Microcosm Assessment of the Effect of an Acute Mercury Contamination Event on the Structure and Activity of Sediment Bacterial Communities

Vanessa Oliveira, Ana P Silva, Bruna Marques, Adelaide Almeida, Newton CM Gomes, Ana I Lillebo* and Angela Cunha*

Department of Biology and CESAM, University of Aveiro, Campus de Santiago, 3810-193 Aveiro, Portugal.

Corresponding author: Angela Cunha, Department of Biology and CESAM, University of Aveiro, Campus de Santiago, 3810-193 Aveiro, Portugal, Tel: +351-234-370-779; Fax: +351-234-426 408; E-mail: acunha@ua.pt

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Abstract

Objectives: In the present study, the effect of acute mercury contamination on the structure and activity of bacterial communities in intertidal mudflats was assessed through a microcosm experiment simulating the mobilization of highly contaminated sub-surface sediments.

Methods: Box-microcosms corresponding to different test conditions were constructed by mixing natural estuarine sediments with high and low concentrations of mercury in defined proportions. The effects on sediment bacteria were characterized by quantifying bacteria using fluorescent in situ hybridization (FISH) technique, assessing the community structural diversity by denaturating gradient gel electrophoresis (DGGE) and analysing descriptors of bacterial activity (extracellular enzymatic activity and leucine incorporation) at the beginning and at the end of a 7-days incubation period.

Results: At the end of the experiment, total abundance of Bacteria was significantly higher in the low-Hg microcosms than in the high-Hg and blended-sediment microcosms. DGGE patterns revealed that the structure of sediment bacterial communities responded to the experimental treatment and to the incubation time. Bacterial activity was inhibited by mercury and that the levels of arylsulfatase and biomass productivity were inversely related with the Hg concentration. The proportion of sulfate-reducing bacteria in relation to total prokaryotes increased at the end of the experiment, which might indicate a differential response of Bacteria and Archaea to confinement and mercury contamination.

Conclusion: Mechanical disturbance of sediments historically exposed to mercury contamination, like bottom trawling or dredging, will cause the mobilization of deeper sediments highly contaminated with Hg which will impact the less contaminated surface sediments. These acute events will impact the structure and activity of bacterial communities and their contributions to the associated biogeochemical cycles, with expectable impacts at the ecosystem level.

Keywords: Mercury; Bacterial communities; Sediment resuspension; Sulfate-reducing bacteria; Bacterial activity; Acute toxicity; Estuary

Introduction

Mercury is a highly toxic element that may be present in aquatic environments as a consequence of anthropogenic activities. By being persistent, easily dispersible and susceptible to bioaccumulations, mercury is considered a global concern pollutant [1]. Although the severe environmental regulations implemented in the last decades have restrained anthropogenic emissions, mercury accumulated in sediments, known as historical contamination, is still a threat to the health of aquatic ecosystems because of the potential release to other environmental compartments (atmosphere, water column, living organism) [2].

Salt marshes are particular environments with a worldwide recognized importance in providing essential ecological functions and services [3,4]. However, as transitional areas, they are more susceptible to large inputs of pollutants, namely mercury, mainly from anthropogenic sources [5,6]. Integrated interventions to minimize the effects of historical contamination by mercury have been set up in coastal ecosystems and wetlands around the world [7-9]. Recent studies have attempted to understand the mechanisms of mercury storage in salt marsh sediments and the role of plants in mercury sequestration and decontamination [10,11]. Long term exposure to mercury is known to cause changes in structure and function of sediment bacterial communities [12-16] and the methylation of mercury to the neurotoxic species methyl mercury (MeHg) is directly related to the heterotrophic activity of anaerobic bacteria. Although iron-reducing bacteria [17] and methanogens [18] can contribute to mercury methylation, and the recent evidence of this capacity in novel bacterial and archaeal strains [19], sulfate-reducing bacteria (SRB) are still regarded as major players in methylmercury production [20-22]. SRB belonging to the family Desulfobacteriaceae (Deltaproteobacteria class) that use acetate as electron donor are recognized as active mercury methylators [20,23].
The Laranjo basin, a 2 km² inner area of the Ria de Aveiro coastal lagoon, was exposed to intense contamination with mercury during the second half of the XX century. Although the industrial processes that caused this situation have changed, and mercury levels in the water column and surface sediments are in now in compliance with environmental regulations, large amounts of mercury are still present in the sub-surface sediments [7,24]. Due to human activities such as bottom trawling or dredging, or because of physical, chemical and biological processes, such as hydrodynamic flows, bioturbation or chemical transformation, Hg can be remobilized [9,25,26]. The mobilization of mercury accumulated in deeper sediments will affect the distribution of mercury within the sediment column, the processes of metal speciation and the transfer between the sediment compartment and the water column [26-29].

The aims of these experiments were: (i) to infer on the impact of a possible dredging on the structure and activity of sediment bacterial communities and (ii) to quantify the sulfate-reduction bacteria, as likely players in processes of Hg methylation. In order to simulate the mobilization of highly contaminated sediments from sub-surface sediments by dredging or other form of mechanical disturbance, control-sediment (low concentration of Hg) from the surface were experimentally mixed with deeper sediments (0.4-0.5 m) from the corresponding site (high concentration of Hg) in a microcosm experiment.

Material and Methods

Study area

The Ria de Aveiro is a temperate shallow coastal lagoon located at Northwest Atlantic coast of Portugal (40°38’N, 8° 44’W). This mesotidal system forms a complex network of channels with extensive intertidal mudflats that get exposed during low tide. In this system, the small inner Laranjo basin is historically contaminated with mercury discharged from a mercury cell chlor-alkali plant located in Estarreja industrial complex and presently, deeper sediments (down to 0.4-0.5 m) show the highest contamination levels [7]. Having in mind the well recorded history of the system, sediment samples were collected at low tide from an unvegetated area adjacent to the salt marsh in the Laranjo Basin. Sediment horizons corresponding to the 0-0.05 m (surface) and 0.4-0.5 m (deeper) were collected in the field, placed in sterile plastic bags, and kept in the cold during transport to the laboratory.

Sediment characterization

In the laboratory, each sediment layer was homogenized and cleaned from shells and other debris. Homogenised sub-samples were analysed for the percentage of fine particles (<0.63 mm), and total mercury concentration (ng Hg g⁻¹). The total mercury concentration in sediments from 0-0.05 and the 0.4-0.5 m sediment horizons was directly determined by atomic absorption spectrometry following thermal decomposition with an Advanced Mercury Analyser (AMA) LECO 254 (Costley et al. 2000). In order to assess the accuracy and precision of the analytical methodology, analysis of certified reference material was carried out (PACS-2 harbour sediment) in parallel with samples [24]. Certified and measured values were in agreement with recoveries of 95% for PACS 2. Mercury quantifications were performed in triplicate and blanks were run in parallel.

Experimental set-up

In the laboratory, the homogenised sediment was distributed in microcosm (9.5 × 9.5 × 4 cm acid-cleaned PET). The test condition (blended 1:4) was achieved by mixing the positive (high-Hg) and the negative (low-Hg) control sediments in a proportion of 1:4 (w:w). Each box-microcosm contained a total of 200 g of sediment and four replicate box-microcosms were prepared for each experimental condition. Microcosms were incubated at room temperature (22°C), at natural light. The biological effects of acute mercury contamination on sediment bacterial communities were assessed by analysing the community structure and descriptors of bacterial activity at the beginning and at the end of the incubation (7 days).

Extracellular arylsulfatase activity

The activity of arylsulfatase was determined fluorimetrically using a Jasco FP-777 fluorometer (Jasco Inc. Easton, MD USA) in sediment suspensions [30]. MUF-sulfate (Sigma-Aldrich, St. Louis MO, USA) was as used as a fluorogenic labelled substrate. The final saturating concentration, established by preliminary kinetic assays, was 2 mM. Sediment suspensions were prepared by adding 100 ml of sterile diluted artificial seawater (salinity 17, same as salinity in situ) to 1 g of fresh sediment and stirred in order to obtain homogeneous sediment suspensions. Six aliquots were transferred to 2 ml microtubes and added of the stock substrate solution. After centrifugation (13,000 × g, 5 min) for the removal of particles, and alkalinisation with 100 μl of a buffer solution (1.384 ml of ammonium, 0.375 g glycin and distilled water to 100 ml, pH 10.5) in order to enhance MUF fluorescence. The remaining three aliquots were incubated at room temperature (22°C) for 3 hours, after which particles were removed by centrifugation, the buffer solution was added and the final fluorescence was read. The rate of substrate hydrolysis was estimated from the variation of fluorescence, standardized to 1 hour incubation and converted to units of substrate concentration with a calibration curve prepared for the fluorescence product (MUF) by the internal standard approach.

3H-Leucine incorporation

3H-Leucine incorporation [31] was used to estimate bacterial biomass productivity (BBP). For each sample, three aliquots of sediment suspension plus one trichloroacetic acid (TCA)-killed control were placed into Eppendorf tubes and incubated with 483 nM of 3H-leucine (Amersham, Biosciences Ltd, Sweden, 45-80 Ci/mmol) for 1 h at room temperature. The saturation concentration and the incubation time were established in preliminary kinetics assays. Incubation was stopped with 80 μl of TCA (100%). Samples were centrifuged (13,000 g for 10 min), the supernatant was rejected, and 1 ml of cold TCA (5%) was added. Between centrifugations, 1 ml of cold TCA (5%) and cold ethanol was added. Finally, 1 ml of scintillation cocktail was added. After a 3 day period of stabilization, the radioactivity was determined in a scintillation counter (Beckman LS 6000 IC; GMI, Inc, Ramsey, Minnesota, USA).

Quantification of bacteria and sulfate reducing bacteria (SRB)

Quantification of Bacteria and SRB was performed by Fluorescent In Situ Hibridization (FISH) with a mixture of the oligonucleotide probes EUB338 [34], EUB338 II and EUB338 III (Daims et al. 1999)
for the Bacteria domain and a mixture of the oligonucleotide probes DELTA 495 a, b, c, covering most Deltaproteobacteria for sulfate reducing bacteria [32,33]. The protocol was adapted from Llobet-Brossa et al. [34], with minor modifications. Samples (0.5 g of fresh sediment) were fixed in 2% formaldehde. Fixed samples were washed with 1 × PBS, centrifuged (13000 × g for 2 min) and stored in PBS/ethanol (1:1) at -20°C. Aliquots of sediment suspension were diluted with PBS, mixed, and cells were collected by filtration on the surface of 0.2 μm pore-size polycarbonate membranes (GE Osmonics Labstore, Minnetonka, USA) [35]. Hybridizations were performed with 21 μl of hybridization buffer (900 mM NaCl, 20 mM Tris/HCl pH 8, 0.01% SDS) for both Bacteria and SRB probes (Eurofins MWG Operon, Ebersberg, Germany) and 3 μl of each probe (5 ng/μl), at 46°C for 90 min. After hybridization, the membranes were washed with a buffer solution for 15 min at 48°C. The membranes were subsequently rinsed with distilled water, air-dried, stained with DAPI, mounted in Citifluor immersion oil solution (Citifluor Ltd, London, UK) and examined under an epifluorescence microscope equipped with a mercury bulb and filter sets 31000 (Chroma) for DAPI detection and 41007a (Chroma) for Cy3 detection. Cells were counted with 1000 × magnification in 10 randomly selected optical fields. Total DAPI counts were an estimate of total number of prokaryotic cells [35].

16S rRNA denaturing gradient gel electrophoresis (DGGE) profiling of the bacterial communities

DNA was extracted from sediment using the E.Z.N.A. Soil DNA kit (Omega Bio-Tek, Norcross, GA, USA) according to the instruction of the manufacturer. PCR amplification was carried out in a Veriti 96-Well Thermal Cycler (Applied Biosystems, Foster City, CA, USA) using DreamTaq PCR Master Mix (2X) purchased from Fermentas (Fermentas Canada Inc., Burlington Ontario, Canada) and nucleotides from IBA GmbH (IBA GmbH, Göttingen, Germany). 16S rRNA gene fragments were amplified with general bacterial primers U27F (5´-AGATTTTGATCTGCTGCGTAC-3´) and 1492L (5´-GGTTACCTTGTGCTGGTGTA-3´) [36]. Reaction mixtures (25 μl) contained 1 μl of bovine serum albumin (2 mg ml⁻¹, diluted 100 x), 0.1 μM of each DNA primer, 12.5 μl HotStartTaq PCR Master Mix (Qiagen, Venlo, The Netherlands) and 1 μl template DNA. The amplification conditions were as follows: initial denaturation (95°C for 10 min); 25 cycles of denaturation (95°C for 45 s), annealing (56°C for 45 s), extension (72°C for 1:30 min); final extension (72°C for 10 min). For all samples, a nested PCR procedure was followed. The primers for the amplification were 685 GC (5´-CCG GGG GCG GCC GCG GCC GCC GGG GGG GCA CGG GCC GAA GCA CGC GAA GAA CCT TAC-3´) and 1401L (5´-CGG TGT GTA CAA GAC CC-3´) [37] and a GC clamp was attached to the 5´ end of the forward primer to increase the separation of DGGE bands in the electrophoretic analysis. Each reaction (25 μl) contained 0.5 μl of acectamide (50%), 0.1 μM of each DNA primer, 12.5 μl HotStartTaq PCR Master Mix and 1 μl of template DNA. The amplification protocol was: an initial denaturation at 95°C for 10 min, 30 cycles of denaturation at 95°C for 1 min, annealing at 53°C for 1 min, extension at 72°C for 1:30 min, final extension at 72°C for 7 min, and cooling to 15°C. Negative controls without template DNA were included. Five μl of PCR products were analysed by electrophoresis on a 1.5% agarose gel with GelRed (Biotium, Hayward, CA, USA) as DNA staining agent. DGGIE was performed using the DCodeTM Universal Mutation Detection System (Bio Rad Laboratories, Hercules, CA, USA). PCR amplicons were loaded onto 6-10% (w/v) polyacrylamide gel in 0.5 × TAE buffer (20 mM Tris-acetate pH 7.4, 10 mM sodium acetate, 0.5 mM Na₂EDTA) using a linear denaturing gradient ranging from 40% to 58%. Electrophoresis was performed for 16 h at 80 V at 60°C in 1 x TAE buffer. The gels were silver-stained [38]. The images were analysed using the software package BioNumerics (Applied Maths, Sint-Martens-Latem, Belgium).

Statistics

The significance of the differences between treatments was assessed by 2-way ANOVA using the SPSS Statistics 17.0 (International Business Machine Corp., New York, USA). Normality was confirmed by Kolmogorov-Smirnov test and equality of variances was checked by the Levene test.

The data matrix of band (band positions and their corresponding intensities) abundance per sample was log₂(x + 1) transformed and a distance similarities matrix constructed using the Bray-Curtis index with the vegdist function in the vegan package in R version 2.11.1 [39]. Variation in bacterial composition among treatments/time was visually assessed with Principal Coordinates Analysis (PCO) using the cmdscale function in R and the Bray–Curtis distance matrix as input. Variation among treatments groups was compared using the Adonis function in vegan. The Adonis function is an analysis of variance with distance matrices using permutations that partitions distance matrices among sources of variation; in this case treatment and time. In the adonis analysis, the Bray-Curtis distance matrix of band composition was the response variable with treatment (high-Hg, low-Hg and blended 1:4) and time (day: 0 or 7) as independent variables. We also included the interaction term between treatment and time. The number of permutations was set at 999; all other arguments used the default values set in the function. Measured microbiological variables, namely biomass productivity (μgGdw⁻¹h⁻¹), arylsulfatase (nmol.gdw⁻¹h⁻¹), number of prokaryotic cells (cells.gdw⁻¹), number of bacteria (cells.gdw⁻¹), number of sulfate reducing bacteria (cells.gdw⁻¹), proportion of EUBP (EUBP abundance/ total prokaryote abundance (NTP)), proportion of SRB (SRBP abundance/ total prokaryote abundance (NTP) and Hg content (ngHg.mgdw⁻¹)) were fit in the PCO ordinations using the envfit function in vegan. The envfit function was also used to test for significant relations between these variables and the PCO ordinations of bacterial community composition using 999 permutations; all other arguments in the function were left as default.

Results

Sediment characterization

The surface sediment (0-0.05 m), used as low-Hg, was characterized by 60.2 ± 3.3% of fine particles (<0.63 mm) and 14.0 ± 0.1 ngHg.mgdw⁻¹. In the deeper high-Hg sediments (0.4-0.5 m) used as the source of mercury for the microcosm experiments, the percentage of fine particles was 59.2 ± 1.5% and the concentration of mercury was 98.9 ± 1.7 ngHg.mgdw⁻¹.

Extracellular enzymatic activity

The variation of arylsulfatase activity in the conditions of the experiment is represented in Figure 1. At the beginning of the experiment, arylsulfatase activity was the highest in samples from the upper sediment layer, taken as low-Hg (9.7 nmol.gdw⁻¹h⁻¹), and lowest in sediments from the 0.4-0.5 m, taken as the high-Hg (3.2
nmol.gdw⁻¹h⁻¹). In blended 1:4 sediment, the potential maximum hydrolysis rate (Hm) was intermediate (6.1 nmol gdw⁻¹h⁻¹) between both control sediments and differences between treatments were statistically significant (two-way ANOVA, p<0.05).

At the end of the experiment, the pattern of variation was similar but there was a statistically significant overall increase (two-way ANOVA, p<0.05) in activity in the low-Hg sediment (15.4 nmol.gdw⁻¹h⁻¹) and in blended 1:4 (11.4 nmol.gdw⁻¹h⁻¹), in relation to the beginning of the incubation. At the end of the experiment, arylsulfatase Hm in the high-Hg microcosms (4.22 nmol.gdw⁻¹h⁻¹) was not significantly different from the initial value.

### Bacterial biomass productivity

The variation of the rate of bacterial biomass productivity is represented in Figure 2. The highest activity rate was observed in the low-Hg (10.9 μgC.gdw⁻¹h⁻¹) and the lowest in the high-Hg (1.2 μgC.gdw⁻¹h⁻¹). In blended 1:4 sediment, biomass productivity was 7.8 μgC.gdw⁻¹h⁻¹. Differences between treatments were statistically significant (two-way ANOVA, p < 0.05) but differences between the beginning and the end of the microcosm experiments were not.

### Prokaryote abundance

#### Table 1: Abundance of prokaryote cells (Bacteria and SRB) ± SD (4 replicates) in mudflat sediments containing different concentrations of mercury, assessed at the beginning (T0) and at the end (T7) of 7-day microcosm experiments.

| Samples               | TPN1 (x 10⁹ cells.gdw⁻¹) | Bacteria | SRB2 | Bacteria/TPN | SRB/TPN |
|-----------------------|--------------------------|----------|------|--------------|---------|
| Low-Hg T0             | 2.2 ± 0.22               | 1.9 ± 0.21 | 0.5 ± 0.11 | 82.8 ± 1.44 | 23.8 ± 4.08 |
| Blended 1:4 sediment T0 | 1.3 ± 0.06               | 0.9 ± 0.09 | 0.4 ± 0.02 | 69.4 ± 5.31 | 32.6 ± 0.72 |
| High-Hg T0            | 1.0 ± 0.06               | 0.5 ± 0.04 | 0.2 ± 0.02 | 53.9 ± 5.93 | 20.0 ± 1.88 |
| Low-Hg T7             | 2.6 ± 0.16               | 2.0 ± 0.22 | 1.1 ± 0.22 | 76.2 ± 4.40 | 40.2 ± 7.39 |
| Blended 1:4 sediment T7 | 1.4 ± 0.08               | 1.0 ± 0.05 | 0.5 ± 0.04 | 73.1 ± 6.71 | 33.7 ± 4.19 |
| High-Hg T7            | 1.0 ± 0.06               | 0.7 ± 0.03 | 0.4 ± 0.10 | 62.6 ± 0.98 | 39.1 ± 10.18 |

TPN: Total number prokaryote; SRB: sulfate reducing bacteria

Total number of prokaryotes was the highest (2.2 ± 0.22 × 10⁹ cells.gdw⁻¹) in low-Hg, the lowest (1.0 ± 0.06 × 10⁹ cells.gdw⁻¹) in high-Hg and intermediate (1.3 ± 0.06 × 10⁹ cells.gdw⁻¹) in the blended 1:4 sediments (Table 1). During the 7-day incubation, the variation in total prokaryote abundance was not statistically significant in any of the treatments (Table 1). On average, Bacteria detected with the probes EUB338, EUB338II or EUB 338III accounted for 70% DAPI stained cells (total prokaryotes). At the beginning of the experiment, the
abundance of Bacteria was significantly higher (two-way ANOVA, p<0.001) in low-Hg (1.9 ± 0.21 × 10^9 cells.gdw⁻¹, 82.8 ± 1.44% TPN), than in high-Hg or blended 1:4 sediments. After 7 days of incubation, the values of total abundance of Bacteria detected with the probes EUB338, EUB338II or EUB 338III were not significantly different from the initial values, with exception of high-Hg (ANOVA, p<0.05). The abundance of Bacteria was still significantly higher in the low-Hg (2.0 ± 0.22 × 10^9 cells.gdw⁻¹, 76.2 ± 4.40% TPN) and lower in blended 1:4 sediments (1.0 ± 0.05 × 10^9 cells.gdw⁻¹, 73.1 ± 6.71% TPN) and in the high-Hg (0.7 ± 0.03 × 10^9 cells.gdw⁻¹, 62.6 ± 0.98% TPN).

SRB accounted for 30% of total prokaryote abundance, on average. At the beginning of the experiment, the proportion of SRB in relation to total prokaryotes was higher in low-Hg sediments than in sub-surface sediments. During the experiment, there was a significant increase in the proportion of SRB in all samples, with exception of blended 1:4 sediment (two-way ANOVA, p<0.001). At the beginning of the experiment, the abundance of SRB in low-Hg (0.5 ± 0.11 × 10^9 cells.gdw⁻¹; 23.8 ± 4.08% TPN) and blended 1:4 sediment (0.4 ± 0.02 × 10^9 cells.gdw⁻¹; 32.6 ± 0.72% TPN) was significantly higher than in the high-Hg (0.2 ± 0.02 × 10^9 cells.gdw⁻¹; 20.0 ± 1.88% TPN) (two-way ANOVA, p<0.001). During the 7-day incubation this pattern persisted but there was an increase in SRB abundance (Table 1).

Community structure

DGGE profiles bacterial 16S rRNA gene fragments revealed differences in the banding pattern between the high-Hg and the other two sediments treatments (low-Hg and blended 1:4 sediments) related with treatment (Adonis: F_{2,23} = 5.080, R² = 0.326, P<0.001), time (Adonis: F_{1,23} = 6.223, R² = 0.221, P<0.001) and the interaction between treatment and time (Adonis: F_{5,23} = 9.060, R² = 0.716, P<0.001). PCO ordination of DGGE profiles is represented in Figure 3. The first two PCO axes explain approximately 55% of the variability of the dataset. Generally, replicates from the same experimental treatment group closely, indicating a good reproducibility of the analysis.

**Figure 3**: Principal Coordinate Analysis (PCO) analysis of 16S rRNA fragments of Bacteria amplified from total community DNA extracted from four replicates of shallow low-Hg (○) deeper high-Hg (Δ) and blended 1:4 sediments (◊) in the beginning (T0: open symbols) and at the end (T7: solid symbols) of the microcosm experiments. Codes are Prod: Hg: Mercury concentration, Day (incubation time), Productivity (µgC.gdw⁻¹.h⁻¹); Aryl: arylsulphatase (nmol.gdw⁻¹.h⁻¹); NTP: Number of cells.gdw⁻¹, EUB: Number of Bacteria cells.gdw⁻¹; SRB: Number of sulfate reducing bacteria cells.gdw⁻¹ and EUBP: % EUB/NTP and SRBP: % SRB/NTP.

The first PCO axis (Figure. 3) reveals that low-Hg and blended 1:4 sediments were characterized by higher arylsulphatase (Aryl; envfit for 1st and 2nd axes: P=0.001), productivity (Prod; envfit for 1st and 2nd axes: P=0.001), total prokaryote abundance (NTP; envfit for 1st and 2nd axes: P=0.001), abundance of Bacteria (EUB; envfit for 1st and 2nd axes: P=0.001), abundance of sulfate-reducing bacteria (SRB; envfit for 1st and 2nd axes: P=0.004) and proportion of Bacteria relation to total prokaryote abundance (EUBP; envfit for 1st and 2nd axes: P=0.001). The high-Hg treatment was characterize by a significant dependence on Hg content and time (Hg and Time; envfit for 1st and 2nd axes: P=0.001). There were no significant associations between the proportion of sulfate-reducing bacteria in relation to total prokaryote abundance (SRBP) and the ordination of the 1st and 2nd axes.

**Discussion**

A microcosm approach, in which a mechanical disturbance was simulated by mixing sediments from the deeper highly contaminated and surface less-contaminated layers, was carried as a realistic approach to investigate the impact of a possible dredging on the structure and activity of bacterial communities, without the use of an exogenous mercury source.

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Hetetrotrophic activity

The descriptors of activity analysed in this work show that bacterial activity was inhibited by mercury contamination and that the levels of arylsulfatase and biomass productivity were inversely related with the concentration of mercury in the sediments. One of the effects of mercury is the inhibition of bacterial heterotrophic activity [40]. Therefore, mercury acute contamination may have an immediate impact on organic matter diagenesis in estuarine sediments.

The bacterial community in the blended 1:4 sediments displayed levels of activity that were intermediate between low-Hg and high-Hg sediments. However, the levels of activity observed in in blended 1:4 sediments cannot be fully explained by processes of conservative mixing of two original communities expressing high (low-Hg) and low (high-Hg) levels of activity. In fact, heterotrophic activity in the blended 1:4 sediments was below the expected (e.g., lower that the weighted average of the values of activity in the two contrasting sediment types), possible indicating that an inhibition at cellular level of the more active community of the upper layer sediments. The comparison of the values observed and those predicted by conservative mixture indicate that the inhibition was immediate (observed in T0) and the inhibition factors were of 28% for arylsulfatase and 13% for biomass productivity. During the incubation, there was a slight recovery of arylsulfatase (13% inhibition at the end of the experiment) but an intensification of the inhibition of biomass production (33% at the end of the experiment). The different pattern of variation of these two descriptors of heterotrophic activity is probably related with the organisms involved. Leucine incorporation is a direct measure of prokaryote biomass productivity, because only prokaryotes are able to take up leucine in the dissolved form. On the contrary, arylsulfatase, although mainly originating from bacteria, is also common in fungi and can be expressed in plants and animals [41] and can be used as an indicator of recovery of contaminated soils [42]. Considering that in sediments, organic matter is mostly available as polymers, extracellular enzymatic activity is the initial and limiting step in organic matter decomposition and recycling [43]. In estuarine and coastal sediments, organic sulfur corresponds to a major fraction of the available sulfur and arylsulfatase activity significantly contributes to sulfate supply [44] for assimilatory or dissimilatory (sulfate reduction) metabolic pathways. In addition, arylsulfatase in soils can be truly extracellular, meaning that a significant fraction may occur adsorbed to sediment particles [45]. Therefore, the inhibition of arylsulfatase activity may be a combined process of direct inhibition of the enzyme and decreased activity of the organisms producing it, some of which may resist and recover [40]. The decrease in bacterial biomass productivity indicates a decrease in prokaryote activity rate from which cell did failed to recover within the time-frame of the experiment.

Community structure

Communities of Bacteria in the deeper (high-Hg) and surface sediments (low-Hg) were structurally distinct, such as expected from a different degree of exposure to mercury [13] and different environmental conditions, namely sediment texture, availability or organic substrates and electron acceptors [46,47]. Also, the influence of plant roots may have contributed to the in situ shaping of the community structure [10]. In sediments chronically contaminated with mercury, there is a selective pressure towards the increase of the relative abundance of bacterial groups that are tolerant to the metal [15], capable of extracellular bioaccumulation [48] or that are, in some way, involved in processes of speciation and transference, such as sulfate-reducing bacteria [14,49,50]. However, in this work, the proportion of SRB in relation to total prokaryotes in the initial surface sediments (low-Hg) was higher than in deeper sediments (high-Hg). Changes in the structure of sediment bacterial communities during microcosm experiments are likely to [51]. The increase in the proportion of SRB in all samples at the end of the experiment is more likely to have been caused by sediment handling, confinement and oxygen depletion, which may have selected for anaerobes, than an effect of mercury contamination, since it occurred in control and test sediments. The decrease in the proportion of Bacteria, in relation to total prokaryotes, in the positive control and in test sediments, and the opposite trend in the negative control, may indicate a differential response of Bacteria and Archaea to confinement and mercury contamination. Significant effects of mercury contamination on archaeal sediment communities have been demonstrated [52]. In this study, Archaea seem to have been even more responsive than Bacteria, in terms of abundance.

At the beginning of the experiment, the communities in low-Hg, high-Hg and blended 1:4 sediments were different between each other. However, low-Hg and blended 1:4 sediments were more similar between each other than with the communities in the heavily contaminated sediments. Community structure in high-Hg changed during the experiment but at the end of the incubation, it was still different from all the other communities, indicating that the effect of mercury on the shaping of the community structure was stronger than other effects related with confinement or oxygen depletion. The structure of the community in the low-Hg sediment was more responsive to the incubation time and changed in a scattered way. Overall, communities in sediments with the lowest mercury load became different from the initial community in these sediments and more similar to the blended 1:4 community. The structure of the blended 1:4 community was the least variable during the incubation. Taking into consideration that the effects of mercury on bacterial activity were immediate, as inferred from the rates of arylsulfatase activity and leucine incorporation in the blended 1:4 sediments at the beginning of the experiment, community structure seems to be much less reactive to mercury than heterotrophic activity. The changes in community structure observed during the experiment are, therefore, more likely triggered by factors associated with confinement, to which the low-Hg (less subjected to the selective pressure of mercury) was more responsive.

Conclusion

The results of this work demonstrate that the contamination of sediments with realistic concentrations of mercury has a higher impact on bacterial activity than on the structure of bacterial communities. Events causing disturbance or re-suspension of contaminated sediments susceptible of mobilizing mercury accumulating in deeper layers to upper or surface layers will impact bacterial communities and therefore their contributions to the associated biogeochemical cycles. Ultimately, when associated to the re-suspension of sediment particles to the water column, this might have an impact at the ecosystem level.

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