Changes in the non-protein thiol pool and production of Dissolved Gaseous Mercury in the marine diatom *Thalassiosira weissflogii* under mercury exposure

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**Abstract**

Two detoxification mechanisms working in the marine diatom *Thalassiosira weissflogii* to cope with mercury toxicity were investigated. Initially, the effect of mercury on the intracellular pool of non-protein thiols was studied in exponentially growing cultures exposed to sub-toxic HgCl₂ concentrations. *T. weissflogii* cells responded by synthesizing metal-binding peptides, named phytochelatins (PCs), besides increasing the intracellular pool of glutathione and γ-glutamylcysteine (γ-EC). Intracellular Hg and PC concentrations increased with the Hg concentration in the culture medium, exhibiting a distinct dose-response relationship. However, considerations of the PCs-SH : Hg molar ratio suggest that also glutathione could be involved in the intracellular mercury sequestration. The time course of the non-protein thiol pool and Hg intracellular concentration shows that PCs, glutathione and γ-EC represent a rapid cellular response to mercury, although their role in Hg detoxification seems to lose importance at longer incubation times. The occurrence of a process of reduction of Hg(II) to Hg⁰ and subsequent production of dissolved gaseous mercury (DGM) was also investigated at lower Hg concentrations, at which the PC synthesis doesn’t seem to be involved. The significant (P<0.01) correlation between the cellular density in solution and the production of DGM suggests that this diatom is capable of directly producing DGM, both in light and dark conditions. This finding has been confirmed by the absence of DGM production in the culture media containing formaldehyde-killed cells. Finally, the relationship between these two different pathways of Hg detoxification is discussed.

**Keywords**: dissolved gaseous mercury; mercury; non-protein thiols; phytochelatins; phytoplankton; *Thalassiosira weissflogii*

1. Introduction

Mercury is one of the most dangerous trace metals because it does not have recognized biological activity and its presence is widespread in the aquatic environment. Hence, aquatic
microorganisms, which are the first levels of the food chain, have developed defence strategies to neutralize the toxic effects of this metal (Barkay et al., 2003; Perales-Vela et al., 2006). Many studies have demonstrated that phytoplankton species can respond to metal toxicity through the production of antioxidant compounds (Pinto et al., 2003) and intracellular metal-binding thiol peptides (Kawakami et al., 2006 and references therein reported). Glutathione and related peptides appear to be the major components of heavy metal detoxification in plants, algae and some yeast species. Glutathione is the main non-protein thiol in animals, plants and protists. It plays an important role in maintaining reducing conditions inside cells and in protecting plants from environmental stress, including oxidative damage and excess of xenobiotic organic compounds or heavy metals. The accumulation of heavy metals in marine microalgae induces the enzymatically-mediated synthesis of intracellular peptides, polymers of glutathione, named phytochelatins (PCs). PCs, with the structure of $(\gamma$-Glu-Cys)$_n$-Gly $(n=2-11)$, are thiol-containing peptides implied in heavy metal detoxification, because of their capability to bind metal ions inside the cells (Grill et al., 1985; Cobbett, 2000). In vitro experiments have shown that PCs protect metal-sensitive enzymes from inactivation and restore the activity of metal-poisoned enzymes (Kneer and Zenk, 1992). Several studies have revealed that PC synthesis is activated both in vivo and in vitro by a wide range of metal ions, including Cd$^{2+}$, Cu$^{2+}$, Pb$^{2+}$, Ag$^+$, Zn$^{2+}$ and Hg$^{2+}$ (Gekeler et al., 1988; Rauser, 1995; Zenk, 1996). A considerable amount of literature has been published on the induction of PCs in phytoplankton exposed to different metals like Cd, Pb, Zn, Cu (Ahner et al., 1995; Morelli and Scarano 2001, 2004; Rijstenbil and Wijnholds, 1996; Le Faucheur et al., 2006) but the detoxification of mercury by PCs has been scarcely documented (Howe and Merchant, 1992; Ahner and Morel, 1995).

In recent years a number of investigations have focused on a detoxification mechanism acting in microorganisms exposed to mercury, that is the ability of bacteria to reduce Hg(II) to
volatile Hg° by means of an enzymatic pathway (Barkay et al., 1991; Nakamura et al., 2001; Rolfhus and Fitzgerald, 2004; Barkay and Wagner-Dobler, 2005; Fantozzi et al., 2009). In contrast, so far, few studies report the formation of Hg° in phytoplanktonic algae (Ben-Bassat and Mayer, 1977; Mason 1995; Devars et al., 2000) and, in addition, this reduction mechanism is largely unknown. It is well known that elemental mercury (Hg°) plays a fundamental role in the biogeochemical cycle of mercury (Schroeder et al., 1989; Horvat et al., 2003) since it constitutes 90% of volatile forms of mercury in natural waters, named Dissolved Gaseous Mercury (DGM). These forms pass from the water into the atmosphere due to their low water solubility and high volatility. The increasing interest in the study on the occurrence of a biotic production of DGM in aquatic environments, suggests that this issue needs to be further investigated.

Field and laboratory studies suggest that, besides bacteria, phytoplankton can play an important role in the processes of the formation of DGM, through an indirect contribution due to the release of biogenic organic matter involved in the photochemical reactions of DGM production, as well as through a cellular direct reduction (Mason et al., 1995; Devars et al., 2000; Lanzillotta et al., 2004; Poulain et al., 2004).

The aim of the present work is to investigate the defence mechanisms against mercury stress in the marine diatom *Thalassiosira weissflogii* by following the pattern of the non-protein thiol pool as well as the production of DGM. In particular, we investigated the intracellular concentration of glutathione, γ-EC and PCs as a function of both external metal concentration and time of exposure. In addition, we measured the production of Hg° in a culture of the same diatom grown in a medium enriched with HgCl₂, in dark and light conditions, in order to evaluate the mercury production process acting in this diatom.

2. Materials and methods
2.1 Chemicals

All reagents were analytical grade: diethylenetriaminepentacetic acid (DTPA), reduced glutathione (GSH), γ-glutamylcysteine (γ-EC), cysteine and monobromobimane (mBrB) were from Fluka; 4-(2-hydroxyethyl)-piperazine-1-propane-sulfonic acid (HEPPS), tris (2-carboxyethyl) phosphine (TCEP), hydrogen peroxide (30% solution) and HgCl₂ were from Sigma; methanesulfonic acid (MSA) was from Merck; HCl, HNO₃ Suprapur grade, acetonitrile, trifluoroacetic acid (TFA), SnCl₂ and formaldehyde (40%) were from Carlo Erba. Solutions of mBrB, SnCl₂ and TCEP were prepared weekly. The solution of SnCl₂ (0.4 M) in 1.2 M HCl was purged with charcoal-filtered air for 1h in order to provide a mercury-free solution. All the reagents were stored in the dark at +4°C. Water was purified by a Milli-Q system (Millipore).

Seawater was collected in an uncontaminated area, 3 miles offshore from the Island of Capraia (Tyrrenian Sea, Italy), by a metal-clean technique, filtered through 0.45 μm membrane filters and stored in the dark at +4°C.

Membrane filters used throughout the experiments were from Millipore.

2.2 Culture conditions and molecular characterization

The marine diatom, Thalassiosira weissflogii (Grunow) Fryxell & Hasle (1977) used in this study (strain 1085/1 isolated from Gorleston-on-Sea, Norfolk, England in 1975) was obtained from the Culture Collection of Algae and Protozoa (CCAP), Dunstaffnage Marine Laboratory, UK (http://www.ccap.ac.uk). Stock cultures were grown in axenic conditions, in natural seawater enriched with the f/2 medium (Guillard, 1975) at one-fifth the reported trace metal concentration, at 21°C and fluorescent daylight (100 μmol photons × m⁻² × s⁻¹) in a 16:8 light-dark cycle. Exponential growth was maintained by inoculating cells into a fresh
sterilized medium, weekly. Cell counts were carried out by means of a Neubauer counting chamber under a microscope.

Since molecular clades of diatoms are often cryptic, with no or few morphological or life history traits that can be convincingly argued to be synapomorphies, we carried out a molecular characterization of the diatom strain used in this study. The DNA was isolated following the standard protocol of Sambrook et al. (1989), modified and optimized for the genomic DNA isolation from protists, as reported by Fokin et al. (2008). The SSU-rRNA gene, universally considered a good species-specific marker, was amplified by PCR using the universal eukaryotic forward primer 18S F9 5’-CTGGTTGATCCTGCCAG-3’ (Medlin et al., 1988) and the 18S R1513 Hypo reverse primer 5’-TGATCCTTYGCAGGTTC-3’ (Petroni et al., 2002). The PCR product was purified and directly sequenced in both directions. The SSU-rRNA gene sequence of the T. weissflogii strain used in this study is available from the GenBank/EMBL databases under the accession number FJ600728.

2.3 Incubation experiments

All the mercury incubation experiments were carried out using, as a culture medium, natural seawater enriched with the f/2 medium lacking the trace metal stock solution. Calculated volumes of the stock cultures of T. weissflogii, at the end of the logarithmic growth phase, were used as inoculum to obtain an initial cell density of $1 \times 10^6$ cells L$^{-1}$.

In a first set of incubation experiments, designed to evaluate the effect of mercury on the growth rate of T. weissflogii, 100 mL culture media were spiked with HgCl$_2$ to the final concentrations ranging from 5 to 750 nM. The cultures were allowed to grow for 6 days during the exponential phase and the growth was monitored by counting cells.

Two different experiments were carried out, with the aim of investigating the pattern of the non-protein thiol pool under mercury exposure. In a 2-day exposure experiment, 1-L cultures
were exposed to HgCl$_2$ concentrations ranging from 5 to 150 nM. At the end of the exposure (cell density was $1-2 \times 10^7$ cells $\text{L}^{-1}$), aliquots of 800 mL and 50 mL of each culture were used for the determination of the non-protein thiols and the intracellular mercury concentration ([Hg]$_{\text{intr}}$), respectively. In a 7-day exposure experiment, a 2-L culture was exposed to 150 nM HgCl$_2$ and, at selected time intervals, from 0 to 7 days, aliquots of 50 mL of the culture were sampled and used for the determination of the [Hg]$_{\text{intr}}$. Moreover, aliquots of the culture from 800 to 200 mL, depending on cell density, were sampled and used for the determination of the non-protein thiols. In the exposure experiments, a control culture (no Hg added) was always used.

The production of dissolved gaseous mercury (DGM) was measured in cultures of T. weissflogii during exponential growth. For this purpose, 500-mL of the culture medium was spiked with HgCl$_2$ to reach an initial concentration of 5 nM and left to stand for 3 days. Before cell addition, the concentration of total dissolved mercury was approximately 65% of the initial one. This procedure was chosen to avoid elevated abiotic DGM production occurring within the first days after mercury addition, as shown in preliminary experiments. After inoculum of T. weissflogii cells, two aliquots of 50 mL of the culture were sampled at 1 day time intervals and used for the measurement of the DGM production and for the determination of the cellular mercury concentration ([Hg]$_{\text{cell}}$), respectively. An additional experiment of DGM production was performed by using T. weissflogii cells treated with formaldehyde according to the following procedure. Mercury-treated cells from 50 mL of a culture at the 4$^{th}$ day of growth (cell density approx. $4-5 \times 10^7$ cell $\text{L}^{-1}$) were collected by filtration (1.2 $\mu$m membrane filters) and re-suspended for 10 min in a solution of 1.6% formaldehyde in seawater. Afterwards, the formaldehyde-killed cells were collected by filtration, re-suspended again in their growth medium and submitted to the measurement of DGM production.
2.4 Determination of total dissolved mercury

Total dissolved mercury concentration ([Hg]_{diss}) was determined in the culture medium whether inoculated or not inoculated with *T. weissflogii* cells, by using the method described elsewhere (Ferrara et al., 2001). In the former culture medium, the cells were removed by filtration (1.2 µm membrane filters) before the Hg measurement. A calculated aliquot of the sample was diluted with distilled water to a final volume of 25 mL, acidified with 100 µL of HNO₃ and photo-oxidized using a UV medium-pressure lamp (90W) for 5 minutes in an ice bath. Mercury was measured using the Atomic Absorption Spectrometer (AAS) Gardis-3, based on the dual gold amalgamation procedure, after adding 200 µL of the SnCl₂ solution and purging with mercury-free air for 3 min at a flow rate of 0.3 L min⁻¹.

2.5 Determination of cellular and intracellular mercury concentration

Mercury-treated cells were collected by filtration onto 1.2 µm membrane filters and used for the determination of the total cellular mercury concentration ([Hg]_{cell}). In order to determine the [Hg]_{intr}, the harvested cells were incubated for 10 minutes with 1 mM EDTA in seawater to remove the metal adsorbed to the cell surface, then rinsed extensively with natural seawater. The cells, whether rinsed with EDTA or not rinsed, were immediately placed in 1 mL of HNO₃ (0.14 M) in water and mixed with 1 mL of concentrated HNO₃ and H₂O₂ (2:1 v/v). The sample was digested at 45° C for 16 h. This mineralization procedure was validated by using a Standard Reference Material (T6) “Fresh Water Plankton”. The results of analysis on the Standard Reference Material was 0.173 ± 0.03 µg g⁻¹ DW, compared with that of 0.186 ± 0.04 µg g⁻¹ DW reported by JRC (Joint Research Centre) of the European Commission. A calculated aliquot of the mineralized sample was diluted with distilled water to a final volume.
of 25 mL, added with 200 µL of the SnCl_2 solution and assayed for mercury concentration by pre-concentration on a gold trap and AAS determination.

2.6 Determination of DGM production

Measurements of the DGM production under both dark and light conditions were accomplished using the experimental apparatus described in detail elsewhere (Fantozzi et al., 2009).

A 50 mL sample was transferred into a 100 mL glass Pyrex purging bottle, showing optical properties elsewhere described (Lanzillotta and Ferrara, 2001) and a good transmittance (85%) for wavelengths > 350 nm.

Prior to the determination of the DGM production, samples were purged for 2 hours in dark conditions in order to eliminate the original DGM content.

DGM production in darkness was obtained incubating the sample contained in the purging bottle for 20 min in the dark; DGM production under light conditions was recorded following the exposure of the purging bottle, containing the sample, for 20 min to the same fluorescent light used for culture growth (100 µmol photons × m^{-2} × s^{-1}).

The DGM produced in the sample was extracted under dark conditions by means of mercury-purified air, used as a carrier gas, and accumulated on a gold trap. Mercury was thermally desorbed heating the trap at 500 °C and determined by Atomic Fluorescence Spectrometry (Tekran 2500 – detection limit 5 × 10^{-4} pmol Hg), using pure argon as a carrier gas. The detection limit of the procedure was 0.05 pM, calculated on the basis of the three standard deviation of the blank. The instrument was calibrated using a 25 µL Hamilton gas-tight micro-syringe to inject elemental mercury saturated air from a mercury vapour generator, kept at a constant temperature (4 °C), onto the gold trap. Preliminary tests were performed to verify the period of incubation within which the DGM production was linear in time. An
incubation time of 20 minutes was selected to obtain a meaningful DGM amount and to be within the linear time range of DGM production.

All the experiments were performed at a constant temperature of 21°C and the purging bottle, together with the Teflon tubing, were pre-cleaned by acid washing every time before the experimental apparatus was involved in a new measurement cycle.

The DGM determinations were replicated 3 times.

2.7 Determination of the non-protein thiols

After incubation, the cells were collected by filtration onto 1.2 µm membrane filters, re-suspended in 1.5 mL of 0.1 M HCl / 5 mM DTPA, then disrupted by sonication (Sonopuls Ultrasonic Homogenizer, Bandelin) for 3 min with a repeating duty cycle of 0.3 s, in an ice bath. The cellular homogenate was centrifuged (20000 g, 45 min) and the supernatant was used for the determination of thiols. Glutathione, γ-EC and PCs were separated and quantified by High Performance Liquid Chromatography (HPLC) after derivatization with the fluorescent tag mBrB, by following the procedure described elsewhere (Morelli and Scarano, 2001), based on the method reported by Rijstenbil and Wijnholds (1996) with some modifications. Briefly, 400 µL of the sample were added to 200 µL of buffer (400 mM HEPPS / 5 mM DTPA, pH 9) and to 20 µL of 10 mM TCEP in order to reduce oxidized thiol groups. After 15 min of incubation, two successive reactions in the dark at 45°C for 15 min were carried out, following the addition of 40 µL of 10 mM mBrB and of 40 µL of 100 mM cysteine, respectively. Finally, 40 µL of 1 M MSA were added to stop the reaction. Analyses were performed on an HPLC system consisting of two Shimadzu LC-10AD pumps, a Rheodyne 7725 injection valve equipped with a 100 µL loop, a fluorescence detector (RF–10AXL, Shimadzu) set at 380 nm excitation wavelength and 470 nm emission wavelength, and an Alltech Alltima (5 µm, 250 mm × 4.6 mm) C-18 reverse-phase column. An acetonitrile
gradient in 0.1% TFA (from 10% to 12% for 15 min and from 12% to 28% for a further 40 min) was used at a flow rate of 1 mL min$^{-1}$. Standard PCs from *Silene vulgaris* (Friederich et al., 1998) were kindly provided by Prof. M.H. Zenk, Munich University (Germany), and were used to check the retention time of phytochelatin oligomers. PC quantification was obtained from the relationship peak area vs concentration of GSH standard solutions. The total cellular PC concentration was expressed as the sum of the $\gamma$-Glu-Cys units quantified in each chromatographic peak of phytochelatins.

3. Results

3.1 Effect of mercury exposure on the growth rate of *T. weissflogii*

The effect of mercury on the growth rate ($\mu$) of *T. weissflogii* was investigated by growing cells in culture media at increasing Hg concentrations (initial cell density $1 \times 10^6$ cell L$^{-1}$). The growth rate of the control culture was about $1.0 \pm 0.1$ doublings day$^{-1}$ ($n=3$). In the range of Hg from 5 to 500 nM the growth rate gradually decreased, reflecting the inhibition of growth under mercury exposure (Fig. 1). Exponential growth was observed in all the cultures during 6 day exposure, but the Hg addition lengthened the lag phase, as also reported by other authors for cultures of *Chlorella* (Ben-Bassat and Mayer, 1975). It was extrapolated that the 50% inhibition of the growth rate occurred at an initial [Hg] = 250 nM, whereas inhibitions lower than 20% occurred for [Hg] $\leq$ 150 nM. In order to avoid toxic effects during mercury exposure, we used well tolerated Hg dosages, never exceeding the dose of 150 nM in the exposure experiments.

3.2 Two-day exposure to mercury

The pattern of the non-protein thiol pool in response to mercury exposure was studied by monitoring the concentration of glutathione, $\gamma$-EC and PCs in cells of *T. weissflogii* exposed
for 2 days to increasing Hg concentrations, from 5 to 150 nM (Fig. 2). *T. weissflogii* cells responded to the Hg exposure by increasing the total level of the non-protein thiol pool. Glutathione was the major thiol, being always present at concentrations higher than those of γ-EC and PCs. Its intracellular concentration increased even at low Hg concentrations. At 150 nM Hg, the amount of glutathione was two-fold with respect to that found in the non-treated cells. The concentration of γ-EC was significantly less than that of glutathione both in the control and in Hg-treated cells, but its level increased with increasing Hg concentration in solution. Hg exposure also induced the synthesis of PCs but, under these experimental conditions, they were detectable at [Hg] ≥ 25 nM. The PC cellular pool increased by following a dose-response relationship until it reached the value of 673 ± 104 amol cell⁻¹ at 150 nM Hg. The cellular pool of peptides was composed mainly of PC₂ (85-100%), the remaining amount being polymerized as PC₃ (0-15%). The predominance of the pentapeptide and the inability to synthesize oligomers with n>3 were found in all the cultures, irrespectively of the Hg dose. Assays of intracellular Hg showed that the metal concentration ([Hg]ₘₐₜ) increased with the Hg exposure, exhibiting a trend similar to that of the PCs. Cellular concentration of thiol groups of PCs was similar to that of intracellular Hg, exhibiting a molar ratio PCs-SH : Hg close to 1. Since in vitro studies have shown that PC₂ binds Hg with a stoichiometry of two SH groups for one metal ion (Mehra et al., 1996), it seems that the amount of PCs synthesized in this diatom during a 2-day exposure is not sufficient to sequester intracellular mercury ions. The finding that cellular glutathione and, to a lesser extent, γ-EC increased in response to Hg exposure, can account for a role in the intracellular mercury sequestration, in addition to PCs.

### 3.3 Time course of the non-protein thiol pool and mercury accumulation
Exponentially growing cultures of T. weissflogii exposed to 150 nM Hg were assayed at time intervals for their intracellular concentration of Hg and non-protein thiols (Fig. 3). The time course of the [Hg]$_{intr}$ showed a rapid uptake of the metal, occurring during the first day of exposure, thereafter there was no further increase at longer exposures. The PC cellular concentration, after reaching a maximum value on the first day, decreased with exposure time, until halved at the 7$^{th}$ day of exposure. This finding indicates that PC synthesis occurs quickly, as soon as the metal is taken up by the cells, thereafter, the lowering of its concentration suggests the occurrence of a process of degradation and/or export, as reported by other authors for Cd-PCs complexes induced in the same diatom (Lee et al., 1996).

Glutathione assays showed a transient increase of its intracellular concentration in the Hg-treated cells compared to that measured in the control culture, occurring during the first 2 days of exposure. An increase of 65 and 137% was calculated on the 1$^{st}$ and 2$^{nd}$ day, respectively. At the end of the experiment, the glutathione level in the Hg-treated cells was restored to values similar to those of the untreated cells. A similar pattern was observed for the $\gamma$EC peptides, which exhibited an increase in the Hg-treated cells compared to the untreated ones of 145 and 103% on the 1$^{st}$ and 2$^{nd}$ day, respectively. In conclusion, the time course of the non-protein thiol pool and Hg intracellular concentration shows that PCs, glutathione and $\gamma$-EC represent a rapid cellular response to mercury. However, at longer incubation times, their role in Hg detoxification seems to lose importance. Since the [Hg]$_{intr}$ remained almost constant during the entire incubation time, and the PC concentration lowered, it can be hypothesized that part of the intracellular Hg initially sequestered by PCs, or possibly by glutathione and $\gamma$-EC, could be transferred to other, more stable intracellular ligands.

Similar incubation experiments carried out at lower Hg concentrations, at which the PC synthesis doesn’t seem to be involved, showed that the intracellular Hg concentration
followed a decreasing trend, starting from the beginning of exposure to longer incubation times. Thus, at \([\text{Hg}] = 5\, \text{nM}\), the \([\text{Hg}]_{\text{intr}}\) decreased from 20.6 ± 2.8 amol cell\(^{-1}\) on the 1\(^{st}\) day, to 6.6 ± 1.1 amol cell\(^{-1}\) on the 7\(^{th}\) day of exposure. This trend can be due, at least in part, to dilution by cell duplication, nevertheless the occurrence of a process of loss of Hg cannot be excluded. In the literature it has been reported that aquatic microorganisms, mainly bacteria but also eukaryotic phytoplankton, are capable of transforming ionic Hg to volatile Hg species, thus the existence of a similar process of Hg transformation could contribute, in our experimental conditions, to the lowering of the \([\text{Hg}]_{\text{intr}}\).

3.4 Production of DGM in cultures of \text{T. weissflogii}

The ability of the marine diatom \text{T. weissflogii} to produce volatile Hg species was assayed by carrying out direct measurements of DGM production in an exponentially growing culture of this diatom, previously spiked with mercury (\([\text{Hg}] = 5\, \text{nM}\)). The pattern of \([\text{Hg}]_{\text{diss}}\) in the presence and absence of cells, together with that of the Hg taken up by the cells (\([\text{Hg}]_{\text{cell}}\)), is reported in Fig. 4. The figure shows that the cell addition dramatically lowers the \([\text{Hg}]_{\text{diss}}\) in solution, concomitantly with an increase in cellular density (see insert). As expected due to cell growth, the fraction of Hg associated to cells (\([\text{Hg}]_{\text{cell}}\)) increases with incubation time. Nevertheless it can be calculated that, during the exponential growth phase, this amount is not sufficient to explain the loss of \([\text{Hg}]_{\text{diss}}\) in solution.

Measurements of DGM were performed both in the whole culture and in the culture medium after removal of cells by filtration, in order to isolate the biotic contribution to the mercury volatilization from the abiotic one, due to the culture medium. Samples were analyzed under dark and light conditions to compare the efficiency of the two DGM production processes.

Table 1 shows the values of DGM production recorded on day 4 of growth of the culture of \text{T. weissflogii}, by using both alive and formaldehyde-killed cells. The results show that a
meaningful DGM production occurred both under dark and light conditions. The DGM production of the culture of *T. weissflogii* with live cells was significantly higher than that measured in the culture medium alone, both in dark and light conditions. On the contrary, the DGM production of the culture with formaldehyde-killed cells exhibited values similar to those obtained after cell removal. These results clearly demonstrate the significant contribution of living cells in mercury volatilization. The DGM production in the culture medium was higher in the light compared to the dark, as expected from the contribution of the biogenic organic matter in photochemical reactions of Hg reduction (Costa and Liss, 1999; Lanzillotta et al., 2004). Our results also show that, in our experimental conditions, the contribution of the live cells to DGM production seems to be independent of the light, being 4.6 ± 0.8 pmol L\(^{-1}\) h\(^{-1}\) in the light and 4.5 ± 0.9 pmol L\(^{-1}\) h\(^{-1}\) in dark conditions.

In order to strengthen the previous findings, we examined the relationship between the percentage of total dissolved mercury transformed in DGM by cells in 1 h (%DGM) and the cellular density in solution, calculated at different times of growth of the culture of *T. weissflogii* (see Fig. 4). Fig. 5 A-B shows a positive and significant correlation between the %DGM and cellular density, both in light and dark conditions (p<0.01). Moreover, we examined the relationship between the same % DGM and the [Hg\(_{\text{diss}}\)] in the culture medium. In this case, no significant correlation was observed (Fig. 5 C-D). The strict dependence of the %DGM on the cellular density, and not on the [Hg\(_{\text{diss}}\)] in the culture medium, confirms the direct cellular contribution to the DGM production in the culture. Moreover, the similarity of the slopes of the regression lines in Fig. 5 A-B supports the finding that the direct contribution of living cells to DGM production was unaffected by the conditions of illumination. These findings are consistent with those of Devars et al. (2000) who found that the ability to volatilize mercury by the freshwater microalga *Euglena gracilis* was independent of light.
4. Discussion

The marine diatom *T. weissflogii* responded to mercury exposure with two distinct mechanisms: the increase of the non-protein thiol pool and the production of DGM.

Our data show that the mercury treatment (5-150 nM, 2 day- exposure) induced a general increase of the non-protein thiol pool: besides glutathione and γ-EC, which are constitutively expressed in the cell, HPLC analysis showed the occurrence of PC synthesis. Although it is well known that marine phytoplankton can synthesize PCs in response to a variety of metal ions (Ahner et al., 1995; Rijstenbil and Wijnholds, 1996; Morelli and Scarano 2001, 2004; Kawakami et al., 2006; Le Faucheur et al., 2006), systematic studies regarding their capability to synthesize PCs in response to mercury are lacking. Howe and Merchant (1992), in a study examining the ability of the green microalga *Chlamydomonas reinhardtii* to produce metal-binding peptides in response to Cd, Hg or Ag, reported that Hg-treated cells exhibited a transient but striking increase in glutathione levels, but were not able to accumulate measurable amounts of PCs. Recently, much more information has become available on the effects of Hg on the non-protein thiol pool in plants (Gupta et al, 1998; Iglesia-Turino et al., 2006; Israr et al., 2006; Rellan-Alvarez et al., 2006). Among these authors, general agreement on the involvement of glutathione in Hg detoxification can be observed. Only one paper (Gupta et al., 1998) reports that, besides glutathione, PCs can play a role in the Hg cellular sequestration in two species of aquatic plants.

Our findings on the time course of the non-protein thiol pool show that glutathione and related peptides (PCs and γ-EC) undergo a rapid synthesis followed by a slower decrease of their cellular concentration at longer exposure times. At the end of the exposure, only the level of PCs, but not that of glutathione and γ-EC, remained altered in the Hg-treated cells compared to the untreated ones. The restoring of glutathione to basal levels (comparable to those measured in the control culture) might imply the occurrence of a process of release of
this thiol. Accordingly, Tang et al. (2005) demonstrated an extracellular release of glutathione by *T. weissflogii* cells under copper stress. Our findings seem to suggest a mechanism in which the Hg taken up by the cells at the beginning of the exposure could form Hg-GSH complexes which might subsequently transfer the metal ion into the newly formed PCs in order to form more stable Hg-PCs complexes. These, in turn, could be released and/or degraded more slowly. The occurrence of a similar mechanism for Hg sequestration is supported by an *in vitro* study demonstrating that GSH can transfer Hg into PCs at increasingly longer chain lengths (Mehra et al., 1996). The initial formation of metal-glutathione complexes followed by a transfer to the metal-induced PCs has been also hypothesized to occur in *Phaeodactylum tricornutum* under Cd or Cu exposure (Morelli et al., 2002; Morelli and Scarano, 2004). In the present study, the substantial stability of the [Hg]$_{\text{intr}}$ concomitant with a decrease of the PC concentration, along with the exposure time, doesn’t exclude that other intracellular ligands might participate in the intracellular sequestration of the metal. In a recent paper, Kelly et al. (2007) reported that a number of eukaryotic algae were able to biotransform Hg(II) into $\beta$-HgS at varying degrees and to accumulate this metal species in the cell. Further studies are needed to clarify this issue.

Our data show alterations of the non-protein thiol pool at [Hg] > 5 nM, whereas at lower concentrations we demonstrated that *T. weissflogii* is capable of transforming mercury, added as HgCl$_2$, into volatile Hg species. Other authors have suggested that eukaryotic microorganisms, besides the prokaryotic ones, can reduce mercury, but only few authors measure DGM production directly. Ben-Bassat and Mayer (1978), Amyot et al. (1994) and Vandal et al. (1991) found a correlation between chlorophyll a concentration and Hg$^+$ formation rate suggesting that there is a link between productivity and Hg reduction. Mason et al. (1995) carried out measurements of DGM production in laboratory monocultures of a number of phytoplankton species, including *T. weissflogii*, and demonstrated their capability
of reducing Hg(II) to Hg°, although the rate of reduction was insufficient to account for the reduction rates observed in incubated field samples. The rate of DGM production measured by these authors in T. weissflogii (0.29 amol cell\(^{-1}\) d\(^{-1}\)) was comparable to that measured in the present study (2.6 amol cell\(^{-1}\) d\(^{-1}\)) at similar cellular density (5-7 × 10\(^7\) cell L\(^{-1}\)), taking into account the ten-fold higher [Hg] which we used.

Very little has been found in the literature on the mechanisms involved in the Hg reduction in eukaryotic microorganisms. Hg° production could involve cell surface reduction, similar to that found for other trace metals (Jones et al., 1987) rather than a gene encoded Hg resistance mechanism, as in the case of prokaryotic microorganisms. Ben-Bassat and Mayer (1977) isolated from crude extracts of the green alga C. pyrenoidosa an intracellular fraction (molecular weight < 1200 Da) responsible for Hg reduction, but its nature remains unknown. Taken together, our results show that T. weissflogii is able to activate a process of reduction of Hg(II) to Hg°, producing measurable amounts of DGM when exposed even at low Hg concentrations ([Hg] = 5 nM). At higher Hg concentrations ([Hg] = 10 -150 nM), the rate of DGM production seems to be insufficient to prevent Hg intracellular accumulation. In this case, the Hg accumulated by the cells would induce a general increases in the actual pool of glutathione and γ-EC, besides inducing an ex-novo synthesis of PCs.

References


Figure captions

**Figure 1.** Percentage of inhibition of growth rate ($\mu$, doublings day$^{-1}$) of *T. weissflogii* cultures under mercury exposure. Initial cell density = $10^6$ cells L$^{-1}$. Exposure time = 6 days. Different symbols refer to two independent experiments.

**Figure 2.** Patterns of glutathione (○), γ-EC (▼), PCs (●) and [Hg]$^{\text{intr}}$ (Δ) in *T. weissflogii* cells exposed for 2 days to increasing Hg concentrations. Initial cell density = $10^6$ cells L$^{-1}$. Standard deviations refer to duplicate experiments.

**Figure 3.** Time course of the intracellular Hg concentration and of the non-protein thiol pool in *T. weissflogii* cells exposed to 150 nM Hg for 7 days. PC concentration is expressed as the sum of the γ-Glu-Cys units. Initial cell density = $10^6$ cells L$^{-1}$. Standard deviations refer to duplicate experiments.

**Figure 4.** Time course of the total dissolved Hg concentration measured in the culture medium either not inoculated (▲) or inoculated with *T. weissflogii* cells (●), together with the cell-bound Hg concentration (○). Culture media contained 5 nM HgCl$_2$ and were let equilibrate for 3 days before inoculum. Standard deviations refer to triplicate experiments. Insert: growth curve of the *T. weissflogii* culture used for the experiment.

**Figure 5.** Correlation between the percentage of total dissolved mercury transformed into DGM by cells in 1 h (% DGM) vs. cellular density (A-B) or vs. [Hg]$^{\text{diss}}$ in the medium (C-D), measured in a culture of *T. weissflogii* exposed to mercury (culture conditions are reported in
the caption of Fig.4). All the DGM production rates are corrected for the abiotic production of the culture medium.
Figure 1
Figure 2
Figure 3
Figure 4
Figure 5

Mean percent DGM in light (A) and dark (B) treatments as a function of cellular density (10^6 cells L^-1) and [Hg]_diss (nM). A positive correlation was observed in light conditions (y = 3.15x + 0.35, r = 0.83*) and a negative correlation in dark conditions (y = 2.41x - 0.03, r = 0.88*).

Mean percent DGM in light (C) and dark (D) treatments as a function of [Hg]_diss (nM). A negative correlation was observed in light conditions (y = -0.45x + 1.65, r = -0.48) and a negative correlation in dark conditions (y = -0.29x + 0.99, r = -0.43).
Table 1. DGM production in cultures of *T. weissflogii* containing either live or formaldehyde-killed cells, as well as in the culture medium after removing cells. Cells were grown for 4 days in a culture medium with 5 nM HgCl$_2$. Cell density = 4-5 × 10$^7$ cell L$^{-1}$.

The experiment was carried out in duplicate.

<table>
<thead>
<tr>
<th></th>
<th>Light</th>
<th>Dark</th>
</tr>
</thead>
<tbody>
<tr>
<td>Culture with live cells (1)</td>
<td>8.7 ± 0.8</td>
<td>4.7 ± 1.8</td>
</tr>
<tr>
<td>Culture with killed cells (2)</td>
<td>4.6 ± 0.7</td>
<td>0.3 ± 0.2</td>
</tr>
<tr>
<td>Culture medium (3)</td>
<td>4.1 ± 0.9</td>
<td>0.2 ± 0.1</td>
</tr>
<tr>
<td>Live cells (1)-(3)</td>
<td>4.6 ± 0.8</td>
<td>4.5 ± 0.9</td>
</tr>
<tr>
<td>Killed cells (2)-(3)</td>
<td>0.5 ± 0.7</td>
<td>0.1 ± 0.2</td>
</tr>
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