-1}$  DW in EU1 for nonexchangeable  
332 and total cadmium, respectively, 115 and 330  $\mu\text{g}\cdot\text{g}^{-1}$  DW  
333 in EU2, and 1350 and 3700  $\mu\text{g}\cdot\text{g}^{-1}$  DW in EU3. Within any  
334 given EU, cadmium concentrations in periphyton without  
335 and after EDTA wash varied during the course of the  
336 experiment: the ratio nonexchangeable/total metal was  
337 about 0.4 for EU3 and increased in EU2 from 0.2 at week 1  
338 to 0.4 at the end of the experiment. In addition, concentra-  
339 tion factors (CFs) of the biofilm for cadmium were  
340 calculated according to Foster (1976) by dividing concen-  
341 trations of cadmium in biofilms (total and nonexchangeable  
342 fractions) by those in water, reflecting an increasing accu-  
343 mulation ability of the biofilms with dissolved metal  
344 concentrations and duration of exposure (Fig. 4). Gross  
345 uptake capacity (i.e., the ratio of total cadmium accumulated  
346 to dissolved metal) increased regularly, from  $21.5 \pm 7.6$  at

**Fig. 3** Biofilm accumulation of total and nonexchangeable cadmium in the experimental units: white bars—EU1, 0  $\mu\text{g Cd}\cdot\text{L}^{-1}$ ; hatched bars—EU2, 10  $\mu\text{g Cd}\cdot\text{L}^{-1}$ ; gray bars—EU3, 100  $\mu\text{g Cd}\cdot\text{L}^{-1}$ . Error bars: standard deviation of three replicates



**Fig. 4** Non-exchangeable (open symbols) and total cadmium (filled symbols) accumulation in biofilms and concentration factor  $[(\mu\text{g}\cdot\text{g}^{-1}, \text{DW})/(\mu\text{g}\cdot\text{L}^{-1})]$  during the course of the experiment. (●) EU1, 0  $\mu\text{g Cd}\cdot\text{L}^{-1}$ ; (○) EU2, 10  $\mu\text{g Cd}\cdot\text{L}^{-1}$ ; (△) EU3, 100  $\mu\text{g Cd}\cdot\text{L}^{-1}$ . Error bars: standard deviation of three replicates. Note the logarithmic scale on the Y axis in the left-hand plot



**Fig. 5** Diatom cell densities enumerated in the glass substrates of each experimental unit: white bars—EU1, 0  $\mu\text{g Cd}\cdot\text{L}^{-1}$ ; hatched bars—EU2, 10  $\mu\text{g Cd}\cdot\text{L}^{-1}$ ; gray bars—EU3, 100  $\mu\text{g Cd}\cdot\text{L}^{-1}$ . Error bars: standard deviation of three replicates

347 week 1 to  $35.1 \pm 1.7$  at week 6 in EU2 (with a regression  
348 coefficient of 0.41;  $R^2 = 0.93$ ) and from  $22.3 \pm 13.0$  at week  
349 1 to  $40.9 \pm 6.1$  at week 4 (regression coefficient, 0.93;  
350  $R^2 = 0.97$ ) before decreasing to  $31.8 \pm 6.5$  in EU3. The  
351 calculations of CF after nonexchangeable cadmium values  
352 also expressed an increasing sorption activity of the cells  
353 grown under 10 and 100  $\mu\text{g Cd}\cdot\text{L}^{-1}$  concentrations: from  
354  $6.4 \pm 0.9$  and  $7.6 \pm 1.1$  (week 1) to  $15.9 \pm 1.2$  (week 6) and  
355  $14.8 \pm 1.9$  (week 4) in EU2 and EU3, respectively, with  
356 regression slopes of 0.25 ( $R^2 = 0.80$ ) and 0.38 ( $R^2 = 0.91$ ).  
357 In EU3, CF values tended to decrease slightly between week  
358 4 and week 6 (CF =  $11.9 \pm 2.7$ ).

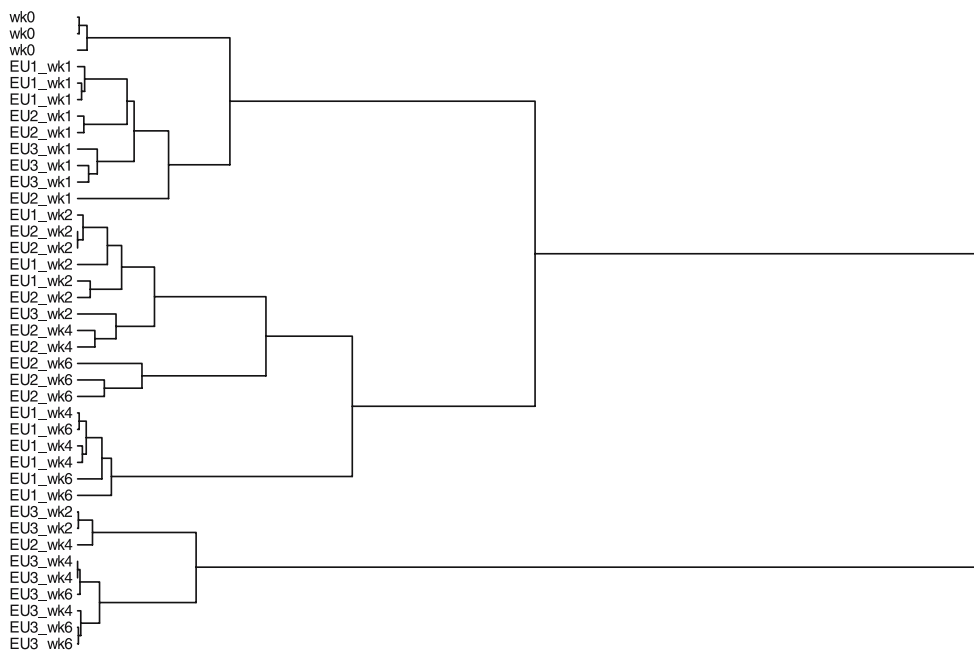
### 359 Diatom Assemblages

360 EU1 and EU2 samples had initial diatom densities of  
361  $2000 \pm 150$  cells·cm<sup>-2</sup> after 1 week of colonization

(Fig. 5). These cell densities increased strongly during the 362  
363 experiment ( $p < 0.01$ ), to  $55,000 \pm 2700$  cells·cm<sup>-2</sup> at  
364 week 6 (average EU1 and EU2 values). Statistically signifi- 365  
366 cant differences ( $p = 0.07$ ) in diatom densities between  
367 EU1 and EU2 were observed only after a 6-week incubation, 368  
369 and not at the other sampling dates. EU3 displayed  
370 dramatically low cell densities for all stages of settlement 371  
372 ( $p < 0.01$  at week 6), compared to control and low cad-  
373 mium contamination units. Data ranged from  $350 \pm 40$  374  
375 cells·cm<sup>-2</sup> at week 1 to  $10,400 \pm 2600$  cells·cm<sup>-2</sup> at week  
376 6. Positive correlations were also observed between diatom  
377 densities and AFDM measurements from the three EUs  
378 ( $R^2 = 0.67$ ;  $p < 0.01$ ); analyses of each EU separately  
379 showed very similar results. 380

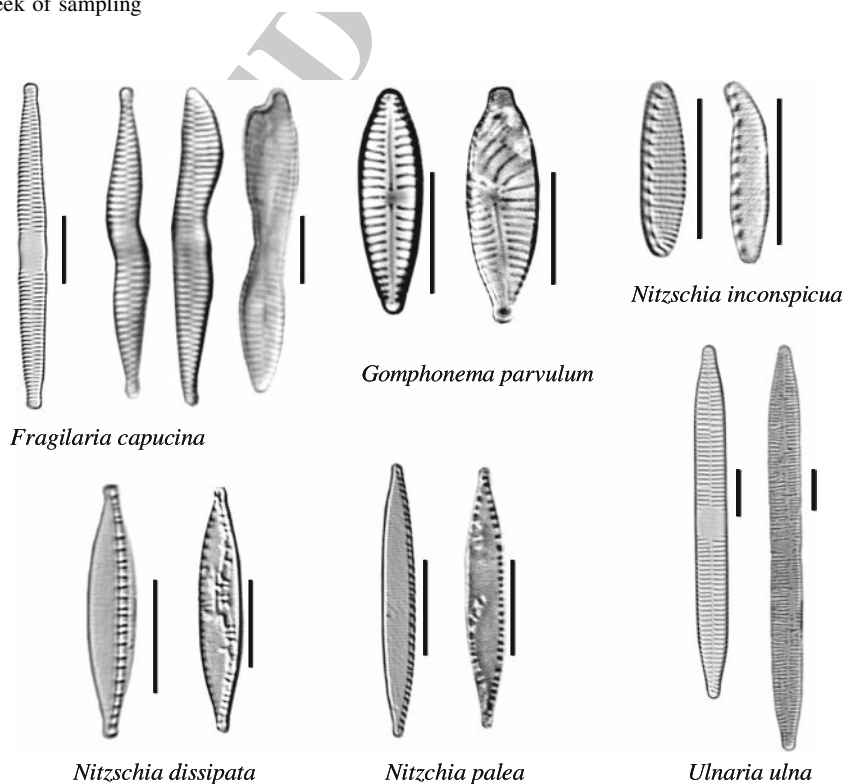
381 About 160 different taxa were identified. In Fig. 6, the  
382 dendrogram based on relative species abundances showed 383  
384 two major clusters, with one comprising diatom commu-  
385 nities grown under 100  $\mu\text{g Cd}\cdot\text{L}^{-1}$  contamination after 2, 386  
387 4, and 6 weeks' exposure, which were strongly dominated  
388 by *Nitzschia palea* (Kützing) W. Smith (from more than 389  
390 50% relative abundances to about 80% at week 6, with a  
391 species richness of  $< 20$  taxa). The other cluster was  
392 divided into two subclusters: one subcluster included 393  
394 communities from the inoculum (week 0) and week 1 for  
395 every EU, and the other subcluster was composed of 396  
397 assemblages from EU1 and EU2 developed after 2, 4, and 6  
398 weeks. Communities from the inoculum (week 0) and from  
399 the first sampling date (week 1) were characterized by an  
400 association of *Navicula lanceolata* (Agardh) Ehrenberg,  
*Navicula gregaria* Donkin, *Surirella brebissonii* Krammer  
& Lange-Bertalot var. *brebissonii*, *Nitzschia dissipata*  
(Kützing) Grunow, and *Achnanthydium minutissimum*  
(Kützing) Czarnecki, while the other subcluster, corre-  
sponding to EU1 and EU2 communities developed after a  
2-week period, contained *Nitzschia palea*, *Eolimna minima*  
(Grunow) Lange-Bertalot, *Planothidium frequentissimum*  
(Lange-Bertalot) Lange-Bertalot, *Navicula gregaria*, and  
*Nitzschia pusilla* (Kützing) Grunow. Naviculaceae pre-  
sented 40% to 70% relative abundances in the control unit





**Fig. 6** Dendrogram of assemblages relationships among diatom communities developed under three cadmium concentrations and sampled on five sampling dates. EU, experimental unit; wk, week of sampling

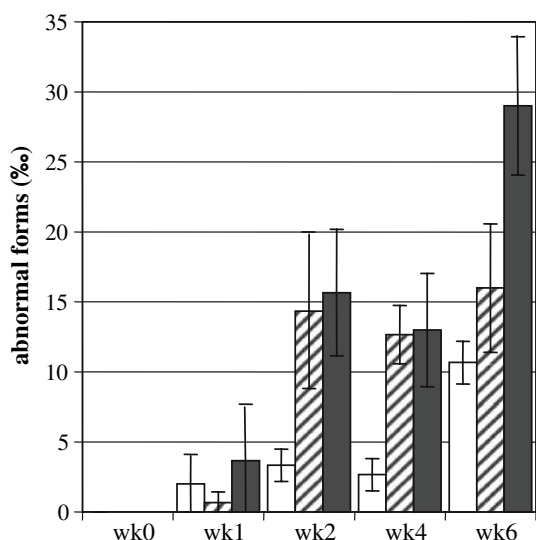
**Fig. 7** Normal and deformed diatoms. Scale bar = 10 μm



401 and tended to decrease under high cadmium exposure,  
 402 unlike Nitzschiaceae, which tended to increase, represent-  
 403 ing more than 80% relative abundances at the end of the  
 404 experiment.

A total of 21 diatom species representing 10 genera 405  
 exhibited morphological deformities (Fig. 7) and were 406  
 quantified. Teratological valves were abundant among 407  
 Raphids (66% of the abnormalities enumerated) belonging 408

Author Proof



**Fig. 8** Average relative abundances of deformed frustules in each experimental unit: white bars—EU1, 0 µg Cd·L<sup>-1</sup>; hatched bars—EU2, 10 µg Cd·L<sup>-1</sup>; gray bars—EU3, 100 µg Cd·L<sup>-1</sup>. Error bars: standard deviation of three replicates

409 to the genera *Nitzschia* (and, occasionally, to the genera  
410 *Caloneis*, *Eolimna*, *Gomphonema*, *Navicula*, *Sellaphora*,  
411 and *Surirella*) and Araphids (33%) such as *Fragilaria* sp.  
412 and a few occurrences of *Meridion*. Abnormalities were  
413 less abundant in Monoraphids from the genera *Planothi-*  
414 *dium*. In Fig. 8 we observed that no teratological forms  
415 were present in the inoculum; in the experimental units,  
416 however, some abnormalities tended to appear, with  
417 increasing abundances in EU2 and EU3, reaching  
418  $16 \pm 4.5\%$  in EU2 and  $35 \pm 5\%$  in EU3 after a 6-week  
419 exposure. The frequency of valve abnormalities in the  
420 control units was quite stable between week 1 and week 4  
421 ( $2.7 \pm 0.7\%$ ) and increased to more than 10% at week 6,  
422 which was, however, lower than the frequencies calculated  
423 in EU2 and EU3, in which combined effects between  
424 duration and level of exposure were highlighted ( $p < 0.05$ ).

## 425 Discussion

### 426 Experimental Setup

427 In this study there were three different experimental units  
428 (EUs) that consisted of six different slides each. To ensure  
429 that the starting conditions (inoculum, physical and  
430 chemical conditions) were the same in the different EUs,  
431 water characteristics and diatom community composition  
432 were determined at week 0. Every EU received a different  
433 treatment (0, 10, 100 µg·L<sup>-1</sup>), but not all slides from each  
434 channel within an EU were completely independent of each  
435 other, and they were consequently considered as subsam-  
436 ples. On the other hand, the analyses performed would not

437 have been possible in  $3 \times 3$  systems, because of the time  
438 involved, especially in the determination of cadmium  
439 concentrations as well as the community composition of  
440 the biofilms. Because the treatments were not “truly”  
441 replicated, the experimental design required the use of  
442 mixed-effects modeling techniques, and the results need to  
443 be taken with caution.

444 It has been reported that cultivated phototrophic biofilms  
445 can develop as small independent ecosystems, showing  
446 poor reproducibility of community composition between  
447 microcosms exposed to comparable treatments (Roeselers  
448 et al. 2006). In the present study, some trends were  
449 observable in all the systems, e.g., nitrogen and phosphorus  
450 decreases or slight increases in pH, likely related to  
451 nutrient consumption and photosynthetic activity of the  
452 primary producers, respectively. However, conductivity in  
453 EU2 increased more than in EU1 and EU3, perhaps  
454 reflecting heterogeneity created by the intrinsic complex  
455 behavior of the microbial communities (Kangas and Adley  
456 1996; Vandermer et al. 2002).

457 Cadmium concentrations used in this study were 0, 10,  
458 and 100 µg·L<sup>-1</sup>, which led to different dynamics depend-  
459 ing on the nominal values imposed in the systems;  
460 however, the absence of experimental data between 10 and  
461 100 µg Cd·L<sup>-1</sup> did not allow any precise evaluation of the  
462 cadmium threshold for negative effects on biofilm biomass  
463 and diatom community structure.

### 464 Sorption Kinetics of Dissolved Cadmium in the 465 Biofilms

466 Cadmium concentrations in the biofilms differed between  
467 EUs and temporally within EUs. Generally, an increase in  
468 metal concentrations in periphyton reflected the duration of  
469 exposure and the dissolved concentrations of this metal in  
470 the culture media of each experimental unit. Hence, the  
471 EU3 biofilms had much higher concentrations at any  
472 sampling time than those of EU2 and EU1 (Fig. 3). It has  
473 been demonstrated that biofilms have a good bioaccumu-  
474 lation capacity (Guanzon et al. 1995; Sunda and Huntsman  
475 1998; Chang and Reinfelder 2000; Hill et al. 2000) and  
476 could be used to monitor river pollution equivalent to metal  
477 measurements in the sediment and suspended solids (Fuchs  
478 et al. 1996). Indeed, heavy metal ions can be entrapped in  
479 the organic matrix and occasionally biosorbed by live cells  
480 (bacteria, fungi, and algae). As observed in this experi-  
481 ment, large amounts of metals assayed in the biofilms were  
482 not actually taken up by the cells; 60% of the metal was  
483 rather absorbed on the cell surface (Torres et al. 1998) and  
484 hence eliminated by the EDTA wash.

485 Metal bioaccumulation reflects both passive and active  
486 modes (Campbell et al. 2002); most studies dealing with

487 the subject describe an initial rapid phase of biosorption  
 488 (Wang and Dei 2001; Hudson 2005), followed by slower  
 489 active metal uptake (Perrein-Ettajani et al. 1999); however,  
 490 most of the studies relative to biofilm bioaccumulation  
 491 capacities describe short-term sorption kinetics. The loga-  
 492 rithmic regressions (Fig. 4) expressed an increasing  
 493 accumulation of cadmium per unit DW during the first 4  
 494 weeks of the experiment, then stabilization at the end of the  
 495 experiment, indicating some kind of steady state, for total  
 496 as well as nonexchangeable cadmium. Increasing cadmium  
 497 sorption (until week 4) with increasing biofilm biomass  
 498 implied a significant contribution of subsurface cells to  
 499 metal sorption, these cells providing additional sorption  
 500 sites (Hill et al. 2000). However, the effectiveness of  
 501 additional biofilm biomass at removing metal was limited  
 502 on the last dates of the experiment when a steady state was  
 503 reached. This may be due to saturation of the metal binding  
 504 sites in the cells (Di Toro et al. 2001; Rijstenbil and Ger-  
 505 ringa 2002), limiting cell absorption capacity, according to  
 506 the decrease in CF at the end of the experiment. It may also  
 507 be explained by the three-dimensional architecture of the  
 508 biofilm, strongly modified under metal exposure. Indeed, in  
 509 copper-treated biofilms, Teitzel and Parsek (2003)  
 510 observed an external layer of dead cells and an increased  
 511 number of live cells toward the substratum. Presumably,  
 512 most of the actively growing cells were in a “protected”  
 513 region of the biofilm, where metal penetration was retar-  
 514 ded. The protective effect of the organic matrix against  
 515 pollution by toxins has been underlined by studies com-  
 516 paring the effects of heavy metals on thin vs. thick biofilms  
 517 (Hill et al. 2000) or on biofilms exposed at various stages of  
 518 development (Ivorra et al. 2000). This protective role of the  
 519 matrix has been attributed to local pH and hypoxia con-  
 520 ditions in the internal layers of the biofilm (Teissier and  
 521 Torre 2002), which modify the redox conditions and,  
 522 possibly, cell bioaccumulation potentials. Moreover, it is  
 523 generally accepted that exopolysaccharides secreted by  
 524 algal and bacterial communities (Pistocchi et al. 1997;  
 525 Decho 2000; Muller et al. 2005) also play a role in binding  
 526 metals, thus reducing their bioavailability and toxicity  
 527 toward live cells.

## 528 Influence of Metal Contamination Level on Biofilms

529 In this study, biofilm accumulation was affected by high  
 530 levels of dissolved cadmium: DW and AFDM were sig-  
 531 nificantly lower in biofilms exposed to a 100  $\mu\text{g Cd}\cdot\text{L}^{-1}$   
 532 contamination (EU3) than those measured in biofilms  
 533 grown in EU1 and EU2. Growth inhibition at high metal  
 534 contamination levels, as shown by the decrease in growth  
 535 rate in EU3, has been widely reported (Conway 1978;  
 536 Conway and Williams 1979; Wong 1987; Guanzon et al.

1994; Payne and Price 1999; Gold et al. 2003a). Not only  
 global metabolism affected by metals (Husaini and Rai  
 1991), but so is cell ultrastructure (endoplasmic reticulum,  
 mitochondria), which seems to be modified by elevated  
 intracellular cadmium concentrations (Wong 1987).  
 Moreover, Guanzon et al. (1994) described perturbations in  
 phosphorus metabolism and cell division over critical  
 concentrations of metal. Takamura et al. (1989) found  
 growth inhibition of freshwater benthic algae at far lower  
 cadmium concentrations than those applied in EU3, and  
 our findings provide added support for the idea that DW  
 and AFDM are better indicators of metal damage to biofilm  
 communities than microbenthic algal photosynthesis  
 (Lehmann et al. 1999; this study, data not shown), on  
 which substances like cadmium and zinc have an unспе-  
 cific mode of action (Lehmann et al. 1999).

Although dissolved cadmium in EU2 exceeded metal  
 concentrations normally regarded as toxic for chemical  
 waste, biomass as assessed by AFDM was comparable  
 between EU1 (reference) and EU2 (10  $\mu\text{g Cd}\cdot\text{L}^{-1}$ ), and  
 DW was even slightly higher in EU2 than in EU1. Nale-  
 wajko (1995) showed that moderate additions of cadmium  
 (up to 5  $\mu\text{g Cd}\cdot\text{L}^{-1}$ ) may result in some growth stimula-  
 tion. Nutrient bioavailability would also be likely to  
 interact with the toxicant and slightly favor biofilm  
 development (Lozano and Pratt 1994), nutrient concentra-  
 tions being higher in EU2 at weeks 4 and 6 compared to the  
 reference (see  $\lambda$ ). Ivorra et al. (2002) noted that increased  
 dissolved phosphorus concentrations might mitigate metal  
 toxicity to biofilms or exert a protective role, by making  
 coordination complexes of trace metals with phosphorus;  
 moreover, Barranguet et al. (2002) described nutrient-  
 stimulated biomass accumulation in photosynthetic bio-  
 films affected by metals. This complicates the assessment  
 of toxic effects on living organisms (and especially dia-  
 toms) embedded in the biofilm.

## Influence of the Metal Contamination Level on Diatom Assemblages

It was expected that differences in cadmium concentrations  
 between the systems would control the quantitative as well  
 as the qualitative characteristics of the diatom assemblages.

Diatom growth played a role in the development of the  
 biofilm by dynamically contributing to the pool of live  
 organisms (export and import of cells) as well as producing  
 polysaccharide exsudates, which are partly responsible for  
 the coherence of the organic matrix. Whatever the stage of  
 colonization, Stevenson and Peterson (1991) estimated  
 substantial daily immigration and emigration rates for  
 diatoms in slow-flowing streams. In the present study,  
 specific growth rates were observed, diatom densities

**Table 2** Metal sensitivity and tolerance of diatom species identified in the experiment, as described in previous mesocosm (M) and field (F) studies

Species	Described as metal-sensitive in <sup>a</sup>	Described as metal-tolerant in <sup>a</sup>
<i>Achnanthes minutissimum</i> (Kützing) Czarnecki	3 (M), 18 (F)	4 (F), 5 (F-M), 7(M), 8(M), 11 (M), 16 (F), 20 (F)
<i>Amphora pediculus</i> (Kützing) Grunow	18 (F)	20 (F)
<i>Eolimna minima</i> (Grunow) Lange-Bertalot		5 (F-M), 6 (F), 17 (M) , 20 (F)
<i>Fragilaria capucina</i> Desmazières	6 (F)	11 (M), 13 (F), 19 (M), 20 (F)
<i>Gomphonema parvulum</i> (Kützing) Kützing	11 (M)	5 (F-M), 6 (F), 10 (M), 18 (F)
<i>Mayamaea atomus</i> var. <i>permitis</i> (Hustedt) Lange-Bertalot		15 (F)
<i>Navicula lanceolata</i> (Agardh) Ehrenberg		20 (F)
<i>Navicula(dicta) seminulum</i> (Grunow) Lange Bertalot		9 (F)
<i>Nitzschia dissipata</i> (Kützing) Grunow	5 (M), 7 (M), 8 (M)	20 (F)
<i>Nitzschia palea</i> (Kützing) W.Smith		1 (M), 2 (M), 8 (M), 12 (F), 14 (F), 18 (F)
<i>Planothidium lanceolatum</i> (Brebisson ex Kützing) Lange-Bertalot		20 (F)
<i>Surirella angusta</i> Kützing		5 (F-M), 6 (F), 21 (M)

<sup>a</sup> Numbers refer to the following papers: 1, Admiraal et al. (1999a, b) ; 2, Admiraal et al. (1999a, b); 3, Blanck et al. (2003) ; 4, Cattaneo et al. (2004); 5, Feurtet-Mazel et al. (2003); 6, Gold et al. (2002); 7, Gold et al. (2003a); 8, Gold et al. (2003b); 9, Ivorra et al. (1999); 10, Ivorra et al. (2002a); 11, Ivorra et al. (2002b); 12, Lai et al. (2003); 13, Lehmann et al. (1999); 14, Medley and Clements (1998); 15, Morin et al. (2007); 16, Nunes et al. (2003); 17, Peres et al. (1997); 18, Sabater (2000); 19, Shehata et al. (1999); 20, Szabó et al. (2005); 21, Takamura et al. (1989)

587 increasing regularly in EU1 and EU2. In the streams  
588 exposed to high cadmium concentrations, growth rates  
589 were diminished due to chemical stress affecting cell sur-  
590 vival, cell division (Perrein-Ettajani et al. 1999), or  
591 reproduction rates (Rott 1991; Peres et al. 1997) or even  
592 modifying migration strategies (Peterson 1996). These  
593 structural and functional perturbations led to the dramatic  
594 reduction we observed in diatom biomass, as described by  
595 Paulsson et al. (2000). As shown in Figs. 2 and 5 and  
596 indicated by correlations, diatom density increases fit  
597 AFDM evolutions well, expressing the elevated quantita-  
598 tive contribution of diatoms to the pool of living cells in the  
599 biofilms (Stevenson 1996). These approximate covariations  
600 may, moreover, be assigned to a limited mortality of the  
601 cells; thus a thin layer of dead cells would probably be  
602 enough to exert a protective role on diatoms in the inner  
603 parts of the biofilms exposed to metals.

604 The diatom assemblages present during the first week  
605 were similar to the common available pool of species  
606 present in the inoculum. Then the assemblages differenti-  
607 ated according to the competitive ability to grow of the  
608 species under elevated metal exposure. However, most of  
609 the dominant taxa identified in this experiment (including  
610 the control unit) had already been reported in metal-con-  
611 taminated environments (see Table 2), even though the  
612 status of many of them is still under discussion. The use of  
613 microcosms enabled us to eliminate external agents such as  
614 grazing or other disturbances, and the only factor that was

615 liable to mitigate toxicity was resource competitive ability  
616 (Tilman 1986). Nitrogen and phosphorus are prerequisites  
617 for growth and development of autotrophic organisms, but  
618 diatoms are found with a broad range of nutrient avail-  
619 ability, conductivity, etc. (Stevenson and Pan 1999). In  
620 EU2, especially, the joint effects of metal contamination  
621 and interspecific resource competition on the composition  
622 of the ultimate community structure must be taken into  
623 consideration. Saprophytic species like *Nitzschia palea*  
624 and, to a lesser extent, *Gomphonema parvulum* (Kützing)  
625 Kützing may have been favored by the nutrient availability,  
626 mitigating the effects of the cadmium at low concentrations  
627 (Barranguet et al. 2002; Morin et al. 2007).

628 The shifts observed in diatom species in EU3 (high-level  
629 cadmium) were also consistent with data found in the lit-  
630 erature. The increased abundance of *Nitzschia palea* in  
631 EU3 coincided with the reported tolerance of this species to  
632 metal pollution (Table 2). The predominance of *N. palea*  
633 concomitant with the effect of high contamination detri-  
634 mental to most of the species initially present (decline in  
635 diatom abundances, decrease in species richness, elimina-  
636 tion of many species from the inoculum) is in agreement  
637 with the observations of Ivorra et al. (2002). A few studies  
638 of metal-tolerant algae, cited by Foster (1982), have shown  
639 that metal resistance is sometimes correlated with  
640 decreased metal uptake. This assumption may provide an  
641 additional piece of explanation for the decrease in the  
642 ability to accumulate cadmium, strong contamination

643 creating a selective environment where the shifts in com- 693  
 644 munity structure toward dominance of a particular species 694  
 645 probably led to reduced accumulation capacity of the 695  
 646 ultimate biofilms. 696

#### 647 Observations of Diatom Deformities 697

648 The communities inoculated into the three experimental 700  
 649 units showed no morphological abnormalities. In EU1 at 701  
 650 the beginning of the experiment, aberrant cells occurred at 702  
 651 low rates (<3%), similar to the rates of “naturally occur- 703  
 652 ring” deformities. From week 2 in EU2 and EU3, however, 704  
 653 there was a larger percentage of valve abnormalities, and 705  
 654 cadmium exposure was identified as the leading stressor 706  
 655 responsible for their increased occurrence. Abnormal cells 707  
 656 had already been noted in sediment cores at various stages 708  
 657 of the geological record from metal-polluted environments 709  
 658 (Ruggiu et al. 1998; Cattaneo et al. 2004), as well as in 710  
 659 fresh-water (Yang and Duthie 1993; McFarland et al. 1997; 711  
 660 Nunes et al. 2003; Szabó et al. 2005; Morin et al. 2007) and 712  
 661 in seawater (Thomas et al. 1980; Dickman 1998). In this 713  
 662 experiment, concentrations of dissolved cadmium were 714  
 663 above the “bad status” thresholds (3 µg Cd·L<sup>-1</sup>) proposed 715  
 664 by the French freshwater quality evaluation system (SEQ- 716  
 665 eau). As the percentage of abnormal cells increased with 717  
 666 increasing cadmium concentrations in the streams, and 718  
 667 with duration of exposure, it seems that cadmium is the 719  
 668 causative agent for deformed diatoms in this experimental 720  
 669 study. Moreover, the experimental units used very soft  
 670 water, with a neutral pH, which would have enhanced  
 671 metal bioavailability. The frequency of deformities in  
 672 diatom communities exposed to cadmium contaminations  
 673 confirmed the observations described previously, in sys-  
 674 tems that were mostly contaminated by metals such as Cd,  
 675 Cu, and Zn. However, the percentages of deformities  
 676 recorded here were very high compared with those in other  
 677 studies. Maybe the high values found in this experiment  
 678 indicated a more attentive consideration to slight altera-  
 679 tions overlooked in other studies.

680 The reasons underlying the appearance of large numbers  
 681 of deformed diatoms in EU1 after 6 weeks were suspected  
 682 not to involve heavy metals. Anomalous diatom cells have  
 683 commonly been described under culture conditions (Estes  
 684 and Dute 1994), in relation to the fact that the cells had  
 685 probably exceeded the lower limits for successful sexual  
 686 reproduction and were on their way to extinction; however,  
 687 deformities happen in roughly 2- to 12-year cycles. Here  
 688 we would be inclined to single out some physical damage  
 689 during cell development, related to crowding on the sub-  
 690 strate. Cells with mechanically induced deformities may  
 691 also be replicated in clonal populations as the cells divide,  
 692 keeping the frequency of aberrations stable or increasing.

The great number of diatom-based indices used in 693  
 Europe (Prygiel et al. 1999) are efficient tools for moni- 694  
 toring quality in rivers: trophic or saprobic levels (Lange- 695  
 Bertalot 1979; Sládeček 1986; Schiefele and Schreiner 696  
 1991; Kelly and Whitton 1995), acidification (Håkansson 697  
 1993), salinity (Ziemann 1991), or general water quality 698  
 (Coste, in Cemagref 1982; Dell’Uomo 1996; Lenoir and 699  
 Coste 1996). However, these methods are not suitable for 700  
 the assessment of metal pollution. A number of diatom 701  
 species identified as metal-tolerant in this study can with- 702  
 stand a wide range of environmental conditions, and further 703  
 investigations are needed to provide a pertinent classifica- 704  
 tion of taxa depending on their resistance, sensitivity, or 705  
 indifference to metal contamination. Diatom deformities 706  
 are certainly a more specific indicator of high metal con- 707  
 centrations, as emphasized by many field studies (Dickman 708  
 1998; Medley and Clements 1998; Morin et al. 2007). This 709  
 experiment confirmed the increased occurrence of such 710  
 abnormalities as a distinctive characteristic of growing 711  
 diatom biofilms exposed to cadmium. The percentage 712  
 aberrant diatoms is already used in the United States 713  
 (Stevenson and Bahls 1999) as a metrics of presumptive 714  
 metal pollution. In France an update to the SPI (Specific 715  
 Pollution Sensitivity Index [Coste, in Cemagref 1982]) is 716  
 currently under test: it takes into account the relative 717  
 abundances of deformities (abnormal diatoms having the 718  
 worst pollution sensitivity values) in order to get a better 719  
 estimate of general water quality. 720

#### 721 Conclusions 721

It is very difficult to assess the effects of metal pollutions 722  
 on diatom biofilms in field experiments. Therefore, this 723  
 study was carried out to obtain an approximation of the 724  
 toxicity of cadmium at different stages in the develop- 725  
 ment of the community using experimental microcosms. 726  
 It is difficult to account for the differences between 727  
 treatments because the physicochemical variables exhib- 728  
 ited similar trends for all treatments, with the exception 729  
 of conductivity, which increased more in one system 730  
 (EU2). Despite this, it was observed that: (i) diatom 731  
 biofilms have a good accumulation capacity, and their 732  
 growth seems to be lowered by cadmium exposure, 733  
 (ii) shifts in species composition and decreases in spe- 734  
 cies richness may be attributed to cadmium exposure, and 735  
 (iii) changes in diatom morphology (deformities) are 736  
 much more frequently observed in cadmium-treated 737  
 microcosms. 738

The present work reveals how diatom communities can 739  
 be used to provide an assessment of metal pollutions. Field 740  
 validation of the observed effects remains an interesting 741  
 subject for further investigations. 742

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