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The chemical species of mercury accumulated by Pseudomonas idrijaensis, a bacterium

from a rock of the Idrija mercury mine, Slovenia.

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#### **Abstract**

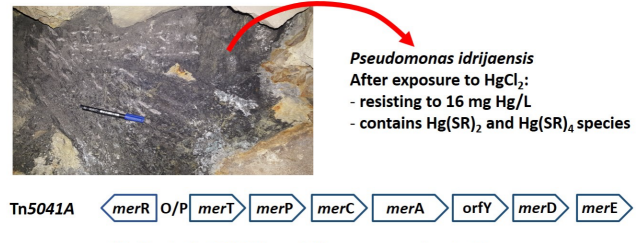
A mercury-resistant bacterial strain has been isolated from a rock of the Idrija mercury mine in Slovenia. The rock had 19 g carbon and 2952 mg mercury (Hg) per kg. Mass spectrometry and DNA sequencing showed that the bacterium belongs to the *Pseudomonas* genus. It is called *Pseudomonas idrijaensis*. This bacterial strain is sensitive to methylmercury (MeHg) like the reference P. aeruginosa strain PAO1, and is resistant to divalent mercury (Hg(II)) in contrast to PAO1. This difference could be attributed to the presence of the mer operon yet deprived of the merB gene encoding the organomercurial lyase, on the basis of whole genome sequencing. The P. idrijaensis mer operon displays the RTPCADE organization and is contained in the Tn5041 transposon. This transposon identified here occurs in other Gram-negative Hg-resistant strains isolated from mercury ores, aquatic systems and soils, including *Pseudomonas* strains from 15,000 to 40,000 years old Siberian permafrost. When P. idrijaensis was exposed to mercury chloride, two intracellular Hg species were identified by high energy-resolution XANES spectroscopy, a dithiolate Hg(SR)<sub>2</sub> and a tetrathiolate Hg(SR)<sub>4</sub> complex. P. idrijaensis had a much higher [Hg(SR)<sub>2</sub>]/[Hg(SR)<sub>4</sub>] molar ratio than bacteria lacking the mer operon when exposed to 4  $\mu$ g Hg<sup>2+</sup>/L - resulting in an intracellular accumulation of 4.3 µg Hg/g dw. A higher amount of the Hg(SR)<sub>2</sub> complex provides a chemical signature for the expression of the dicysteinate Mer proteins in response to mercury toxicity.

# **Highlights:**

- 1. Newly discovered bacterium *Pseudomonas idrijaensis* from a rock of the Idrija mine.
- 2. The rock sample was rich in quartz and mica and contained 2952 mg of mercury per kg.
- 3. Pseudomonas idrijaensis is resistant to 80 µM of mercury chloride (16 mg Hg/L).
- 4. P. idrijaensis is endowed with the mer operon, contained in the transposon Tn5041.
- 5. P. idrijaensis accumulates bis and tetrathiolate species: Hg(SR)<sub>2</sub> and Hg(SR)<sub>4</sub>.

# **Graphical abstract:**

# Rock from the Idrija mine, Slovenia, containing 2952 mg Hg(II)/kg



Similar to Tn5041B from *P. fluorescens*, extracted from a 15-40 thousand year old Siberian permafrost soil

#### 1. Introduction

Although mercury deposits are globally distributed in 26 mercury mineral belts, nearly three quarters of the total world's production has originated from just five mercury belts (Rytuba, 2003). Almadén, the most important mercury belt comprising 11 mercury deposits in central Spain, has produced over one-third of the world's mercury. The second largest mercury mine in the world was Idrija, Slovenia. Idrija is located about 50 km west of the Slovenian capital Ljubljana. The mine is 1500 m long and 300 to 600 m wide and extends below the surface of the Idrija valley in the NW – SE direction. It has produced more than 12.7 Mt of ore with 145,000 t Hg (average content of 1.13 % Hg) since 1490 (Mlakar, 1974). The production of mercury stopped in 1988 and the mine finally closed in 1995. Idrija still contains 10 % of the world's known mercury reserves (Brinck and van Wambeke, 1975). Idrija was considered for centuries to be the center of scientific and technological progress in the region. Mining operations in more than 150 orebodies extended vertically over 360 m (+ 330 to - 33 m) on fifteen levels (Lavrič and Spangenberg, 2003). Ore minerals are cinnabar, metacinnabar, native mercury, pyrite and scarce barite (Palinkaš et al., 2008). The host rocks and ore itself contain organic matter as disseminated kerogen, black solid hydrothermal bitumen, and greenish idrialite, a complex mixture of polycyclic aromatic hydrocarbons containing nitrogen and sulphur-bearing compounds (Lavrič and Spangenberg, 2001). Half a millennium of Hg production has resulted in elevated mercury content in all environmental compartments (Gosar et al., 2016). The Soca River (Isonzo River in Italy) and its tributary, the Idrija River, have drained for centuries the Hg-enriched deposits of the Idrija mine, resulting in trapping of Hg in the sediments of the Gulf of Trieste, that in turn constitute a net source of MeHg (Faganeli et al., 2003). On the banks of the Idrija River, the total Hg and MeHg concentrations in soils are ranging from 0.25 to 1650 mg/kg and not detected to 0.44 mg/kg, respectively (Tomiyasu et al., 2017). Hg levels in foodstuffs from the Idrija mine area were higher compared to those in food from non-contaminated areas (Miklavčič et al., 2013). Retired Idrija mine workers and Idrija residents occupationally unexposed contained much higher Hg concentrations in their tissues (sampled by autopsy) than control individuals (Falnoga et al., 2000). It was shown that the long-term occupational exposure to Hg endured by the Idrija mine workers caused renal dysfunction (Franko et al., 2005). In 2012, Almadén and Idrija were included to the Mercury World Heritage list of the UNESCO. Total Hg concentration in contaminated soils has been reported to vary between 8.4 and 415 mg/kg (Kocman et al., 2004) and between 0.3 to 973 mg/kg (Gosar et al., 2006).

Hg reduces the diversity of bacterial communities in soil, and this decrease is directly proportional to the Hg concentration in soil (Rasmussen et al., 2000). To survive such a hostile environment bacteria are endowed with the *mer* operon. This operon provides bacterial resistance to inorganic and organic Hg through the action of the mercuric reductase encoding *merA* gene and the organomercurial lyase encoding *merB* gene (Barkay et al., 2003; Mathema et al., 2011). Genes of the *mer* operon (*merA* and *merB*) have been previously detected in water samples downstream of the mine, not upstream (Hines, 2000). The Hg-resistant bacterium *Acinetobacter idrijaensis* has been isolated from a soil on the Idrija site. It contains in its genome the *merB*, *merA*, *merP*, *merT*, and *merR* genes (Campos-Guillén et al., 2014). This strain is able to grow in the presence of high concentrations of HgCl<sub>2</sub> (200 mg/L, 737 μM) and catalyzes the volatilization of Hg (*merA* gene).

Here, we document a new bacterial species named *Pseudomonas idrijaensis*. It has been recovered from a rock of the Idrija mine situated approximately 104 m below the surface and containing as much as 2952 mg Hg/kg. We show that this bacterial strain can thrive in such a toxic environment because it contains the *mer* operon in its genome. Another goal of the present study was to address the question related to the chemical forms of the Hg atoms

within the bacterial cells after exposure to inorganic Hg and the possible influence of the *mer* operon on these chemical forms.

#### 2. Materials and methods

#### 2.1. Sampling and characterization of the rock

A piece of black shale/siltstone from the Upper Ladinian formation was sampled aseptically in April 2017 at the third mine level, 110 m below the entrance to Borba Shaft situated 226 m above sea level (Figure S1). The rock was put in a sterile glass bottle and transferred to the laboratory within 4 h. We chose this rock because it contained both native elemental mercury and organic matter. The air temperature in mine was 13 °C and the air humidity 84 % at the time of sampling.

The chemical composition of the bulk sample was determined by the ACME Analytical Laboratory, Canada. Major oxides were determined by ICP-MS after fusion in lithium borate and digestion in dilute nitric acid (Table 1). Aqua Regia digestion and ICP-ES/ICP-MS analysis were used for selected elements (As to Zn in Table 2). Carbon and sulfur were determined using a LECO analyzer. The mineral composition was determined by X-ray powder diffraction (XRD) using a Philips diffractometer (graphite monochromator, CuKα radiation, proportional counter). The mineral phases were identified using the Powder Diffraction File (1996) data system and the Panalytical XPert HighScore (v. 1.0d) program package. Phyllosilicates were identified using published methods (Moore & Reynolds, 1989). Semi-quantitative estimates of minerals were obtained from measurements of the relative peak intensities. Relative abundances are presented with Xs (Table S1). The pH value of the rock sample was determined after dispersion of its powder (1:2.5) in distilled water.

# 2.2. Isolation and characterization of Pseudomonas idrijaensis

Ten grams of rock from the Idrija mine were crushed in a mortar and the powder suspended in peptone water. The suspension was concentrated on sterile membrane filters with 0.45 µm pore size and placed on Nutrient Agar (Biolife). Plates were incubated aerobically at 22°C for 72h, after which the growth of colonies was checked. Only one morphological type of colony was grown. It was recultivated in pure culture. The isolate was characterized initially by routine bacteriological techniques.

Before whole genome sequencing the isolated bacterium was identified by matrix-assisted laser desorption ionization-time of flight mass spectrometry - MALDI-TOF MS, as described in the Supporting Information (SI, paragraph 1).

# 2.3. Determination of mercury chloride minimum inhibitory concentration (MIC) and resistance tests

The MICs values to mercury chloride exposure were determined for the *P. idrijaensis* and *P. aeruginosa* PAO1 strains. The PAO1 strain is a reference strain known to be sensitive to mercury. Bacterial cultures were grown overnight in Luria-Bertani (LB) medium and diluted to an OD<sub>600</sub> of 0.01. Then 2 ml of the diluted overnight culture were dispensed in 10 ml plastic tubes containing 50 μl of a blend made up with water and a stock solution of HgCl<sub>2</sub> in order to reach final concentrations between 0 and 100 μM HgCl<sub>2</sub>. The stock solution of HgCl<sub>2</sub> was prepared at a concentration of 20 mM (4 g/L). Each final concentration was assayed in triplicate for each bacterial species. After 16 h of incubation at 37 °C, cell growth was monitored by spectrometry (absorbance at 600 nm) and MICs values were registered.

Resistance tests to HgCl<sub>2</sub> and monomethylmercury chloride (MeHgCl) were carried out at the surface of agar-containing LB medium. 50 µl bacterial cultures in overnight LB medium were spread over plastic Petri dishes containing 15 ml agar-jellified LB medium. 15, 30, 150, and 300 µg of Hg from 4 and 40 g/L HgCl<sub>2</sub> and from 5 and 50 g/L MeHgCl stock

solutions were spiked at the center of the Petri dishes and the boxes incubated at 37°C for 20 h. The same assay was repeated at 15°C for 48 h. The growth inhibition was read as the diameter of the zone of inhibition.

#### 2.4. Genomic DNA extraction

From a fresh streak of *P. idrijaensis* on an agar-jellified LB medium, 6 ml of liquid LB medium were inoculated and grown overnight at 37°C. The bacterial genomic extraction followed a published recipe (Wilson, 2001). Briefly, the bacteria were lysed with SDS and proteinase K, and the genomic DNA treated with CTAB/NaCl, then extracted with chloroform/isoamylic alcohol (24:1) and afterward with phenol/chloroform/isoamylic alcohol (25:24:1). The genomic DNA was precipitated with 0.6 volume of isopropanol, and after centrifugation washed with 70 % ethanol, then dried. The genomic DNA was dissolved in 10 mM Tris, 1 mM EDTA (TE buffer pH 8.0). After quantification of its concentration by UV spectrometry at 260 nm (NanoDrop 2000 apparatus, Thermo Fisher Scientific), the genomic DNA was diluted in TE to a concentration of 0.1 mg/mL.

# 2.5. Whole genome sequencing

The *P. idrijaensis* genome has been sequenced by the Genome Scan company (Leiden, The Netherlands) using Illumina next generation sequencing technology (Illumina Hiseq 4000). Data were assembled with Spades v3.11.1.

# 2.6. Exposure to $HgCl_2$ for XANES spectroscopy

From a fresh streak of *P. idrijaensis* on an agar-jellified LB medium, 3 ml of liquid LB medium were inoculated and grown overnight at 37°C. Then 50 ml of LB were inoculated with that culture and grown at 37°C for 8 h. With this starting culture, 2 L of LB were

inoculated and grown overnight at 37°C. The resulting turbidity was 0.96 at 600 nm, corresponding to 0.288 g biomass (one unit of OD<sub>600</sub> corresponds to 0.3 g/L bacteria). The 2 L culture was split in two 1 L bottles and centrifuged for 10 min at 5000 rpm in a JLA8.1 rotor (Beckman-Coulter Avanti JXN-26 centrifuge). Each cell centrifugate was washed with 250 ml of sterile minimally complexing medium (MCM buffer). The composition of the MCM buffer was slightly modified from Dahl et al. (2011): 20 mM 3-(N-morpholino)propane sulfonic acid (MOPS buffer), 0.41 mM MgCl<sub>2</sub>, 6 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, and 10 mM glucose. Each cell pellet was resuspended in 500 ml of MCM and spiked with HgCl<sub>2</sub> from a 20 mg/L HgCl<sub>2</sub> stock solution. 100 µl were added to one cell batch and 1 ml to the other batch yielding nominal concentrations of 4 and 40 µg Hg/L, respectively (19.9 and 199 nM). The two cultures were incubated at 37°C for 195 min, then each culture was split in two and centrifuged for 10 min at 5000 rpm in a JLA8.1 rotor. Two cell pellets (one exposed to 4 and the other to 40 µg Hg/L) from split 1 were used to analyze mercury speciation in whole bacterial cells (Tot fraction). These cell pellets were washed with 45 ml of MCM and centrifuged in 50 ml Falcon tubes at 4500 rpm for 10 min (in swinging buckets n° 8172, using the centrifuge Heraeus Labofuge 400R, Thermo Scientific). The two remaining other cell pellets (one exposed to 4 and the other to 40 µg Hg/L) were used to analyze mercury speciation in the intracellular bacterial compartment (Intra fraction). To remove mercury from the cell membrane and cell wall without compromising the cytoplasmic membrane integrity, a two-step washing procedure was used (Thomas & Gaillard, 2017). Cell pellets were resuspended in 200 mL of 50 mM EDTA and 100 mM oxalate solution (pH = 7.5) and after 10 min of reaction were centrifuged for 10 min at 5000 rpm in a JLA8.1 rotor. The centrifugates were resuspended in 50 mL of a 10 mM reduced glutathione and 3 mM ascorbate solution (pH = 7). After 5 min of reaction, the suspensions were centrifuged in 50 ml Falcon tubes at 4500 rpm for 10 min, and the cells washed with 45 ml MCM then centrifuged in 50 ml Falcon tubes at 4500 rpm for 10 min. The four cell preparations were lyophilized after 1 h of storage at -80°C. The four final bacterial biomasses weighed about 250 mg.

# 2.7. Measure of Hg concentration

Mercury in the four prepartions was quantified with a DMA-80 mercury analyzer (Milestone Dual-cell). The detection limit was 0.003 ng Hg and accuracy was confirmed by analysis of the IAEA-436 reference from the International Atomic Energy Agency. The recommended value is  $4.19 \pm 0.36$  mg kg<sup>-1</sup> d.w. and the determined value was  $3.98 \pm 0.13$  mg Hg kg<sup>-1</sup> dry weight (n = 21).

# 2.8. XANES identification of the mercury species in P. idrijaensis exposed to HgCl<sub>2</sub>

Mercury L<sub>3</sub>-edge high energy-resolution XANES (HR-XANES) spectra were measured at 10-15 K with high-reflectivity analyzer crystals (Rovezzi et al., 2017) on beamline ID26 at the European Synchrotron Radiation Facility (ESRF). Data were analyzed against a large database of spectra for mercury minerals (α-HgS, β-HgS, β-HgS<sub>NP</sub>), Hg(II) complexes in natural organic matter, and Hg(II) and methylmercury (MeHg) model complexes with thiolate ligands (Manceau et al., 2015a, 2015b, 2016, 2018, 2019; Bourdineaud et al., 2019). All reference spectra were considered as a basis for identification, but only diagnostic spectra are discussed herein.

The proportions of the Hg species were evaluated using least-squares fitting (LSF) of the experimental spectra with linear combinations of the component spectra. The regression analysis was performed by minimizing the normalized sum-squared residual  $NSS = \Sigma[(y_{exp} - y_{fit})^2]/\Sigma(y_{exp}^2)$ . The precision of the percentage of a fit component was estimated to be equal to the variation of its optimal value when the fit was degraded by 20% (i.e., 20% increase of NSS).

#### 3. Results

# 3.1. Characterization of the rock

The pH value of the rock powder was 3.67. Quartz and mica are the main constituents (Table S1), in agreement with chemical analysis (Tables 1 and 2). They are followed by kaolinite and gypsum. Pyrite, goethite and rozenite are minor (Table S1). Goethite, rozenite and gypsum are weathering products of pyrite. No mercury-bearing phases such as cinnabar and metacinnabar were identified in the sample. Yet, this rock contains a high level of mercury, 2952 mg/kg, and several other toxic metals and metalloids at a concentration above 10 mg/kg: As, Mo, Pb, Sb, U, and V (Table 2).

# 3.2. Characterization of P. idrijaensis

The bacterial isolate recovered from the rock possesses the following characteristics: Gram-negative rod, oxidase and catalase positive, no reaction on the Kligler Iron Agar. The MALDI-TOF MS identification yielded 10 consecutive score values form 1.198 to 1.462 for *Pseudomonas* spp (Table S2). The findings indicate reliable identification of the genus *Pseudomonas*, but unsuccessful species identification. The isolate proved to be sensitive to all tested antibiotics (Table 3).

Over the course of the search for the *mer* operon genes and before processing to the whole genome sequencing, a 2278 bp fragment was amplified using the R1 and A5 primers targeting the *mer*R and *mer*A genes, respectively (SI, paragraph 2). After cloning and sequencing, this fragment appeared to be unrelated to the *mer* operon because the R1 probe had hybridized to both extremities of the amplified fragment. The closest sequence retrieved from a blast inquiry of GenBank was that of the formate dehydrogenase  $\alpha$  subunit of *Pseudomonas alkylphenolica*, strain KL28 (GenBank accession number CP009048) (Mulet et

al., 2015). The two DNA sequences were identical at 94 %. After whole genome sequencing the *P. idrijaensis fdo*G2 gene proved to determine the synthesis of an 808 residues protein best matching with its *P. alkylphenolica* counterpart with a 97.6 % identity. These genetic data confirm the mass spectrometry identification of *Pseudomonas* spp. Due to highly repetitive sequences, part of the *P. idrijaensis* genome could not be sequenced and that part included the 16S rRNA genes. Therefore, the classical identification through 16S rRNA sequence comparisons could not be performed.

3.3. Resistance of P. idrijaensis to mercury chloride and sensitivity to methylmercury chloride

After 16 h of incubation in liquid LB medium at 37 °C, the highest concentrations of

HgCl<sub>2</sub> for which no impact on cell growth was observed were 5 μM for P. aeruginosa PAO1

and 80 μM for P. idrijaensis. The MICs of HgCl<sub>2</sub> were 10 μM for PAO1 and 90 μM for P.

idrijaensis. The growth surfaces were three- to six-times less inhibited than those of PAO1

depending on the amount of Hg(II) (Table 4). Thus, P. idrijaensis is clearly resistant to

Hg(II). However, the two strains featured the same sensitivity to methylmercury chloride

(Table 4). Results were independent of the temperature of growth (37 and 15 °C).

# 3.4. Identification of the mer operon genes in P. idrijaensis

The resistance tests suggested that the *P. idrijaensis* strain harbored at least the *mer*R, *mer*P, *mer*T and *mer*A genes encoding the mercuric reductase in response to Hg(II) exposure, whereas *mer*B encoding the organomercurial lyase was absent. Before processing to the whole genome sequencing we first obtained PCR evidences of the presence of the *mer* operon within the bacterium's genome (SI, paragraph 3 and Figure S2). The amplified fragments revealed that the organization of the *mer* operon was RTPCA, with *mer*R transcribed in opposite direction relative to *mer*TPCA (Fig. S2).

The whole genome sequencing of *P. idrijaensis* confirmed the existence of a *mer* operon with the genes organization deduced from PCR analysis. However, it also contains the merD and merE genes (Fig. 1). The mer operon is contained within the Tn5041 transposon. A blast of scaffold 5 from P. idrijaensis matched the Tn5041 sequence from a Pseudomonas species isolated from the soil of the Khaidarkan mercury mine, Kirghizia, Central Asia (Kholodii et al., 1997). Out of 14907 nucleotides composing the transposon, only 7 did not co-occur in the two *Pseudomonas* species. Their locations in the Tn5041 sequence are shown with asterisks in Fig. 1, along with some remarkable genomic characteristics of this sequence. The extended organization of the mer operon is RTPCAYDE with an open reading frame orfY intercalated between the merA and merD genes. The function of the hypothetical protein encoded by orfY is unknown. The transcriptional regulatory signals involved in the mer genes expression are present and intact. They lie within the intergenic region between merR and merT. There is a potential binding site for MerR repressor, ccgtacatgactacgg, with its inverted repeats (underlined), and there are – 35 and – 10 signals allowing transcription of merTPCAYDE in one direction and those for the merR gene transcription in the opposite direction. Noteworthy is the existence of a second merR gene, called merR2, located 51 nucleotides downstream from the right arm of the Tn5041 transposon materialized by the terminal inverted repeat (TIR) (Fig. 1). The encoded protein is 90 % identical to a MerR family of transcriptional regulators contained in the genome of *Pseudomonas* sp. ERMR1:2, a strain isolated in India, East Rathong glacier, West Sikkim (protein accession number: PAM82489; nucleotidic accession number: NKJI01000010). The MerR2 protein contains only one cysteine residue versus 4 for the genuine MerR regulator. The function of MerR2 as a metal regulator has not been proven yet.

#### 3.5. Mercury speciation after exposure to mercury chloride

The dry weight Hg concentrations of the bacterial masses analyzed by HR-XANES were  $[Pi-4-Tot] = 6.7 \pm 0.2 \mu g Hg/g$ and  $[Pi-4-Intra] = 4.3 \pm 0.2 \mu g Hg/g$ for the total and intracellular preparations exposed to 4  $\mu$ g Hg/g, and [Pi-40-Tot] = 61  $\pm$  1  $\mu$ g Hg/g and [Pi-40-Intra] =  $42 \pm 4 \mu g$  Hg/g for those exposed to 40  $\mu g$  Hg/g. The four HR-XANES spectra feature a near-edge peak at 12279.3 eV (indicator region "A" in Fig. 2a) characteristic of linear dithiolate complex (Hg(SR)<sub>2</sub>) (Manceau et al., 2019). The straighter the RS-Hg-SR angle, the sharper is this peak. The trailing spectral edge of Pi-4-Intra is shifted to lower energy in the 12320-12340 eV interval (indicator region "B"). The shift is indicative of longer Hg-S bond distance and means that higher coordination complex, such as Hg(SR)<sub>3</sub> and Hg(SR)<sub>4</sub>, coexists with the main Hg(SR)<sub>2</sub> complex (Fig. 2b). This Hg coordination effect, described previously with a comparison of the Hg(Cys)<sub>2</sub>, Hg(D-penicillamin)<sub>3</sub>, and Hg(Cys)<sub>4</sub> HR-XANES spectra (Manceau et al., 2019), is explained physically by the Natoli rule (Bianconi et al., 1983). The distinctive difference of Pi-4-Intra is better seen in its comparison with Pi-4-Tot (Fig. 2c). The shift in energy observed in region B is accompanied by a decrease in intensity of the near-edge peak in region A. This is because the RS-Hg-SR angles of the three-coordinate or four-coordinate complexes present in Pi-4-Intra and detected in region B are not linear (Fig. 2b). The coordination structures of Hg in Pi-4-Tot and Pi-4-Intra are examined in more detail below.

In Pi-4-Tot, Hg(II) is coordinated linearly to two thiolate ligands (Fig. 2c). The RS-Hg-SR bond axis from a Hg(SR)<sub>2</sub> complex is perfectly straight only when the Hg coordination is strictly two. The proximity to Hg atoms of oxygen and nitrogen donors from typically carbonyl and amine groups bends the RS-Hg-SR angle (Fig. 3a). On the HR-XANES spectra, this is reflected phenomenologically in a decrease of the near-edge peak intensity or sharpness, similarly to the bonding of a third (Hg(SR)<sub>3</sub>) and fourth (Hg(SR)<sub>4</sub>) thiol, but the effect here is more subtle because oxygen and nitrogen are weaker ligands. Moreover, the

attenuation of the near edge peak occurs without modification of region B because the variation of the RS-Hg-SR angle caused by O/N donors does not change the Hg-S distance, in contrast to the addition of a third (Hg(SR)<sub>3</sub>) and fourth (Hg(SR)<sub>4</sub>) sulfur ligand (Fig. 2b). The effect of secondary Hg....O/N interactions on HR-XANES is illustrated in Fig. 3b-d with the Hg[SR]<sub>2</sub>, the Hg[SR+O]<sub>2</sub>, and the Hg[SR+N]<sub>2</sub> coordinations from the Hg(Cys)<sub>2</sub> pH 3, Hg(GSH)<sub>2</sub> pH 7.5, and Hg(Cys)<sub>2</sub> pH 11.5 references (Bourdineaud et al., 2019). Pi-4-Tot has a near-edge intensity intermediate between those of the Hg[SR]<sub>2</sub> and the Hg[SR+O/N]<sub>2</sub> coordinations, which suggests that Hg is secondarily bonded to one O or N ligand on average. This bonding environment is a signature for biologic matter for it is not detected in natural organic matter (Manceau et al. 2015a).

Regarding Pi-4-Intra, the shift in energy of region B (Fig. 2c) has been observed previously in fish exposed to Hg(II)-doped dissolved organic matter (DOM) (Fig. 2a) (Bourdineaud et al., 2019):  $84 \pm 8\%$  of the Hg in fish is two-coordinate (Hg(SR)<sub>2</sub>) and  $16 \pm 8\%$  is four-coordinate (Hg(SR)<sub>4</sub>). A linear fit of Pi-4-Tot and Hg(Cys)<sub>4</sub> to Pi-4-Intra shows that the tetrathiolate species amounts to  $13 \pm 6$  at%, or  $4.3 \times 0.13 = 0.6 \mu g$  Hg/g, inside the bacteria (Fig. 4b). The bacterium and fish spectra differ, however, in region A, the Pi-4-Intra near-edge being more intense than the near-edge of the fish spectrum. This difference was examined further by measuring the spectra of *Escherichia coli* and *Bacillus subtilis* exposed to  $4 \mu g$  Hg/L. The two spectra were statistically identical, therefore averaged to obtain one spectrum representing bacteria with no *mer* operon. This spectrum is also distinctive from Pi-4-Intra in region A (Fig. 4c). The Pi-4-Intra spectrum was best-fit as a linear combination of 43% of the *E. coli-B. subtilis* spectrum plus 56% of the Hg(Cys)<sub>2</sub> spectrum ( $\pm$  6 at%) (Fig. 4d). Thus, higher amounts of the mercury dithiolate species distinguishe *P. idrijaensis* equipped with the *mer* operon. The two Hg(SR)<sub>3</sub> references from our database (Manceau et al., 2019) were both unsuitable models.

# 4. Discussion

The rock sample contains four times less Hg (2952 mg/kg) than the Idrija deposit on average (Mlakar, 1974), and almost seventeen times less than the Almadén mine on average (5%, Higueras et al., 2011). The Hg content of 2952 mg/kg is, however, two times higher than the highest Hg content of 1650 mg/kg reported in soil near the Idrija mine (Tomiyasu et al., 2017).

Comparison of the amounts of Hg added to the living bacteria and measured in the biomass after 195 min exposure shows that bacteria took up almost all the added Hg and volatilized little Hg. In the 40 µg Hg/L experiment, 0.288 g biomass was exposed to 20 µg Hg(II) in 500 ml giving an initial Hg concentration equal to 69 µg Hg/g bacteria dw. The initial Hg concentration was 6.9 µg Hg/g bacteria dw in the 4 µg Hg/L experiment. At the end of the experiment,  $[Pi-40-Tot] = 61 \pm 1 \mu g Hg/g dw$  and  $[Pi-4-Tot] = 6.7 \pm 0.2 \mu g Hg/g dw$ . Thus, bacteria sorbed and retained 89% and 97% of total Hg in the exposure medium. Furthermore, comparison of total and intracellular Hg concentrations shows that two thirds of the Hg diffused into the bacterial cell. Thus, P. idrijaensis efficiently uptakes Hg. 1012 dry bacteria weigh 0.3 g, corresponding to 3.33 x 10<sup>12</sup> bacteria/g dw. After exposure to 40 μg Hg/L, 42 μg Hg/g inside the cells represent  $(42 \times 10^{-6} \times 6 \times 10^{23}) / (200.6 \times 3.33 \times 10^{12}) = 37.8 \times 10^{3}$  atoms of Hg/bacterial cell. In turn, knowing the volume of a bacterial cell, 1.67 µm<sup>3</sup> during the exponential phase of growth in a rich medium (Kubitschek and Friske, 1986), we can calculate an internal Hg concentration of  $(37.8 \times 10^3) / (6 \times 10^{23} \times 1.67 \times 10^{-15}) = 37.7 \mu M$ resulting in a bioaccumulation factor of 189. After exposure to 4 µg Hg/L, 4.3 µg Hg/g inside the cells represent 3.9 x 10<sup>3</sup> atoms of Hg/bacterial cell, making 3.9 µM Hg, therefore a bioaccumulation factor of 196. Such high bioaccumulation factors likely result from the intervening action of the three transporters MerC, MerE and MerT.

A sharper near-edge peak of the HR-XANES spectrum distinguishes *P. idrijaensis* equipped with the *mer* operon. This feature indicates a higher content of the linear coordination of the Hg atom. The three-coordinate Hg-MerR complex induces expression of genes encoding for a series of membrane-bound proteins (MerT, MerC, MerE, MerF, MerH) and the periplasmic protein MerP, which coordinate Hg(II) linearly (MerT, MerC, MerF, MerP; Nascimento and Chartone-Souza, 2003; Serre et al., 2004; Silver and Hobman, 2006; Steele and Opella, 1997; Wilson et al., 2000; Sahlman et al., 1997), or are assumed to do so (MerE and MerH; Kiyono et al., 2009; Schué et al., 2009). Therefore, the enhanced near-edge structure of *P. idrijaensis*, explained by a richer content in Hg dithiolate complex, is probably a signature for the *mer* operon. Consistent with this hypothesis, the least-squares fitting of Pi-4-Intra with the *E. coli + B. subtilis* reference spectrum suggests that 57 % of the Hg is bound to a dicysteine Mer protein (Fig. 4d). This molecular form of mercury, induced by the *mer* operon, could be viewed as a detoxification mechanism of its own.

Divalent mercury is reduced to gaseous Hg(0) by the action of the homodimeric mercuric reductase MerA (Fox and Walsch, 1982; Schiering et al., 1991; Hamlett et al., 1992). The coordination of Hg(II) in each monomer is two-coordinate, and its transfer to the catalytic active site occurs via a T-shape (SR<sub>2+1</sub>) intermediate complex (Lian et al., 2014). Figure 4e shows that Hg(0) has a distinct HR-XANES spectrum characterized by intense Hg-Hg oscillations in the top-edge region. This species is clearly absent in Pi-4-Intra.

The nature of the intracellular four-coordinate species is uncertain. The sulfur ligand can be thiolate, as in metalloprotein  $Hg(SR)_4$  complex and in  $Hg_x(SR)_y$  metallothionein cluster (Manceau et al., 2019), or inorganic sulfide, as in  $\beta$ -HgS (Thomas and Gaillard, 2017; Thomas et al., 2018; Manceau et al., 2018). The second hypothesis was tested by exposing P. *idrijaensis* to 10 times as much Hg (i.e., 40  $\mu$ g Hg/L) as Pi-4-Intra to see if more  $\beta$ -HgS precipitated. The Pi-40-Intra spectrum has a trailing edge shifted to higher energy relative to

Pi-4-Intra, which means less β-HgS, thus dismissing the β-HgS hypothesis (Fig. 4f). The four coordinate species may be linked to the bacterial metallothionein (MT) SmtA/BmtA. This MT, discovered in cyanobacteria, is also present in the *Pseudomonas* genus (Blindauer et al., 2002; Blindauer 2011). The NMR structures give a Zn<sub>4</sub>Cys<sub>9</sub>His<sub>2</sub> cluster for *Synechococcus* SmtA (Blindauer et al., 2001), and Zn<sub>3</sub>Cys<sub>9</sub>His and Cd<sub>4</sub>Cys<sub>9</sub>His clusters for *P. fluorescens* PflQ2 (Habjanič et al., 2018). High similarities of structure were observed between the Zn<sub>4</sub>SmtA and Cd<sub>4</sub>PflQ2 clusters and the [Cd<sub>4</sub>Cys<sub>11</sub>] cluster of the α-domain in rat MT2 (Blindauer et al., 2001; Habjanič et al., 2018), and between the Zn<sub>3</sub>PflQ2 cluster and the [Cd<sub>3</sub>Cys<sub>9</sub>] cluster of the β-domain in human MT2 (Habjanič et al., 2018).

The *mer* operon of *P. idrijaensis* is contained in the Tn5041 transposon discovered in a *Pseudomonas* strain (KHP41) from the Khaidarkan mercury mine in Kirghizia (Kholodii et al., 1997). Tn5041 most likely acquired the *mer* operon from a transposon belonging to the Tn21 subfamily of the Tn3 transposon family (Kholodii et al., 1997). *A. idrijaensis*, the other strain found in soil of the Idrija mine, does not possess the Tn5041 element. This bacterium is endowed with two *mer* operons simply separated by the gene of a hypothetical protein and possessing only one *mer*R gene. The organization of the two *mer* operons is ED<sub>1</sub>BR and D<sub>2</sub>TPAG with the genes E, D1, B, D2 transcribed in one direction and the genes R, T, P, A, G transcribed in the opposite direction. Two *mer*D genes are present, and the relative MerD1 and MerD2 proteins display 61 % of identity. The MerD protein from *P. idrijaensis* is 53 % and 40 % identical to *A. idrijaensis* MerD1 and MerD2, respectively.

The Tn5041-like elements can be divided in 12 types, based on the genetic sequences isolated from several strains originating from mercury mines (Khaidarkan, Caucasus, Carpathians), Russian and American soils, and Russian rivers (Kholodii et al., 2002). Since the Tn5041 transposons from *P. idrijaensis* and *Pseudomonas* KHP41 differ by only 7 nucleotides over 14907, the new transposon identified here is assigned to the Tn5041A type

like to the one from the reference strain KHP41. P. putida DRA525 isolated from an artisanal gold mining camp in Mozambique also harbors a Tn5041-hosted mer operon (Drace et al., 2018). It differs from the *P. idrijaensis* transposon by only one nucleotide and therefore also belongs to the Tn5041A type (indicated in Fig. 3). Hg-resistant bacteria endowed with Tn5041 were not always isolated from soils or sediments polluted by mercury. This transposon has also been found in *Pseudomonas* bacteria isolated from Edoma suite sediments of the Siberian permafrost of Kolyma Lowland (late Pleistocene icy complex). The lowland is formed of fluvio-lacustrine loamy sediments aged from 15,000 to 40,000 years (Petrova et al., 2002; Mindlin et al., 2005). Some of the samples contained as little as 6 to 35 µg Hg/kg, therefore were uncontaminated. The Tn5041 transposon of one permafrost strain (P. fluorescens ED23-70) has been sequenced. Compared with Tn5041A from KHP41, it contains two substituted nucleotides at positions 2875 and 3773, and has one additional short 47 nucleotides-long region ( $\rho$ ) in the left arm at position 653-699 just after the putative attachment (att) site. This transposon was assigned to the B type Tn5041B (Mindlin et al., 2005). Recently, another Hg-resistant bacterium endowed with Tn5041A, the Pseudomonas sp. strain 1239, has been isolated from a soil sampled in the Loire valley, a location that is not affected by mercurial pollution (Crovadore et al., 2018). The sequence of the strain 1239 Tn5041A is identical to that of P. putida DRA525. In fact, the presence of the mer operon within a bacterial genome is not compulsory linked to a Hg-polluted environment: indeed, when seeking for bacteria sharing identical MerD, MerE and OrfY protein sequences with P. idrijaensis, the Blast search allowed retrieving 49 strains all belonging to the Pseudomonas genus. Among these 26 were clinical human isolates and 23 were environmental isolates. Among the latter, one had been isolated from a dairy product, 17 had been collected from sites unrelated to mercurial contamination, and only 5 had been sampled from Hg-polluted sites: one in the Khaidarkan mine (KHP41), one in a gold camp in Mozambique (DRA525), and 3 in the Oak Ridge Field Research Center (MPBC4-3, MPR-R5A and MPR-R5B) (Table **S**3).

Although the soil and rocks from the gold mining camp in Mozambique are devoid of antibiotics, as is also the case for those of the Idrija mine, *P. putida* DRA525 can resist to both Hg(II) and antibiotics. Resistance to antibiotics is conferred through the MexE-MexF-OprN multidrug efflux system (Drace et al., 2018). Co-resistance to both heavy metals enzymatically and antibiotics through efflux pumps is common (Baker-Austin et al., 2006) and probably explains why so many clinical isolates are also endowed with Hg resistance (Table S3). *P. idrijaensis* contains in its genome the MeA-MexB-OprM multidrug efflux system (one *mex*A gene, two *mex*B genes, and four *opr*M genes), which should confer broadspectrum antibiotic resistance (Li et al., 2015). However, *P. idrijaensis* does not resist to known antibiotics, perhaps because it lacks the regulator gene *mex*R. The three genes *mex*A, *mex*B and *opr*M may not be expressed for they are not organized in an operon.

#### 5. Conclusion

While the Tn5041 element occurs most commonly on plasmids, it is inserted in the chromosome of *P. idrijaensis* like for the *Pseudomonas* strain KHP41, *Pseudomonas* sp. strain 1239 and *P. putida* DRA525 (Kholodii et al., 1997; Crovadore et al., 2018; Drace et al., 2018). This mobile element is old since it has been found in *P. fluorescens* bacteria isolated from Siberian permafrost dated to the Pleistocene, about 15,000 to 40,000 BP (Petrova et al., 2002; Mindlin et al., 2005). The permafrost soil contained trace amounts of Hg like the Loire valley soil (from which has been isolated the strain 1239), which indicates that mercury contamination pressure is not compulsory to select this mercury resistance transposon once inserted in the chromosome. The presence of the *mer* operon proved to influence the mass distribution of the mercurial chemical species accumulated within bacterial cells after a challenge with inorganic Hg, with a relative enrichment in the dithiolate Hg(SR)<sub>2</sub> complex.

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<b>Table 1.</b> Chemical composition (in wt. %) of the rock										
SiO <sub>2</sub>	Al <sub>2</sub> O <sub>3</sub>	Fe <sub>2</sub> O <sub>3</sub>	MgO	CaO	Na <sub>2</sub> O	K <sub>2</sub> O	TiO <sub>2</sub>	$P_2O_5$	LOI*	Sum
		4.13			0.50	2.78	0.62	0.15	12.9	99.75
* loss on ignition (1000°C).										

<b>Table 2.</b> Content of sulfur, carbon and selected trace elements in the rock. <sup>a</sup>															
S	C	As	Ba	Bi	Cd	Co	Cu	Hg	Mo	Ni	Pb	Sb	U	V	Zn
2.61	1.91	24.6	258	3.1	< 0.1	9.9	23.3	2 952.0	11.3	9.8	21.1	15.7	11.2	118	41
<sup>a</sup> S and C in wt. %, As to Zn in mg/kg.															

**Table 3.** MIC and zone diameter values of antibiotics against environmental isolate of *P. idrijaensis*.

Antimicrobial		MIC	Zone diameter
category	Antibiotic	(mg/L)	(mm)*
carbapenems	MEM - meropenem	1	27
	IMI - imipenem	1	32
fluoroquinolones	CIP - ciprofloxacin	$\leq 0.25$	30
aminoglycosides	GEN - gentamicin	≤ 1	28
	AMK - amikacin	$\leq 2$	30
	NET - netilmicin		27
cephalosporins	CAZ - ceftazidime	$\leq 1$	18
	FEP - cefepime	$\leq 1$	30
penicillins	TZP - piperacillin/tazobactam	8	26
polymyxins	CST - colistin	$\leq 0.5$	0.75

<sup>\*</sup> determined by E-test. Isolate was sensitive to all tested antibiotics according to EUCAST criteria.

**Table 4.** Compared sensitivity of *P. idrijaensis* to inorganic and organic mercury.

Quantity of mercury (μg) <sup>a</sup>	Growth inhibition zone	Ratio <sup>b</sup>	
	P. aeruginosa PAO1	P. idrijaensis	
$HgCl_2$			
15	$20 \pm 1$	8 ± 1 *	6.2
30	23 ± 1	11.5 ± 1.5 *	4.0
150	$33 \pm 2$	17 ± 1 *	3.8
300	40 ± 2	22 ± 1 *	3.3
MeHgCl			
15	$23 \pm 2$	$26 \pm 1$	0.8
30	$28 \pm 1$	$32 \pm 4$	0.8
150	$46 \pm 2$	$48 \pm 2$	0.9
300	$59 \pm 1$	$55 \pm 3$	1.1

<sup>&</sup>lt;sup>a</sup> A solution containing the indicated quantity of mercury was laid at the centre of the Petri dish.

<sup>&</sup>lt;sup>b</sup> Ratio of surfaces of growth inhibition between PAO1 and *P. idrijaensis*.

<sup>\*</sup> Asterisks indicate significant differences between PAO1 and *P. idrijaensis* growth inhibitions (P < 0.05, n = 3).

# Legends to figures

**Figure 1.** Organization of the *mer* operon of *P. idrijaensis* inside the Tn5041A transposon. The numbers above the arrows symbolizing the Tn5041 elements indicate the positions of the genes and open reading frames (orf, putative genes coding for hypothetical proteins), and some remarkable genetic elements in the 14907 bp-long transposon. The numbers below the arrows indicate the length in bp of the genetic elements. Arrows point in direction of the transcription. XerC2 is an integrase; KfrA is a DNA binding protein involved in the transposition process; OprD2 is an outer membrane porine; TnpC is a repressor of the tnpA gene; TnpA is a transposase of the Tn3-family transposon. The two uncoding TIR sequences are terminal inverted repeats that flank the transposon on its extremities. att5041 is the attachment site. INTA5041 is a 195 bp-long truncated group II intron. R' is a truncated copy of merR.  $\sigma\lambda$  and  $\kappa\gamma$  are relics of mobile elements:  $\sigma\lambda$  is a defective element showing some homology with the insertion sequences IS2 and ISRm1, and κγ is a defective minitransposon of the Tn3-family with terminal inverted repeats. The  $\sigma\lambda$ , R' and  $\kappa\gamma$  elements overlap with orfs 15723 and 1573. The regulatory cis elements O/P are the promoter and operator of the mer operon. The locations of the 7 nucleotides distinguishing Tn5041A of P. idrijaensis and Pseudomonas sp. from the Khaidarkan mine are indicated with asterisks. The substitutions are 811: A/G (A in P. idrijaensis and G in the Khaidarkan bacterium); 2557: T/A; 2558: T/A; 4252: C/T; 8346: C/T; 13835: A/T; 13836: A/T. The symbol ^ indicates the location of the single nucleotide distinguishing Tn5041A of P. idrijaensis and both P. putida DRA525 and Pseudomonas sp. strain 1239. It is located at position 14729: G/C (P. idrijaensis/DRA525).

**Figure 2.** HR-XANES spectra of *P. idrijaensis* cultures exposed to 4 μg (Pi-4) and 40 μg (Pi-40) Hg/L (a,c), and ball-and-stick representation of the Hg(SR)<sub>2</sub>, Hg(SR)<sub>3</sub> and Hg(SR)<sub>4</sub> complexes (b). The Hg-S distances are from Manceau and Nagy (2008).

**Figure 3.** Ball-and-stick representation (a) and HR-XANES spectra (b-d) of three dithiolate complexes differing in the second coordination shell compared to the spectrum of P. *idrijaensis* exposed to 4  $\mu$ g Hg/L (Pi-4-Tot fraction). The ball-and-stick models are from Bourdineaud et al., 2019. Only the bonding environment of Hg is represented for Hg(glutathione)<sub>2</sub>.

**Figure 4.** Comparison of the HR-XANES spectrum from the intracellular fraction of P. idrijaensis exposed to 4  $\mu$ g Hg/L (Pi-4-Intra) with the spectrum from fish exposed to Hgdoped DOM (a), with the average spectrum of the intracellular fractions of E. coli and B. subtilis exposed to 4  $\mu$ g Hg/L (c), and with elemental Hg (e). Linear least-squares fit to the Pi-4-Intra spectrum with  $87 \pm 6\%$  Pi-4-Tot and  $13 \pm 6\%$  Hg(Cys)<sub>4</sub> at pH 11.9 (b), and with  $43 \pm 6\%$  E. coli + B. subtilis and  $57 \pm 6\%$  Hg(Cys)<sub>2</sub> at pH 7.5 (d). Comparison of the Pi-4-Intra spectrum with the intracellular spectrum from P. idrijaensis exposed to 40  $\mu$ g Hg/L (Pi-40-Intra) (f).

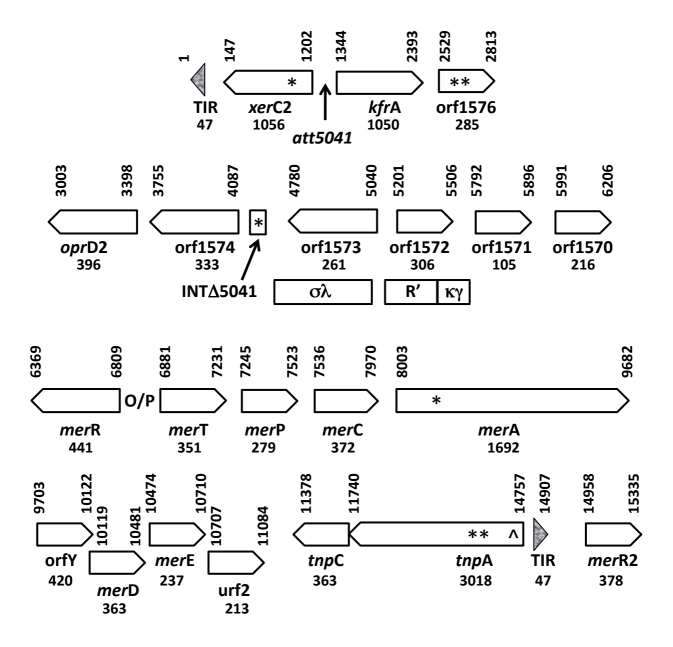


Figure 1

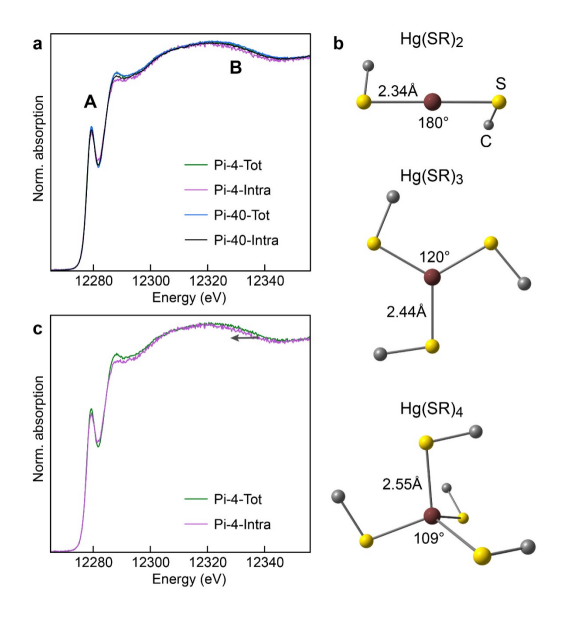


Figure 2

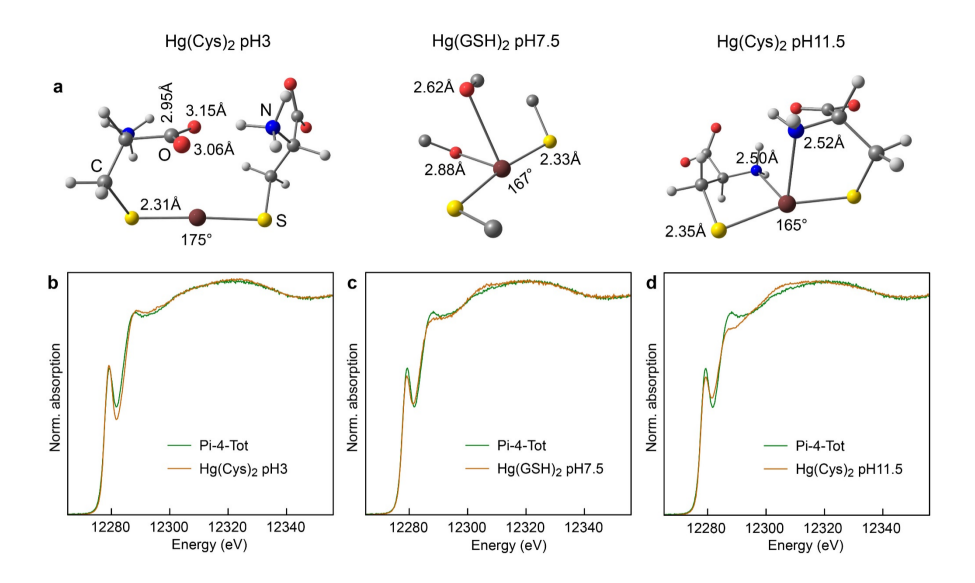


Figure 3

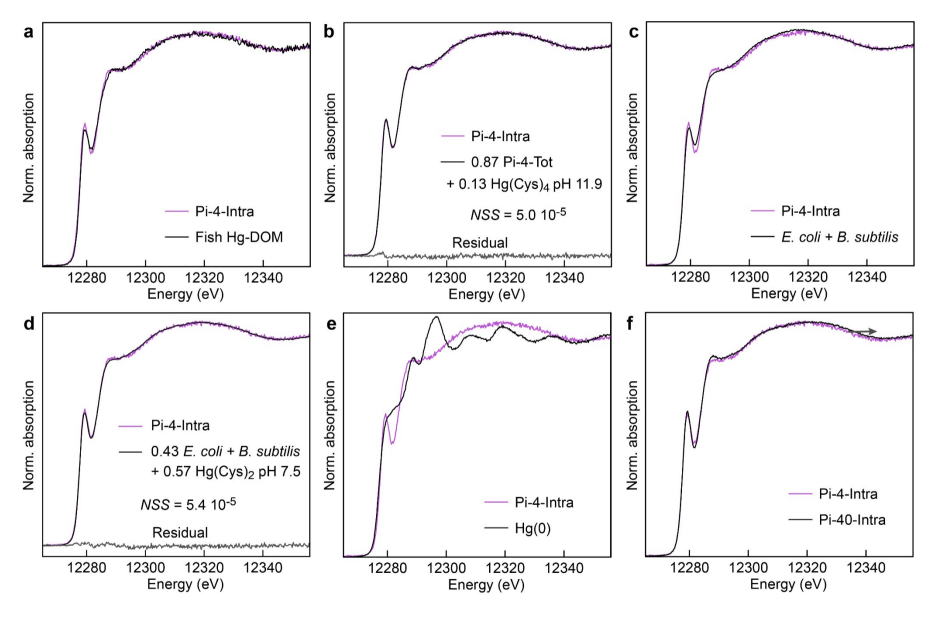


Figure 4

**Supporting Information** 

The chemical species of mercury accumulated by *Pseudomonas idrijaensis*, a bacterium

from a rock of the Idrija mercury mine, Slovenia.

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1

## 1. Characterization of *Pseudomonas idrijaensis* by mass spectrometry

Before whole genome sequencing the isolated bacterium was identified by matrix-assisted laser desorption ionization-time of flight mass spectrometry - MALDI-TOF MS (software version 3.0, Microflex LT, Bruker Daltonics) after extraction of biomass with formic acid (Singhal et al. 2015). A characteristic peptide mass fingerprint (mass-to-charge ratio (m/z) of 2-20 kDa) of the isolate was compared with the spectra of known microbial isolates contained in the MALDI Biotyper 3.0 software. The antibiotic susceptibility profile of the isolate was determined according to minimum inhibitory concentration (MIC) values obtained by the Vitek2 system (bioMérieux) and gradient dilution E-test (bioMérieux). Values were interpreted according to EUCAST criteria (2018) for clinical isolates of *Pseudomonas* sp.

#### 2. Cloning of a 2278 bp fragment from the chromosome

A PCR using the R1 and A5 probes has been performed with 0.1 µg of P. idrijaensis genomic DNA as described below in the paragraph 3. The 2278 bp amplified mer fragment was excised from a band of a 1 % low-melting temperature agarose gel, and purified on a column containing a silica membrane (kit NucleoSpin Gel and PCR Clean-up, Macherey-Nagel, reference 740609.250). The concentration of the purified fragment was assessed by UV spectrometry at 260 nm. The fragment was ligated to the plasmid vector pCR4-TOPO (Invitrogen Thermo Fisher Scientific, reference 450030) as follows: 1 µl of plasmid (10 ng), 2 μl of fragment (36 ng), 1 μl of salt solution (1.2 M NaCl, 60 mM MgCl<sub>2</sub>), and 2 μl of sterile milliQ water were mixed then let rested for 30 min at ambient temperature. Thereafter, Escherichia coli TOP10 cells (Invitrogen Thermo Fisher Scientific, reference C404010; genotype: mcrA,  $\Delta(mrr-hsdRMS-mcrBC)$ , Phi80lacZ(del)M15,  $\Delta lacX74$ , deoR, recA1, araD139, Δ(ara-leu)7697, galU, galK, rpsL(SmR), endA1, nupG) rendered chemically competent for transformation through CaCl<sub>2</sub> treatment were transformed with 2 µl of the ligation mixture. Transformants were selected on ampicillin-containing jellified LB medium (50 µg/ml). Positive recombinants were selected using PCR: 10 colonies were picked out from the Petri dish and each was suspended in a mixture made of 10 µl of sterile milliQ water, 0.5 µl (50 ng) of M13 forward primer, 0.5 µl (50 ng) of M13 reverse primer, and 12.5 µl of 2 x concentrated DreamTaq blend. The thermocycling program used was as follows: initial denaturation step at 94°C for 10 min, 36 cycles at 94°C for 60 s, 55°C for 60 s, and 72 °C for 140 s, followed by a final extension step at 72°C for 10 min. Two transformants were positive

since they gave amplified fragments of about 2700 bp on 1 % agarose gel electrophoresis. These recombinants were cultured on LB medium containing ampicillin ( $50 \mu g/ml$ ) and the harbored plasmids were purified after alkaline lysis on columns containing a silica membrane (kit NucleoSpin plasmid, Macherey-Nagel, reference 740588.250). The fragments contained in these two plasmids were sequenced using M13 forward and reverse primers (GATC Biotech, Eurofins Genomics Company).

#### 3. PCR detection of the *mer* operon genes

Before processing to the whole genome sequencing, the *mer* genes were detected with the following mer oligonucleotide probes: R1, R2, T1, T2, P1, P2, C1, C2, F1, F2, A0, A1, A2, A5, A6, B1, B2, D1, D2, and D3 (Liebert et al., 1997). Letters correspond to the initial of the cognate mer gene (A: merA, B: merB; C: merC; D: merD; F: merF; P: merP; R: merR; T: merT). R1, T1, C1, C2, A1, A5 and A6 are sequences found in the plasmid NR1 from Shigella flexneri with the mer operon organization merRTPCAD (GenBank accession number: K03089). P2, F1, F2 and A2 are sequences found in the plasmid pMER327/419 from Pseudomonas fluorescens with the mer operon organization merRTPFAD (GenBank accession number: X73112). R2, T2, P1, A0, D2, and D3 are sequences found in the plasmid pVS1 from Pseudomonas aeruginosa with the mer operon organization merRTPAD (GenBank accession number: X03406). B1, B2, and D1 are sequences found in the plasmid pDU1358 from Serratia marcescens with the mer operon organization merRTPABD (GenBank accession number: M15049). Two new probes of the merA gene (A1s-nF and A5nR) were designed recently (Ní Chadhain et al., 2006). They are made of degenerated oligonucleotides in order to render them universal. They proved efficient in detecting the merA gene in Pseudomonas species (Ní Chadhain et al., 2006).

The PCRs were performed in a final volume of 20 μl. The mixture contained 1 μl (0.1 μg) of *P. idrijaensis* genomic DNA, 7 μl of sterile milliQ water, 1 μl of each of the two 10 μM oligonucleotide probes (0.5 μM final concentration), and 10 μl of 2 x concentrated DreamTaq blend. This formulation contains the DreamTaq DNA polymerase in an optimized buffer, and dATP, dCTP, dGTP and dTTP, 0.4 mM each, and 4 mM MgCl<sub>2</sub> (Thermo Scientific, reference MAN0012702). The thermocycling program used is called MER1: initial denaturation step at 94°C for 1 min, 36 cycles at 95°C for 30 s, 56°C for 30 s, and 72 °C for 3 min, followed by a final extension step at 72°C for 10 min. Amplified DNA fragments were observed using agarose gel electrophoresis at a concentration in agarose of 1.0 or 1.2 %).

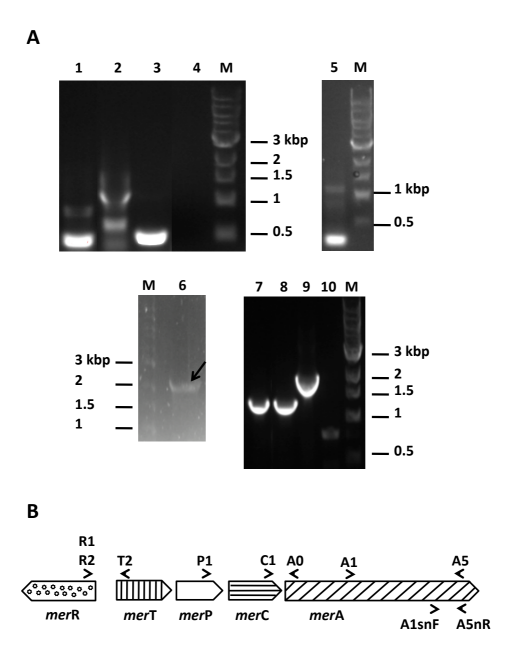
Genetic proof to support the presence of the *mer* operon within the *P. idrijaensis* genome

was obtained with the amplification of the expected 285 bp amplicon using the A1s-nF and A5-nR primers targeting the merA gene (Fig. S2, lane 5). The presence of the merA gene was confirmed with the A1 and A5 primers resulting in the amplification of a fragment of about 1200 bp (Fig. 2, lane 2). In contrast, primers B1 and B2 targeting merB failed to amplify a cognate fragment, in agreement with the lack of resistance to MeHgCl (Fig. S2, lane 4). The organization of the *mer* operon was decrypted with a set of primer couples (Liebert et al., 1997). The R2-T2 primer couple amplified a fragment of about 250 bp (Fig. S2, lane 1), meaning that the merT gene is adjacent to the merR gene. The P1-A0 primer couple amplified a fragment of about 500 bp (Fig. S2, lane 3), meaning that between the genes mer P and merA was lying the gene merF or merC (in the case of the absence of one of these genes, a fragment of about 100 bp would have been expected). The C1-A5 primer couple amplified a 1750 bplong fragment, whereas no fragment was amplified with the F1-A5 primer (Fig. S2, lanes 9 and 10). This means that the gene in between merP and merA is merC. The presence of the supplementary gene merC between merP and merA was further confirmed with the P1-A5, R1-A0, and R2-A0 couples. 2100, 1300, and 1300 bp-long fragments were amplified, respectively, instead of 1650, 850, and 850 bp-long fragments if merC were absent (Fig. S2, lanes 6 to 8). Thus, the organization of the *mer* operon is RTPCA, with *mer*R transcribed in opposite direction relative to *mer*TPCA (Fig. S2, operon scheme).

## 4. Supplementary Figures



**Figure S1.** Photograph of the rock sample. It is an Upper Ladinian black shale/siltstone from the third level of the Idrija mine situated 104 m below the Borba Shaft entrance situated 226 m above sea level. A pencil indicates the scale.



**Figure S2.** Detection of the *mer* genes in the *P. idrijaensis* genome. A: The agarose gels show the PCR amplification of the *mer* fragments from the purified genomic DNA. Lane M: molecular weight ladder. Lane 1: R2 and T2 primers; lane 2: A1 and A5 primers; lane 3: P1 and A0 primers; lane 4: B1 and B2 primers; lane 5: A1snF and A5nR primers; lane 6: P1 and A5 primers (the arrow points out the amplified band); lane 7: R1 and A0 primers; lane 8: R2 and A0 primers; lane 9: C1 and A5 primers; la ne 10: F1 and A5 primers. B: Organization of the *mer* operon deduced from the PCR analysis. The locations of the probes' hybridization sites are shown with truncated arrows. The direction of the arrows indicates the orientation the primer: right for forward and left for reverse.

# **5. Supplementary Tables**

Table S1. Semi-quantitative mineralogical composition of the rock Idrija.

Quartz	Micaceous minerals	Kaolinite	Gypsum	Pyrite	Goethite	Rozenite
XXXX	XXX	XX	XX	X	X	X

X: relative abundance of minerals based on X-ray diffraction (no quantitative value is assigned to X).

**Table S2.** The MALDI-TOF MS identification results for *P. idrijaensis*.

Rank	Matched pattern	Score value	NCBI identifier
1	Pseudomonas graminis DSM 11363T HAM	1.46	<u>158627</u>
2	Pseudomonas chlororaphis ssp chlororaphis DSM 50083T HAM	1.40	<u>333</u>
3	Pseudomonas caricapapayae LMG 2152T HAM	1.37	<u>46678</u>
4	Pseudomonas cichorii DSM 50259T HAM	1.27	<u>36746</u>
5	Pseudomonas chlororaphis ssp aurantiaca CIP 106718T HAM	1.25	<u>333</u>
6	Pseudomonas syringae ssp syringae DSM 6693 HAM	1.22	<u>317</u>
7	Pseudomonas lutea LMG 21974T HAM	1.21	243924
8	Pseudomonas savastanoi ssp savastanoi LMG 2209T HAM	1.20	<u>29438</u>
9	Pseudomonas fluorescens DSM 50090T HAM	1.19	<u>294</u>
10	Pseudomonas abietaniphila CIP 106708T HAM	1.37	<u>89065</u>

**Table S3.** *Pseudomonas* strains sharing identical MerD, MerE and OrfY protein sequences with *P. idrijaensis*.

Strain	Source or sampling location	Reference	
Pseudomonas sp. KHP41	Soil of Khaidarkan mercury mine, Kirghizia, Central Asia.	Kholodii <i>et al.</i> , 1997.	
Pseudomonas sp. ABFPK.	Clinical isolate, USA.	Segre and Mullikin, 2017 (unpublished).	
Pseudomonas sp. NBRC 111118.	Human clinical strain resistant to various antibiotics (Japan).	Shimodaira <i>et al.</i> , 2017, (unpublished).	
Pseudomonas sp. NBRC 111119.	Isolated from human urine, resistant to various antibiotics (Takarazuka, Hyogo, Japan).	Shimodaira <i>et al.</i> , 2017, (unpublished).	
Pseudomonas sp. NBRC 111127.	Isolated from human urine, resistant to various antibiotics (Wakayama, Japan).	Shimodaira <i>et al.</i> , 2017, (unpublished).	
Pseudomonas sp. NBRC 111142.	Isolated from <mark>human urine</mark> , Tochigi, Japan.	Shimodaira <i>et al.</i> , 2017, (unpublished).	
Pseudomonas sp. ANT_H4 and ANT_H14	Isolated from a petroleum-contaminated soil collected near the petroleum pumping and storage warehouse at the Henryk Arctowski Polish Antarctic Station, King George Island, Admiralty Bay.	Styczynski <i>et al.</i> , 2018 (unpublished).	
Pseudomonas sp. 1239.	Isolated from soil samples from the lower Loire Valley, France.	Crovadore <i>et al.</i> , 2019.	
Pseudomonas sp. FSL W5-0299.	Isolate causing color defects in dairy products. Sampled on an agitator drain in a cheese-processing plant,	Reichler <i>et al.</i> , 2019.	

# USA.

Pseudomonas MB-Isolation source: Lake Michigan sp. Batrich et al., 090624 freshwaters, Montrose Beach, 2019 Chicago, USA. Spencer et al., *Pseudomonas* sp. Groundwater from a contaminated MPBC4-3, MPR-R5A and 2019 watershed. Oak Ridge Field MPR-R5B (unpublished); Research Center, Tennessee, USA. Marshall, 1983 P. abietaniphila KF701 Furukawa *et al.*, and KF717 1989; Fujihara et Biphenyl-contaminated soil in al., 2015 Kitakyushu, Japan. P. abietaniphila ATCC Varghese et al., Isolated from a lagoon, British 2019 700689. Columbia, Canada. (unpublished). **ATCC** Isolated from an industrial water aeruginosa Xu et al., 1998; 700888 (ERC1). system, USA. Chugani et al., 2012. P. aeruginosa AZPAE 14381. Clinical isolate from an intraabdominal infection, Bilbao, Spain. Kos *et al.*, 2015. Resistant to meropenem, levofloxacin, and amikacin. Does possess the β-lactamase genes  $bla_{OXA-46}$  and  $bla_{OXA101}$ . P. aeruginosa AZPAE Clinical isolate from an intraabdominal infection, Shatin, China. 14404. Sensitive to meropenem, Kos et al., 2015. levofloxacin, and amikacin. Does not possess β-lactamase genes. Clinical isolate from a reproductive P. aeruginosa **AZPAE** Kos et al., tract infection, Besançon, France. 14816. 2015. Sensitive to meropenem, levofloxacin, amikacin and colistin. Does not possess  $\beta$ -lactamase genes.

P. aeruginosa AZPAE 14858.	Clinical isolate from an intra- abdominal infection, Paris, France. Sensitive to meropenem, levofloxacin, amikacin and colistin. Does not possess β-lactamase genes.	Kos et al., 2015.
P. aeruginosa AZPAE 14872.	Clinical isolate from a reproductive tract infection, Victoria, Argentina. Resistant to meropenem and levofloxacin. Sensitive to amikacin and colistin. Does possess the $\beta$ -lactamase genes $bla_{OXA-2}$ .	Kos et al., 2015.
P. aeruginosa AZPAE 14921.	Clinical isolate from a reproductive tract infection, Paris, France. Sensitive to meropenem, levofloxacin, amikacin and colistin. Does not possess β-lactamase genes.	Kos et al., 2015.
P. aeruginosa AZPAE 14941.	Clinical isolate from an intra- abdominal infection, Hong-Kong, China. Sensitive to meropenem, levofloxacin, amikacin and colistin. Does not possess β-lactamase genes.	Kos et al., 2015.
P. aeruginosa AZPAE 15002.	Clinical isolate from an intra- abdominal infection, Bilbao, Spain. Resistant to meropenem, levofloxacin, and amikacin. Does possess the $\beta$ -lactamase genes $bla_{OXA-46}$ and $bla_{OXA101}$ .	Kos et al., 2015.
P. aeruginosa WH-SGI- V-07167.	Isolated from a human wound, France.	van Belkum <i>et al.</i> , 2015.
P. aeruginosa WH-SGI- V-07371.	Hospital isolate, France.	van Belkum <i>et al.</i> , 2015.
P. aeruginosa WH-SGI- V-07415.	Hospital isolate, France.	van Belkum <i>et al.</i> , 2015.
P. aeruginosa WH-SGI-V-07709.	Hospital isolate, USA.	van Belkum <i>et al.</i> , 2015.

P. aeruginosa 15.111b	P. aeruginosa 15.111b (Cardiff University, UK) is a preservative resistant isolate derivating from the strain PA14 which was originally isolated from a human burn wound.	Weiser <i>et al.</i> , 2017 (unpublished). Cullen <i>et al.</i> , 2015 for the description of the reference strain PA14.
P. aeruginosa ENV-681.	Isolated from manure, Tartu, Estonia.	Laht <i>et al.</i> , 2018 (unpublished).
P. aeruginosa FFUP-PS-65.	Clinical isolate from human urine, Portugal.	Botelho <i>et al.</i> , 2018.
P. aeruginosa AR_0440 and AR_0355.	Antibiotic resistant isolates, US Food and Drug Administration, USA.	Benhamed <i>et al.</i> , 2018 and 2019 (unpublished).
P. aeruginosa AUS511 and AUS527.	Environmental isolates from an Australian River, and a home, respectively, Brisbane, Australia.	Ref: Jeukens et al., 2019.
P. aeruginosa PA3.	Clinical isolate from a bronchial aspirate, bacteriological ward of hospital Bichat, Paris, France.	Mammeri H., 2019 (unpublished).
P. coronafaciens LMG5060.	Isolated from an oat leaf, Avena sativa, UK.	An et al., 2015.
P. fluorescens AU11518.	Isolated from a cystic fibrosis patient, USA.	Scales <i>et al.</i> , 2015.
<ul><li>P. mandelii NBRC</li><li>103147 (CIP105273,</li><li>DSM17967).</li></ul>	Isolated from mineral water, France.	Verhille <i>et al.</i> , 1999; Hosoyama <i>et al.</i> , 2019.
P. monteilii CD10-2.	Duodenal mucosa of celiac disease patient with type 1 diabete, India.	Kaur <i>et al.</i> , 2017 (unpublished).
P. putida DLL-E4.	Paranitrophenol degrader. Isolated from methyl-parathion-polluted soil, China.	Liu <i>et al.</i> , 2003; Hu <i>et al.</i> , 2014.
P. putida ATH-43.	Isolated from soil sediments at a military base, Antarctica, Chile.	Rodriguez-Rojas <i>et al.</i> , 2016.

	Mercury and tellurite resistant bacterium. Resistant to Cd <sup>2+</sup> , Cu <sup>2+</sup> , CrO <sub>4</sub> <sup>2-</sup> , and SeO <sub>3</sub> <sup>2-</sup> , and several antibiotics including streptomycin, cefotaxime, kanamycin, and chloramphenicol.	
P. putida DRA525.	Isolated from an artisanal gold mining camp in Mozambique.	Drace et al., 2018.
P. putida 12917.	Clinical isolate from human stools, Paris, France. Resistant to carbapenem.	Liapis <i>et al.</i> , 2019.
P. putida 142223.	Clinical isolate from human urine, France. Resistant to carbapenem.	Liapis <i>et al.</i> , 2019.
P. syringae pv. coronafaciens Pcn3113.	Isolated from an oat leaf, Avena sativa, England.	Dillon et al., 2019.
P. veronii 1YdBTEX2.	Isolated from soil highly contaminated with benzene, Czech Republic.	Junca and Pieper, 2004; de Lima- Morales <i>et al.</i> , 2013.

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