Methylation of Mercury in Earthworms and the Effect of Mercury on the Associated Bacterial Communities

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Abstract

Methylmercury compounds are very toxic for most organisms. Here, we investigated the potential of earthworms to methylate inorganic-Hg. We hypothesized that the anaerobic and nutrient-rich conditions in the digestive tracts of earthworm’s promote the methylation of Hg through the action of their gut bacteria. Earthworms were either grown in sterile soils treated with an inorganic (HgCl2) or organic (CH3HgCl) Hg source, or were left untreated. After 30 days of incubation, the total-Hg and methyl-Hg concentrations in the soils, earthworms, and their casts were analyzed. The impact of Hg on the bacterial community compositions in earthworms was also studied. Tissue concentrations of methyl-Hg in earthworms grown in soils treated with inorganic-Hg were about six times higher than in earthworms grown in soils without Hg. Concentrations of methyl-Hg in the soils and earthworm casts remained at significantly lower levels suggesting that Hg was mainly methylated in the earthworms. Bacterial communities in earthworms were mostly affected by methyl-Hg treatment. Terminal-restriction fragments (T-RFs) affiliated to Firmicutes were sensitive to inorganic and methyl-Hg, whereas T-RFs related to Betaproteobacteria were tolerant to the Hg treatments. Sulphate-reducing bacteria were detected in earthworms but not in soils.

Introduction

Mercury (Hg) is a naturally occurring metal, which is released in the environment by numerous natural and anthropogenic processes [1,2]. Mercury is relatively stable in the atmosphere and can spread over the entire globe before returning to the earth’s surface. In soil, Hg is highly immobile and accumulates in the top layer, mainly by binding to organic matter especially to thiol groups [3]. Tipping et al. [4] reported a critical limit of 3.3 mg Hg kg⁻¹ organic matter corresponding to 0.13 mg Hg kg⁻¹ soil. At lower concentrations, it is assumed that there are no harmful effects on soils organisms. By applying this definition, 60% of 34 natural forest soils studied in Switzerland would exceed this critical limit [5].

Methylmercury compounds (CH₃Hg-R; methyl-Hg) are the most toxic Hg compounds for humans [6]. Methylation of Hg occurs through biotic and abiotic processes, although biotic processes are most important [7]. Sulphate-reducing bacteria (SRB) under anaerobic conditions seem to be of particular importance to methylate Hg [8–10]. In all SRB, enzymes that catalyze the reduction of sulphite to sulphide were found. Sulphite reductases enzymes (EC 1.8.99.3) consist of at least two polypeptides, encoded by the dissimilatory sulphite reductase genes dsrA and dsrB [11]. The presence of SRB in environmental samples is commonly analyzed by targeting the dsrAB genes [12–14].

The methylation and bioaccumulation of Hg have been well studied in aquatic ecosystems because consuming Hg-contaminated fish may lead to humans being poisoned. In contrast, studies of Hg, and in particular of methyl-Hg, in terrestrial ecosystems are rare. Over 90% of the invertebrate biomass in soils may consists of earthworms [15]. Earthworms play an important role in many soil-forming processes [16]. They also serve as a substantial food source for several higher organisms, such as birds and moles. Earthworms in forest soils are known to accumulate Hg and methyl-Hg [5,17]. Bioaccumulation factors (BAF) of Hg in earthworms were between 1 and 15 whereas BAF for methyl-Hg ranged from 15 to 191 [5]. The lipophilic property of methyl-Hg results in more efficient bioaccumulation than inorganic-Hg, which may explain why there are considerably higher BAF for methyl-Hg than for inorganic-Hg. Another possibility for the high BAF in earthworms is, that inorganic-Hg is methylated in earthworms, e.g. due to the activity of the microbiota in their digestive tracts. In particular, unique conditions prevail in the earthworm gut, which is anaerobic, with large amounts of easily available carbon, and these may favour the anaerobic growth of microorganisms [18,19].

In this study we tested this possibility and hypothesized that the conditions in earthworms’ digestive tracts favour the methylation of Hg by their gut-inhabiting bacteria. Earthworms (Lumbricus terrestris L.) were either grown in sterile soils treated with mercury(II)chloride (HgCl₂), with methylmercurychloride...
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Materials and Methods

Ethic statement
The research institute WSL has a general permit to use the area of their surrounding for scientific purposes. No endangered or protected species were involved in the experiment.

Experimental design
Laboratory experiments with Lumbricus terrestris L. and soils treated with inorganic-Hg, with methyl-Hg or without any Hg compounds were conducted (Figure 1). The experiments were performed in 3.3 L high density polyethylene boxes filled with 2 kg dw sterile soil. In the main experiment (Figure 1A) three approaches were used: a) soils without earthworms to examine the abiotic methylation and demethylation of Hg-species in the soil, b) soils without earthworms, but with a suspension obtained from rinsing the earthworms body surface to examine a possible Hg-methylation in the soils driven by microorganisms carried in by earthworms and c) soils with earthworms to examine the methylation of Hg in the earthworms. Sterile soil was necessary to test the ability of the microbes in the earthworm gut to methylate Hg. Otherwise, methyl-Hg in soils produced by microbes would have been taken up directly by earthworms without forming it in the gut of earthworms. Sterile soil is expected to have an impact on the "mutualistic digestive system", however, preliminary experiments showed that growth and mortality of earthworms grown in sterile soil were not affected (data not shown).

In a separate experiment, casts (excreted feces) were used as controls to study a possible methylation of Hg by organisms introduced into the soil by earthworms (Figure 1B). We analyzed earthworm cast also, because it offer very different environmental conditions for microbial growth than the surrounding soil [20].

The experimental soil was collected in a forest close to the Swiss Federal Research Institute WSL, Birmensdorf. Soil properties (pH, C/N, clay-silt-sand content) were determined according to Fal [21] before the soils were treated. The soil was dried at 105 °C for two days, sieved (4 mm), homogenized and autoclaved three times. Soil aliquots were incubated for three days on petri dishes containing MMN-Agar media to test the sterility of the soils. After autoclaving, soil aliquots were pooled to three samples for analyses of mercury. Soil aliquots were removed (iii). Immediately after removing the earthworms a cast was removed (iv). In a separate experiment (B), casts (excreted feces) were used as controls to study a possible methylation of Hg by organisms introduced into the soil by earthworms. Earthworms were incubated for one week in soils treated with inorganic-Hg (i) before they were placed into new boxes containing soils treated with Hg (ii). After three days, the earthworms were removed (iii). Immediately after removing the earthworms a cast sampling period has been started for 28 days (iv).

Figure 1. Experimental design. Two experimental assays (A and B) were performed: In the main experiment (A) sterile soil was either treated with inorganic Hg (+inorganic–Hg), methyl-Hg (+methyl–Hg) or without Hg (−Hg) (ii). The soils were incubated abiotically, with earthworms or with a earthworm rinsing suspension for 30 days in the dark at 15 °C (iii). At the end of incubation, the earthworms were removed from the soil and the Hg and methyl-Hg concentrations in the soil and earthworms were determined (iii). In a separate experiment (B), the casts (excreted feces) were used as controls to study a possible methylation of Hg by organisms introduced into the soil by earthworms. Earthworms were incubated for one week in soils treated with inorganic-Hg (i) before they were placed into new boxes containing soils treated with Hg (ii). After three days, the earthworms were removed (iii). Immediately after removing the earthworms a cast sampling period has been started for 28 days (iv).

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and the water loss was compensated by adding sterilized Milli-Q water.

L. terrestris was chosen as the model organism as it is very abundant in soils of deciduous forests in Switzerland [17] and the fraction of methyl-Hg compared to the total-Hg in its tissue is high [5]. Juvenile earthworms were obtained from a commercialsource (National Bait Inc., Canada). Before incubation of the earthworms in sterile soils, the earthworms were grown in the untreated collected forest soil (not sterilized, not Hg treated) for six weeks for adapting the earthworms and their gut-inhabiting bacteria to the new condition. Thereafter, the earthworms were kept in boxes with several layers of moist filter papers without feeding for 6 days at 15 °C to let them empty their guts. To clean the body surface of
the earthworms and to obtain a suspension of mucus, soil and microorganisms adhering to the earthworm’s body surface, each individual was rinsed in a 0.8% NaCl solution. Thereafter, six earthworms were taken and pooled to three samples for analysing the initial total and methyl-Hg concentrations. The earthworms contained about 0.1 mg Hg kg$^{-1}$ dw and 5.1 µg methyl-Hg kg$^{-1}$ dw (4.9% methyl-Hg of total-Hg) at the beginning of the experiments.

In the experiments with earthworms, six individuals were put into each box. Equal experiments were conducted without earthworms and with a suspension obtained from earthworm surfaces (Figure 1A, ii). Freeze-dried lettuce was used as a food source, and 0.5 g was added to each box in all experiments (also in experiments without earthworms). The boxes were incubated at 15°C in the dark for 30 days. We assumed that the soil treatment should not harm the earthworms according to Ernst and Frey [22] and Lock and Janssen [23]. All experiments were performed in four replicates. The soils were sampled at the beginning and at the end of the experiments. The soil in each box was totally mixed before and after incubation and at each time point several aliquots were taken and pooled together. At the end of experiments the earthworms were starved for six days on several layers of moist filter paper until their gut contents had emptied (Figure 1A, iii). The six earthworms per each box were pooled (resulting in n = 4 per treatment), placed in liquid nitrogen and stored at −20°C until further treatment. Before chemical and microbial analyses, the frozen earthworms were lyophilized and milled.

To study and collect casts, ten earthworms were incubated as described above in soil treated with inorganic-Hg for one week (Figure 1B, i). Thereafter, the earthworms were placed into new boxes containing soil treated with inorganic-Hg (Figure 1B, ii) and after three days, they were removed (Figure 1B, iii). The earthworm-free boxes were kept in the dark with constant water content for 28 days. To observe a potential ageing effect, subsamples of earthworm casts were sampled (about 500 mg) at the day of removing earthworms (cast was excreted between 0 and 3 days ago), after 2, 7, 15 and 28 days (Figure 1B, iv). The collected samples were stored at −20°C. The samples were lyophilized, milled and stored in the dark until analysis. The cast experiment was investigated in four replicates.

Mercury analyses
The total-Hg (inorganic + organic Hg compounds) and methyl-Hg concentrations were measured in all samples. The total-Hg concentrations in the samples were determined by using an atomic absorption spectrophotometer according to the manufacturer’s instructions (Advanced Hg Analyser; AMA 254, Altec s.r.l., CZ). The methyl-Hg concentrations were determined by Gas Chromatography-Atomic Fluorescence Spectroscopy (GC-AFS) according to Liu et al. [24]. The accuracy of the total-Hg and methyl-Hg analyses was checked against certified reference materials (TORT-2 for biotic and ERM CC580 for soil samples) and its recovery ranged between 95 and 106% for Hg and between 83 and 112% for methyl-Hg.

DNA extraction and PCR of 16S rRNA and dsrA genes
Genomic DNA in the earthworms and soils was extracted using a modified bead-beating method described in Frey et al. [25]. Approximately 500 mg of earthworm tissue or soil sample were processed with a BioFastPrep system (ThermoSavant). The extracted DNA was quantified with Pico Green (Invitrogen, Carlsbad, CA, USA) and stored at −20°C. DNA aliquots extracted from earthworm and soil samples (5 ng µl$^{-1}$) were pre-treated with BSA at 95°C for 4 min to remove PCR inhibitors. Bacteria in earthworms were amplified by a polymerase chain reaction (PCR) targeting the 16S rRNA genes similarly to that described in Frey et al. [26]. Twenty ng of pretreated DNA was added to 20 µl PCR reaction mix containing 1x PCR buffer, 0.5 mM MgCl$_2$, 400 µM dNTP, 0.6 mg ml$^{-1}$ BSA and 0.05 U µl$^{-1}$ Hot star Taq polymerase (Qiagen), 0.2 µM of the forward primer 27F (5'-AGAGTTTGATCMTGGCTCAG-3') and of the reverse primer 1378R (5'-CGGTGTGTA-CAAGGCCCCGGGAAACG-3') were prepared. For T-RFLP analyses the forward primers were fluorescently (FAM) labelled. The PCR reactions were conducted in a Veriti Thermal Cycler (Applied Biosystems, Foster City, USA) and were started by an initial denaturizing step for 15 min at 95°C, followed by 35 cycles of the following steps: 95°C for 45 s, 51°C for 45 s and 72°C for 2 min. The reaction was finished by an extension step for 5 min at 72°C.

SRB were analyzed in soil and earthworm samples using for DNA amplification primers targeting the dsrA subunit [13] followed by agarose gel electrophoresis similar as described by Yu et al. [27]. Two separate PCR were conducted using two different reverse primers. We used the forward primer DSR1F (5’-TGTAGTTGACTGACGATC-3’) and the reverse primers DSR4R (5’-GCGTGTGTA-CAAGGCCCCGGGAAACG-3’) described by Santillano et al. [14]. The PCR reagents were similar to those used previously for the 16S rRNA genes. After an initial step at 94°C for 15 min, 42 cycles of the following steps were performed: 94°C for 45 s, 55°C for 1 min and 72°C for 2 min. The reaction was finished at 72°C for 5 min and the PCR products were verified by agarose electrophoresis and subsequently analysed under UV illumination.

T-RFLP profiling of total bacterial community in earthworms
16S rRNA amplicons were digested with 0.2 U MspI restriction enzymes according to the manufacturer’s recommendations (Thermo Fischer Scientific, Waltham, USA). The digested DNA was purified with the Montage SEQ$^{\text{TM}}$ Sequencing Reaction Cleanup Kit (Millipore, Billerica, USA) according to the manufacturer’s instructions. For capillary electrophoresis, 1 µl of the purified digestion product was mixed with 12.9 µl HiDi formamide (Applied Biosystems) and 0.1 µl of ROX500 DNA fragment length standard (Applied Biosystems) before heating at 95°C for 2 min. T-RFLP profiles were performed using the ABI Genetic Analyzer 310 (Applied Biosystems) and then analyzed using the software GeneScan V3.1. and Genotyper V2.5. (Applied Biosystems) according to Frey et al. [26]. The size and

### Table 1. Soil properties and Hg and methyl-Hg concentrations (mean ± SD) in the soil used in the laboratory experiments.

<table>
<thead>
<tr>
<th>Soil type</th>
<th>pH</th>
<th>C/N</th>
<th>Clay [%]</th>
<th>Silt [%]</th>
<th>Sand [%]</th>
<th>Hg tot (mg kg$^{-1}$)</th>
<th>Methyl-Hg (µg kg$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cambisol</td>
<td>7.0</td>
<td>10.8</td>
<td>25</td>
<td>39</td>
<td>36</td>
<td>0.09 ± 0.00</td>
<td>0.32 ± 0.16</td>
</tr>
</tbody>
</table>

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relative abundance were defined for peaks between 50 and 500 bps by applying a threshold value of 100 fluorescence units. The data for the T-RFLP analyses were standardized by calculating the relative abundance of each T-RF as described by Zumsteg et al. [20].

Cloning and sequencing of 16S rRNA and dsrA genes

One clone library was generated from the 16S rRNA genes collected from earthworms to study the composition of the total bacterial communities, and one clone library was performed with dsrA genes from earthworms. For the clone libraries, the DNA extracted from earthworms from all Hg treatments (soil-Hg, soil+inorganic-Hg, soil+methyl-Hg) was pooled for the PCR of 16S rRNA and dsrA genes separately, similar as described before except for using unlabelled primers. The PCR products were purified with Montage PCR Centrifugal Filter Devices (Millipore) and quantified using UV-VIS spectroscopy (NanoDrop Spectrophotometer ND-1000, Wilmington, USA).

The amplified and purified 16S rRNA and dsrA genes were cloned into pGEM-T Easy Vectors according to the manufacturer’s instructions (Promega, Wisconsin, USA). We selected 400 colonies (384 for 16S rRNA genes and 16 for dsrA genes) and conducted PCR using vector-specific primers (M13F and M13R) as described by Frey et al. [29].

T-RFLP analysis of the colony PCRs was conducted as described above. The T-RFLP profiles of the clones were compared with the profiles of the environmental samples (earthworms). By overlapping the whole community profile with the clone profile, the clone could be assigned to a precise fragment size category. Clones yielding the selected fragments of interest (dominant T-RFs or T-RFs which were affected by the three soil Hg treatments) were sequenced. In total, 136 clones (120 clones for 16S rRNA and 16 clones for dsrA genes) were sequenced using the 3730XL DNA sequencer (Applied Biosystems) and with the forward primer 27F (5’-AGAGTTTGATCMTGGCTCAG-3’) and the reverse primer 907R (5’-CTACGGGCTACCTTGTTACGACT-3’) for 16S rRNA genes, or the primer SP6 (5’-ATTAGTTGACACTATAG-3’) and T7 (5’-TAAACGACTCACTATAGGG-3’) for dsrA genes. The sequences were then checked and manually edited in the software BioEdit (V.7.1.3.0 by Tom Hall), chimera checked (http://comp-hio.anu.edu.au/bellerophon/bellerophon.pl) and a BLAST search was conducted in the NCBI database (http://www.ncbi.nlm.nih.gov) and the ribosomal database project (rdp.cme.msu.edu). The 16S rRNA gene sequences were deposited in Genbank under accession numbers between JX183735 and JX183794, the dsrA gene sequences were aligned under the accession numbers between JX461240 and JX461242.

Sequences were aligned with the ClustalW sequence alignment in BioEdit. Phylogenetic trees were calculated using Bayesian inference with the program MrBayes (ver. 3.2) [30] and the LG+G model. A Markov chain Monte Carlo simulation was run for 2,000,000 generations. Trees were visualized using the software FigTree (ver. 1.3.1).

Statistical analyses

The statistical analyses of total-Hg and methyl-Hg concentrations in the samples were performed with the program STATISTICA (StatSoft, Tulsa, USA). Calculations for statistic significance (p values) of total and methyl-Hg concentrations in soils and earthworms from the different experimental approaches and in the cast experiment were performed using the Kruskal-Wallis test (p≤0.05). The effects of the three soil Hg treatments on the abundance of particular T-RFs were tested by ANOVA and Post-hoc Tukey-HSD test using STATISTICA (StatSoft). A Bray Curtis similarity matrix [31] from square-root-transformed T-RFLP data was calculated and Principal Coordinate Analysis (PCoA) was performed to estimate the Hg treatments (soil-Hg, soil+inorganic-Hg, soil+methyl-Hg) on the bacterial community structure using the software Primer 6 v.6.1.13 and Permanova v.1.0.9 [Perimer-E, UK]. The significance of the effects of Hg treatments on the community structures (T-RFLP profiles) was tested by performing Permutational MANOVA analyses.

Results

Hg and methyl-Hg contents

After the incubation, the concentrations of total-Hg in soils were not significantly different (p<0.05) from the initial concentrations in all experiments. Before incubation, the methyl-Hg concentration in soils treated with methyl-Hg was similar as the total Hg concentration (data not shown). The concentrations of total-Hg and methyl-Hg in soils at the end of the experiments with and without an earthworm rinsing suspension (abiotic experiments) were similar. However, the concentrations of methyl-Hg in soils treated with methyl-Hg decreased by about 60% during the experiments (data not shown).

The concentrations of total-Hg and methyl-Hg in the earthworms increased in all experiments (Table 2). The initial concentrations of methyl-Hg in earthworms was about 5 μg methyl-Hg kg⁻¹ soil dw. At the end of the experiments, the concentrations of methyl-Hg in earthworms were significantly higher in experiments with inorganic-Hg (75 μg methyl-Hg kg⁻¹ soil dw) and with methyl-Hg (16 613 μg methyl-Hg kg⁻¹ soil dw) than without Hg (11 μg methyl-Hg kg⁻¹ soil dw) (p = 0.035 and 0.001 respectively; Table 2). The concentrations of total-Hg in the casts were about 25% lower than in the surrounding soils, whereas the concentrations of methyl-Hg in the casts were similar to the concentrations in the soils but did not change significantly (p<0.05) over time (Table 2).

Bacterial community profiling

The bacterial community structures in the earthworms were strongly influenced by the Hg treatments (Table 3; Figure 2). Bacterial T-RFLP profiles in earthworms grown in soils not treated with Hg are marginally significantly different from those in earthworms grown in soils treated with inorganic Hg (p = 0.06; Permutational MANOVA) and were significantly different to those in earthworms from soil treatments with methyl-Hg (p = 0.03; Permutational MANOVA). The T-RFLP profiles between the inorganic and methyl-Hg treatments were different but not on a significant level (p = 0.12). The numbers of T-RFs in earthworms grown in soil treated with methyl-Hg were lower (26 ±5) as in experiments with Hg (28 ±5) and without Hg (41 ±6; data not shown). Eight T-RFs (11%) in earthworms were significantly decreased in experiments with inorganic-Hg compared to experiments without Hg (Table 3). In methyl-Hg treated soils, eleven T-RFs (16%) in earthworms were significantly less and two T-RFs (3%) significantly more abundant compared to experiments without Hg (Table 3). Overall, the bacterial T-RFLP profiles in earthworms grown in soils without Hg were clearly different from those in earthworms grown in soils with methyl-Hg, according to PCoA analyses (Figure 2).

Bacterial community composition

The T-RF profiles of the earthworm samples were compared to the T-RF profiles of the clones. At least five clones yielding the selected fragments of interest (dominant T-RFs or T-RFs which
were affected by the three soil Hg treatments; Figure 3) were sequenced. Sequence analysis confirmed that cloned sequences were similar and always matched the same closest relatives. The cloned sequences were related to the phyla Firmicutes, Actinobacteria and Proteobacteria (Figure 4). Eight T-RFs (119 bp, 145 bp, 151 bp, 153 bp, 166 bp, 291 bp, 298 bp and 319 bp) belong to the phylum Firmicutes. The cloned sequence of T-RF 151 bp (clone SEst1) was most similar (99%) to a Brevibacillus sp. strain KZ17 (FJ481959) isolated from wasps. In earthworms, the abundance of this T-RF (151 bp) tended to decrease in experiments with inorganic-Hg and methyl-Hg (Figure 3). Cloned sequences of the T-RFs 153 bp (clone SEs2) and 166 bp (clone SEs3) were closely related to each other, and were most similar (97%) to a Bacillus sp.

Table 2. Total-Hg and methyl-Hg concentrations (mean ± SD; n = 4) in soils, earthworms and in casts.

<table>
<thead>
<tr>
<th>Soil*</th>
<th>Treatment</th>
<th>Hg tot [μg kg⁻¹]</th>
<th>Methyl-Hg [μg kg⁻¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td>−Hg</td>
<td>without</td>
<td>101±5.0</td>
<td>0.31±0.06</td>
</tr>
<tr>
<td>−Hg</td>
<td>with</td>
<td>92±0.2</td>
<td>0.34±0.04</td>
</tr>
<tr>
<td>+inorganic-Hg</td>
<td>without</td>
<td>6 145±547</td>
<td>0.81±0.26</td>
</tr>
<tr>
<td>+inorganic-Hg</td>
<td>with</td>
<td>5 640±401</td>
<td>1.26±0.89</td>
</tr>
<tr>
<td>+inorganic-Hg</td>
<td>with rinsing suspension</td>
<td>6 035±523</td>
<td>1.00±0.55**</td>
</tr>
<tr>
<td>+methyl-Hg</td>
<td>without</td>
<td>700±18</td>
<td>250±37</td>
</tr>
<tr>
<td>+methyl-Hg</td>
<td>with</td>
<td>594±52</td>
<td>301±267</td>
</tr>
<tr>
<td>+methyl-Hg</td>
<td>with rinsing suspension</td>
<td>682±4</td>
<td>288±164</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Earthworms*</th>
<th>Treatment</th>
<th>Hg tot [μg kg⁻¹]</th>
<th>Methyl-Hg [μg kg⁻¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td>−Hg</td>
<td>218±57</td>
<td>11±2</td>
<td></td>
</tr>
<tr>
<td>+inorganic-Hg</td>
<td>24 271±5 531</td>
<td>75±35</td>
<td></td>
</tr>
<tr>
<td>+methyl-Hg</td>
<td>15 582±3 085</td>
<td>16 613±7 770</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Casts***</th>
<th>Age [d]</th>
<th>Hg tot [μg kg⁻¹]</th>
<th>Methyl-Hg [μg kg⁻¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–3</td>
<td>5 248±864</td>
<td>1.0±0.2</td>
<td></td>
</tr>
<tr>
<td>2–5</td>
<td>4 563±615</td>
<td>1.5±0.3</td>
<td></td>
</tr>
<tr>
<td>7–10</td>
<td>4 630±622</td>
<td>1.2±0.5</td>
<td></td>
</tr>
<tr>
<td>15–18</td>
<td>4 165±821</td>
<td>1.2±0.8</td>
<td></td>
</tr>
<tr>
<td>28–31</td>
<td>4 619±149</td>
<td>0.9±0.4</td>
<td></td>
</tr>
</tbody>
</table>

*Sampled after 30 days of incubation; **n = 3; ***Soil treated with inorganic-Hg.

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Figure 2. PCoA of bacterial T-RFLP profiles in earthworms. The influence of the Hg treatment (symbols) was estimated with Permutational MANOVA.
doi:10.1371/journal.pone.0061215.g002

Table 3. Numbers and percentages of total numbers of T-RFs (in parentheses) in earthworms after 30 days which were significantly different (decreasing or increasing) between the soil treatments (−Hg, +inorganic-Hg; +methyl-Hg).

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Decreased</th>
<th>Increased</th>
</tr>
</thead>
<tbody>
<tr>
<td>−Hg versus +inorganic-Hg</td>
<td>8 (11)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>−Hg versus +methyl-Hg</td>
<td>11 (16)</td>
<td>2 (3)</td>
</tr>
<tr>
<td>+inorganic-Hg versus +methyl-Hg</td>
<td>0 (0)</td>
<td>2 (3)</td>
</tr>
</tbody>
</table>

Total number of T-RFs in earthworms: n = 71.
doi:10.1371/journal.pone.0061215.t003
strain KZ_AaIM_Mm2 (GU726177) isolated from mosquitoes. The abundance of both T-RFs in earthworm samples was significantly lower in experiments with inorganic-Hg and methyl-Hg than in those without Hg. Overall, T-RFs affiliated to the phylum Firmicutes seem to be sensitive to inorganic-Hg and methyl-Hg.

Cloned sequences of three T-RFs (164 bp, 278 bp and 279 bp) are phylogenetically affiliated to Actinobacteria (Figure 4). Cloned sequences of T-RF 278 bp (clone SEs4) were most similar (98%) to a Microbacterium sp. strain YT0620 (AB376082). T-RF 278 bp was significantly more abundant in earthworms grown in soils without Hg than in soils treated with inorganic-Hg or methyl-Hg.

We found four T-RFs belonging to Alphaproteobacteria (396 bp), Betaproteobacteria (450 bp, 482 bp) or Gammaproteobacteria (87 bp) that were more abundant in earthworms from soils treated with inorganic-Hg or methyl-Hg than in soils without Hg. The T-RF of 450 bp was of special interest because it was clearly more abundant in earthworms grown in soils treated with methyl-Hg, where it was the most dominant T-RF, than in soils without Hg or with inorganic-Hg. Cloned sequences of T-RF 450 bp (clone SEst3) were most similar (99%–100%) to an environmental clone (AJ543436) affiliated to Acidovorax sp. and to a clone LT 1 (FJ214171) related to Verminephrobacter sp. which were both obtained from earthworms.

We also tested the DNA extracts obtained from earthworm and also from soils to detect the presence of SRB with specific PCRs targeting the dsrA genes. In all earthworm but not in soil samples (+Hg, +inorganic-Hg, +methyl-Hg) weak PCR products appeared (data not shown). Cloned sequences were most similar to the dsrA subunit of Desulfovibrio vulgaris strain RCH1 (similarity between 97 and 99%; CP002297).

Discussion

This study demonstrates that earthworms provide suitable conditions for the methylation of inorganic-Hg. In the biotic experiments with earthworms, the concentrations of methyl-Hg in soils are expected to decrease with time due to the strong uptake and retention of methyl-Hg in earthworms, and due to the decrease of methyl-Hg concentrations in soils measured in experiments without earthworms. Therefore, in these experiments, the concentrations of methyl-Hg in the soils should decrease (initial soil concentration minus loss due to accumulation by earthworms and evasion/demethylation of methyl-Hg) during incubation. This was, however, not observed here. Instead, the concentrations of methyl-Hg in soils were slightly higher (1.2 µg methyl-Hg kg⁻¹ soil dw) at the end of the experiments. Hinton and Veiga [32] showed that earthworms are potentially able to methylate Hg. They dissolved Hg⁰ in tannic acid, mixed it with silica sand and paper substrate, and let earthworms (Eisenia fetida) grow in it. They analyzed the methyl-Hg concentrations and found a ratio of methyl-Hg to total-Hg up to 160 times higher in earthworm tissues than in the acid and the substrate. In our study, we have chosen, in contrast, a natural habitat (soil) for the earthworms and Hg²⁺, the most dominant Hg species in natural soils [33]. We were able to show, that inorganic-Hg is methylated in the earthworm itself, and not by the bacteria introduced into the soils by the earthworms. This finding is based on the following evidence. Firstly, the earthworms contained about six times higher concentrations of methyl-Hg if they grew in soils treated with inorganic-Hg than in soils without Hg. Secondly, the concentrations of methyl-Hg in earthworm casts were similar to the concentrations in the soils and did not change over time. Thirdly, the earthworm rinsing solution did not enhance the methyl-Hg concentrations in the inorganic-Hg treated soils and thus, microorganisms introduced into the soil (cast/rinsing solution) did not methylate Hg in the soils. Uncertainties of the measurements of methyl-Hg occur, since it...
renders difficult to separate chromatographically methyl-Hg from inorganic-Hg in samples with very high Hg concentrations. Another uncertainty is that methyl-Hg can be formed artificially during the analytical procedure as pointed out in [34]. Despite these uncertainties we were able to demonstrate evidence that inorganic-Hg is methylated in earthworms itself, and not by the bacteria introduced into the soils by earthworms.

Mercury methylation has been observed in terrestrial invertebrates. Limper et al. [35] tested the ability of termites (Mastotermes darwinesis) to methylate inorganic-Hg. They found an in vivo methylation of Hg in termites and identified SRB (Desulfovibrio intestinalis) in their intestinal tract as important players in the formation of methyl-Hg. In our study, SRB (D. vulgaris) were also detected in earthworms but not in soils. Therefore, we cannot exclude that SRB may have played a role in the methylation of the Hg in our study. Desulfovibrio vulgaris has also recently been shown to facilitate the methylation of inorganic-Hg in freshwater sediments [7].

The occurrence of methylation in earthworms has potentially implications for the Hg cycle in soil ecosystems as it enriches the methyl-Hg pool in soils, and consequently also in the food web. In addition, biomethylation is not only restricted to anaerobic

Figure 4. Phylogram with the phylogenetic relationships of 16S rRNA gene sequences. The phylogenetic tree shows the bacterial strains and environmental clones most closely to the T-RFs in Figure 3. Whenever possible, closest strains were used for the calculation of the tree but when no strain was available (e.g. many species of *Mollicutes* could not be isolated so far) the closest clone was used. The tree was calculated by Bayesian inference using sequences of 898 bp lengths and shows the affiliation between the clones and closest related sequences of NCBI. The clones of our study are bold marked. Only bootstrap values above 0.9 are given. The scale bar represents 0.1 (10%) of sequence divergence. doi:10.1371/journal.pone.0061215.g004
conditions in soils as usually reported [8,9], but also to well-drained soils.

The impact of inorganic-Hg in soils on soil bacterial community structures has already been reported [4,36–41]. It is also well known, that Hg pollution changes the gut bacterial community in soil invertebrates [42–44]. However, as far as we know, the effects of methyl-Hg on the bacterial community structures in soils and earthworms have never been studied. Here, we showed that the bacterial community structures in the earthworms were clearly affected by Hg. Several T-RFs (87 bp, 278 bp, 279 bp, 319 bp, 450 bp and 482 bp) in the earthworms were affected by Hg and these cloned sequences had closest relatives to sequences retrieved from intestine tracts of earthworms [45–49]. T-REF 319 bp (clone E1) was slightly increased with inorganic and methyl-Hg and cloned sequences were phylogenetically affiliated to *Mollicutes* obtained from earthworms. *Mollicutes* within the *Firmicutes* contain no cell walls and contain a reduced genome size as a consequence of a reductive or degenerative evolution process [50–52]. They obtain nutrients from their host cells (earthworm) parasitically. Therefore, the higher tolerance of *Mollicutes* to Hg could be due to the lower number of genes and enzymes affected by Hg, and/or their supply with nutrients from host cells. When the earthworms were grown in soils with methyl-Hg, T-REF 450 bp became dominant (increase from about 7% to 27% of the total abundance). The cloned sequence of T-REF 450 bp (clone SEst5) was closely related to sequences affiliated to *Acidovorax* sp. (AJ543436) and *Fernminephrobacter* sp. (EFJ214171) within the *Betaproteobacteria* retrieved from earthworms. *Fernminephrobacter* are *Gram*-negative bacteria symbiotically colonizing the nephridia of the earthworms *Eisenia fetida*. Interestingly, they have been described as obligate aerobic organisms occurring under low oxygen conditions [53]. In contrast, *Acidovorax* sp. are able to grow outside of earthworms [53–55]. Nephridia are the paired excretion organs of invertebrates and are comparable to the kidneys of vertebrates. It was suggested a long time ago that nephridia are symbiotically colonized by bacteria [48,56], but the functions of the associated bacteria are not well studied. These symbionts promote the degradation of proteins [56]. *Fernminephrobacter* sp. may also play a role in the detoxification of inorganic-Hg or methyl-Hg in our earthworms. A wide range of bacteria belonging to the phyla of *Firmicutes*, *Actinobacteria*, and *Proteobacteria* are known to be Hg resistant [57,58]. Bacterial resistance to Hg is mainly associated with the presence of mercury resistance (*mer*) operons [59]. The mercuric reductase enzyme (*merA*) catalyzes the conversion of Hg(II) to the volatile Hg(0). The *merB* enzyme degrades organic Hg compounds to the less toxic form Hg(II) [60]. However, *mer* genes are predominant in aerobic environments and were rarely found in obligate anaerobes [61] assuming that this mechanism is hardly important for this study. Other unknown mechanisms for Hg(II) reduction are suggested under anaerobic conditions [62,63].

Another mechanism for microbial Hg resistance is based on the methylation of Hg(II) [64]. Methyl-Hg-chloride may diffuse through the cell membrane, and as a result the concentrations of Hg in the organisms probably decrease. However, the mechanisms which could be important in this study are not known and need to be further investigated.

In our study, the concentrations of methyl-Hg in soils at the start and after incubation did not vary significantly (p<0.05) in experiments with or without inorganic-Hg. Interestingly, in all soils treated with methyl-Hg, the concentrations of methyl-Hg decreased (initial: 0.75 mg methyl-Hg kg−1 soil dw and end: almost 0.3 mg methyl-Hg soil dw) by about 60% during the experiments (data not shown). Bacteria are able to demethylate Hg-species [60,65,66]. Abiotic decomposition of methyl-Hg due to photodegradation has also been reported in surface water [67]. In the abiotic experiments with sterile soil treated with methyl-Hg, the concentrations of methyl-Hg decreased by about 60%, even though the boxes were stored in the dark so that a photodegradation of methyl-Hg in soils could be excluded. Likewise, the concentrations of total-Hg in these soils did not decrease, which indicates that methyl-Hg in the soil was degraded by other factors than photodegradation.

**Conclusion**

We have been able to show that the gut of earthworms provides suitable conditions for the methylation of inorganic-Hg. Control experiments (abiotic; with earthworms rinsing suspension; cast) strongly supported our finding that inorganic-Hg is methylated in earthworms itself, and not by the bacteria introduced into the soils by earthworms. SRB may play a role in the methylation of inorganic-Hg in our earthworms because they were found only in the earthworms and not in the soils. The transformation of inorganic-Hg to the much more toxic form of methyl-Hg may not only occur under anaerobic soil conditions, but also in the anaerobic guts of earthworms inhabiting aerobic soil environments. The occurrence of biomethylation in earthworms may have implications for the Hg cycle in soil ecosystems. This process may enrich the methyl-Hg pool in soils and, consequently, also may enrich methyl-Hg in the food web.

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**Author Contributions**

Conceived and designed the experiments: SR OD IB BF. Performed the experiments: SR BF BL. Analyzed the data: SR BF BL. Contributed reagents/materials/analysis tools: BL BF IB. Wrote the paper: SR BF.

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