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Mercury, organic matter, iron, and sulfur co-cycling in a ferruginous meromictic lake

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Abstract

Mercury (Hg) speciation in natural waters is controlled by redox conditions and microbiological activity. Water columns of meromictic lakes have large and stable redox chemical and biological gradients that allow the investigation of many Hg chemical transformations. In this study, Hg speciation (elemental $Hg = Hg^0$, methylated Hg = MeHg) and partitioning between truly dissolved (< 3 kDa), colloidal (< 0.45 μ m and > 3 kDa), and particulate (> 0.45 μ m) fractions, were determined throughout a high-resolution water column profile in the ferruginous meromictic Lake Pavin (Massif Central, France) in July 2018. Total Hg concentrations (THg) in water ranged between 0.4 to 8.8 pmol L⁻¹. The particulate phase represented 10 to 70% of the THg, with a peak found in the mesolimnion associated with the particulate organic carbon maximum. In the mesolimnion, the colloidal fraction represented 12 to 68% of THg, and the highest value was found at the top of the sulfidic zone, whereas the truly dissolved Hg species $(70 \pm 9\%)$ dominated in all the rest of the sulfidic zone. MeHg ranged from less than 10% of THg in the oxic mixolimnion to more than 90% in the anoxic monimolimnion. The Hg methylation was most active within the suboxic zone where iron and sulfate reduction are occurring. These results, associated with those of the partition of organic matter (OM), sulfur, and iron, in conjunction with thermodynamic calculations, allow us to present a conceptual scheme for the Hg cycle in the lake. Atmospheric Hg deposited onto surface waters of the lake is partially photo-reduced and returned to the air, another part is scavenged by biogenic particulate matter and conveyed at depth by settling organic material. Water stratification and redox changes create a sequence of reactions from oxic to ferruginous waters where Hg is successively (i) desorbed from particulate OM where mineralization occurs, (ii) adsorbed onto iron-oxy(hydr)oxides, (iii) desorbed where they dissolved, (iv) precipitate as HgS, (v) methylated, and (vi) reduced as Hg⁰ in the deepest part of the lake. In brief, the (micro)biological uptake, OM, iron and sulfur recycling, through associated microbial consortia, control the Hg cycling in the Pavin waters.

Keywords: Mercury; methylmercury; redox; iron; sulfur; organic matter; Lake Pavin

1 1. INTRODUCTION

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Mercury (Hg) is present in natural waters mainly in two oxidation states (0 and II). The zerovalent form is the dissolved gaseous elemental Hg (Hg⁰), and the divalent forms (Hg^{II}) are often tightly bound to sulfur (S) and carbon (C) (Wang et al., 2022; Xia et al., 1999). Hg^{II} forms preferentially strong complexes with reduced inorganic sulfur and organic matter (OM) via sulfhydryl (-SH) groups (Dyrssen and Wedborg, 1991; Riccardi et al., 2013). It also forms covalent bonds with C (Barone et al., 1996), which leads to the natural occurrence of the highly toxic methylated Hg (MeHg) compounds, including its two forms; i.e., monomethylmercury (MMHg), and the volatile dimethylmercury (DMHg). On the other hand, Hg-S affinity explains the dominant association of Hg with natural OM enriched in reduced S functional groups (Loux, 1998). The biogeochemical cycle of Hg in natural waters involves numerous thermal, photo, and biogenic reactions. Among them, two reversible and competitive Hg transformations are crucial from an aquatic ecology point of view: oxidation-reduction and methylationdemethylation. Indeed, reduction of Hg^{II} leads to the formation of volatile Hg⁰ that can escape from the aquatic environment, and methylation leads to the formation of MMHg that, through its biomagnification in aquatic food webs, is harmful to fish consumers. For this latter reaction, microbial activity is crucial and largely dominates abiotic production (Cooper et al., 2020; Parks et al., 2013; Regnell and Watras, 2018; Ullrich et al., 2001). These reactions occur particularly in hypoxic and anoxic environments (Fitzgerald et al., 2014; Mason et al., 2012), making the Hg cycling in natural waters strongly dependent on redox conditions (Mason et al., 1999). In addition to reactions occurring in the dissolved phase, Hg may precipitate as sulfur minerals (e.g., cinnabar) in euxinic environment, and is involved in heterogeneous reactions, such as sorption onto various minerals, including iron minerals like goethite (Kim et al., 2004; Slowey and Brown, 2007) and mackinawite (Jeong et al., 2007). Numerous studies have documented the main Hg changes in speciation and partitioning where strong redox gradients are present, such as in surficial sediments of lakes and coastal zones (Feyte et al., 2010; Gobeil and Cossa, 1993; Merritt and Amirbahman, 2007). However, the complexity and diversity of Hg transformations are difficult to distinguish in sedimentary deposits where redox changes occur at millimeter or centimeter-scale. On the contrary, they are easily characterized at high resolution in water columns where redox gradients may extend over several meters. Indeed, Hg enrichment and major changes in speciation have been identified at the interfaces between oxic and anoxic waters (Cossa and Coquery, 2005; Guédron et al., 2020b; Iverfeldt, 1988; Lamborg et al., 2008; Mercone et al., 1999; Muresan et al., 2008). The high-resolution (1 cm) study of a

stratified water column in a tropical dam lake, showed prominent Hg methylation intricately 34 linked with Hg⁰ production at the chemocline (Muresan et al., 2018). Both reactions were found 35 to be fueled by nutrients released episodically during the decomposition of settling OM or 36 inhibited by dissolved OM and inorganic compounds continuously transported from the deeper 37 layer. Hence, the chemocline acts as both an accumulation and recycling domain for settling 38 MeHg-loaded organic particles. 39 The water column of meromictic lakes is permanently stratified. Lake Pavin (France) is a good 40 example of such a situation; it offers a unique natural geochemically well-described laboratory 41 42 (Bura-Nakic et al., 2009; Busigny et al., 2016; Jézéquel et al., 2016; Viollier et al., 1997), where 43 Hg transformations can be studied in detail. Only one study has previously described the distribution of the Hg species in Lake Pavin (Cossa et al., 1994). With the analytical tools 44 available at that time, the authors described the MeHg formation at the redoxcline and suggested 45 46 an influence of Fe and Mn recycling on Hg partitioning and speciation. More recently, HgS nanoparticles have been identified in the monimolimnion (Miot et al., 2016). The present study 47 48 is the first to focus on Hg distribution, speciation, and partitioning in the oxic-anoxic transition zone of a meromictic lake using 1 m vertical resolution profiles and which documents their 49 50 relation to Fe, S, and OM redox cycling. In addition, the use of thermodynamic calculations 51 codes on this dataset allowed us to better understand dissolved speciation and solution equilibrium state with respect to mineral phases in order to construct a comprehensive 52 biogeochemical model for Hg cycling in the lake. 53

2. MATERIAL AND METHODS

2.1. Study site

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Lake Pavin (45°55'N, 2°54'E, Fig. 1) is a trachytic maar, located in the Massif Central (France), at an altitude of 1197 m. The lake was formed ~ 6900 years ago (Chapron et al., 2010; Juvigné et al., 1996) during a strong phreatomagmatic explosion (Thouret et al., 2016). The lake is a nearly circular depression of *ca*. 750 m in diameter and 92 m depth, with an area of 0.445 km² and a truncated cone topography. Its watershed is estimated to be 1.24 km² (Michard et al., 1994). The average discharge of Lake Pavin is 50 L.s⁻¹ (Thouret et al., 2016). Because of its geometry, wind protection, and sub-lacustrine springs, the mixing of the Pavin water column is depth-limited (Bonhomme et al., 2016; Busigny et al., 2016). Lake Pavin is a ferruginous meromictic lake, composed of two main layers: (i) the mixolimnion, which extends from the surface to 50-60 m depth and is oxygenated due to the seasonal mixing, and (ii) the permanently anoxic monimolimnion, which extends down to the lake bottom (92 m) and contains very high

concentrations of dissolved elements which contribute to the stability of the layer (Martin, 1985; Michard et al., 1994; Viollier et al., 1997). The residence time of the waters is estimated between about 9 years in the mixolimnion and 100 years in the monimolimnion (Jézéquel et al., 2016). In between, is a transition zone, called the mesolimnion (sometimes considered as the upper part of the monimolimnion), which is characterized by a strong decrease in oxygen and redox potential, and a strong increase in specific conductivity (Lopes et al., 2011; Busigny et al., 2021). It is worth noting that the vertical positions of the oxycline, the redoxcline and the chemocline in the water column may change by a few meters depending on seasons (Bonhomme et al., 2011).

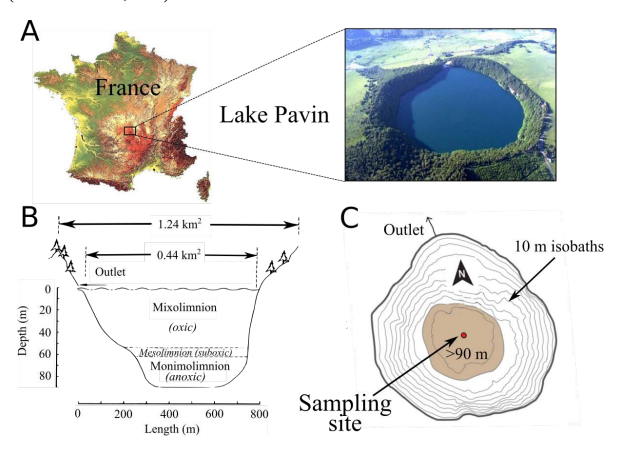


Figure 1. Lake Pavin: A) location in the Massif Central (France) and aerial photography (Credit: D. Jezequel, CNRS Images), B) morphological characteristics of the lake, and C) bathymetric map and sampling site location adapted from Chassiot et al (2016).

2.2. Sample collection and storage

Sampling was performed in July 2018 from the middle of the lake (Fig. 1) on a floating platform using a PFA pneumatic pump (AstiPureTM, Saint-Gobain) with a 100-meter-long polyethylene (0.9 cm diameter) tubing. Samples were collected without any contact with the atmosphere at 1 m depth-intervals in the mesolimnion (52-65 m), and 10 m depth intervals elsewhere. Redox potential (Eh), conductivity, and pH were measured *in situ* with probes connected to a WTW

multi 340i. Filtrations for Hg speciation, anions, major, trace, and redox-sensitive elements were performed on site onto hydrophilic 0.45 µm PTFE filters (OmniporeTM). Filters were then used for the analysis of total particulate Hg (THg_P), and particulate major and trace elements. Filtered and unfiltered samples for total Hg (THg_F and THg_{UNF}) and methylmercury (MeHg_F and MeHgunf) analyses were collected and stored into acid-cleaned FEP Teflon bottles following ultra-clean techniques (Cossa and Gobeil, 2000). In the mesolimnion, additional samples were ultra-filtrated on site on pre-cleaned membranes (<3 kDa, Vivaspin®, polyethersulfone (PES), by centrifugation (6900 rpm, 4°C during 60 min), for the determination of the "truly-dissolved" THg (THg_{UF}) (Guédron et al., 2016). The colloidal part of the THgF, i.e., THg_{coll}, was then obtained from the difference between THg_F and THg_{UF}. All samples for Hg analysis were acidified (0.4 % v/v) on-site with HCl (Optima[®] grade), except those for dissolved gaseous mercury ($DGM = Hg^0 + DMHg$) determinations, which were kept unfiltered in the dark at +4°C until the analysis. Samples for major and trace elements were stored in polyethylene vials and acidified with HNO₃ (1 % v/v, Suprapur[®] grade). Samples for dissolved organic carbon (DOC) analyses were filtered through glass fiber filters (<0.7 µm, GFF). Filters were then cut in equal parts used for the analysis of particulate organic carbon (POC), THgP and particulate methylmercury (MeHg_P). DOC samples were stored in borosilicate vials and acidified with HCl (0.2 % v/v). All samples were wrapped in double bags and kept in the dark at +4 °C until analysis.

2.3. Mercury speciation and partitioning

Filtered, unfiltered, and ultra-filtered THg, and DGM in water were measured by CV-AFS (Tekran®, model 2500) following the published US-EPA Standard Method N° 1631 (Bloom, 1996; Cossa et al., 2003; USEPA, 2002). The detection limit was 0.5 pmol L⁻¹ for a 40-mL water aliquot and the reproducibility varied between 5 and 15 % (N = 6) according to the concentration level in the samples. The accuracy of THg measurements was tested using the certified reference material (CRM) ORMS-5 (certified value = 26.2 ± 1.3 pg g⁻¹) from the National Research Council of Canada. DGM was quantified by purging 500 mL of unfiltered water with Hg-free argon within two hours after sample collection (Cossa et al., 2017). Filtered and unfiltered MeHg (= MMHg + DMHg) were quantified following the derivatization method based on an aqueous ethylation, and purge-trap on a Tenax®, gas chromatography, and cold vapor atomic fluorescence spectrometry (CV-GC-AFS) with a Tekran® detector (model 2700) following the US-EPA 1630 protocol (USEPA, 1998). The precision of analysis was determined by measuring replicates of a control standard (RSD < 10 %, N=20), and the absence

of matrix effect was verified using the standard addition technique (Guédron et al., 2011; 120 Guédron et al., 2017). Total particulate Hg and MeHg (THg_P, MeHg_P) concentrations were 121 determined with the same method as for solution, after the digestion of Hydrophilic PTFE 122 membranes with HNO₃:HCl mixture (2:1 v:v, 70 °C, 48 h) for THg_P, and after the digestion of 123 GF/F membranes with HNO₃ (6 N, 70 °C, 24 h) for MeHg_P. The QA/QC was checked with two 124 CRMs, MESS-3 and IAEA-405, for THg and MeHg respectively. Recovery was $100 \pm 10\%$ for 125 the two methods, and precision was better than 15% for $THg_P(n = 6)$ and better than 5% for 126 127 $MeHg_P(n = 5)$.

2.4. Major, trace elements, and organic matter analyses

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Particulate major and trace elements were quantified on an aliquot of the digestion solution used for the THg_P determination (PTFE filters). Total Fe and Mn concentrations and trace elements (Cu, Cd, Co, Ni, Zn, Pb, Cr) were quantified with an ICP-MS (Thermo[®], X-series 2). Precision was assessed by replicate analysis of CRM SLRS-4, with values better than 7% for all studied elements. Major (Ca, K, Mg, Na, P, S, Si, Sr) were quantified by inductively coupled plasma atomic emission spectrometry (ICP-AES, Varian 720 ES model). The precision was better than \pm 5%. Ferrous iron (Fe^{II}) was determined on-site using the ferrozine method (Viollier et al., 2000) with a portable spectrophotometer. Sulfides ($\Sigma H_2S = H_2S + HS^- + S^{2-}$) samples were measured on-site with a portable spectrophotometer following Cline's method (Cline, 1969) on filtered water stabilized directly after filtration with 0.01 M zinc acetate (Tisserand et al., 2021). Calibration was performed using an external standardization from a solution prepared with Na₂S.9H₂O and titrated by an iodometric method (USGS, 2002). Anions [(i.e., chloride (Cl⁻), sulfates (SO₄²⁻), nitrates (NO₃⁻), and total dissolved phosphates (TDP)] were measured by ion chromatography (IC 332, Metrohm®), with a precision better than 1 %. DOC was measured with a TOC-VCSN analyzer (Shimadzu[®]). POC and was quantified with a mass spectrometer (SERCON®) after combustion of GF/F filters at 1000 °C (Raimbault et al., 1999). The detection limit and precision were estimated at 10 µg and 5%, respectively. Finally, suspended particulate matter (SPM) was reconstructed from the organic and inorganic content of particles collected on filters (Prahl et al., 1997).

2.5. Thermodynamic calculations

WHAM 7 (Tipping, 1994) and PHREEQC 3.0 (Parkhurst and Appelo, 2013) softwares were combined following the method of crossed modeling described in Rigaud *et al.* (2013) for the determination of (i) aqueous Hg^{II} and MeHg speciations, and (ii) mineral phase equilibrium accounting for redox gradient and dissolved OM complexation. The THERMODDEM default

database (https://thermoddem.brgm.fr/) was used and implemented with all reactions involving dissolved Hg and MeHg known in the literature [Tab. SI.1 and 2 (Blanc et al., 2018; Drott et al., 2013; Feyte et al., 2012; Skyllberg, 2008; Smith and Martell, 1976)]. Calculations were firstly performed with WHAM and the calculated apparent thermodynamic constants (K_{app}) of the complexation reactions between aqueous Hg^{II} and MeHg and humic acid (HA) and fulvic acids (FA) were then added to the database for PHREEQC calculations. Temperature, pH, dissolved molar concentrations in dissolved inorganic Hg (THg-MeHg) and MeHg, HA, FA, Hg, Fe, Mn, ΣH₂S, major cations and anions were defined as input parameters in both models (Tabs. SI.3 and 4). HA and FA molar concentrations were calculated from Eq. 1 and 2:

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$$HA_{\text{mol }L}^{-1} = 2 \times 0.1 \times DOC_{\text{mg }L}^{-1} \times 10^{-3} / M_{\text{HA-g mol}}^{-1}$$
 (Eq. 1)

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$$FA_{\text{mol }L}^{-1} = 2 \times 0.9 \times DOC_{\text{mg }L}^{-1} \times 10^{-3} / M_{\text{FA-g mol}}^{-1}$$
 (Eq. 2)

- 164 considering a dissolved organic matter to DOC ratio of 2:1, and a FA:HA mass ratio of 9:1 with
- $M_{HA} = 650 \text{ g.mol}^{-1}$ and $M_{FA} = 2000 \text{ g.mol}^{-1}$ (Buffle and Chalmers, 1988; Malcolm, 1985).
- Saturation indices (SI) were calculated for 47 mineral phases containing Hg, Fe, S, and Mn,
- including 11 Hg mineral phases (Tab. SI.2).

3. RESULTS

3.1. Water column chemistry

Figure 2 illustrates the successions of the redox (Fig. 2A) and physical (Fig. 2B) structures of the water column of Lake Pavin during the sampling period. The mixolimnion was spread between 0 and 53 m depth and the monimolimnion from 53 m down to 92 m including the mesolimnion between 54 m and 65 m. The aerobic limit was observed at 53 m (Fig. 2). The water conductivity was stable in the mixolimnion with an average value of $60 \mu S cm^{-1}$, and increased in the mesolimnion to reach 470 $\mu S cm^{-1}$ in the monimolimnion (Fig. 2C). Major elements concentration profiles in the filtered fraction depicted similar patterns parallel to the conductivity (Fig. SI.1). The pH decreased from 7.0 at the surface to 6.1 at the bottom of the lake, with a marked gradient in the suboxic layer (Fig. 2C). Redox values (Eh) were high in the mixolimnion, and started to decline with increasing depth from 40 m down to 58 m, *i.e.*, the bottom of the suboxic zone (Fig. 2C). Suspended particulate matter (SPM) was quite homogenous and low in the mixolimnion, and increased markedly around the oxycline as well as at the bottom of the lake in the nepheloid layer (Busigny et al., 2014) up to 0.96 mg L⁻¹ (Fig. 2D). DOC concentrations (Fig. 3A) averaged 90.4 \pm 13.6 μ mol L⁻¹, and increased gradually

from the top of the monimolimnion up to 354 μ mol L⁻¹ at the bottom of the lake. POC concentrations increased in the mesolimnion from an average concentration of $11.5 \pm 5 \mu$ mol L⁻¹ in the mixolimnion to $21.4 \pm 2 \mu$ mol L⁻¹ in the monimolimnion. The high value at 90 m (~ 39 μ mol L⁻¹) suggests that particles from the nepheloid layer were sampled. Total dissolved phosphates (TDP) were only measurable below 65 m depth and increased rapidly in the monimolimnion up to 0.5 mmol L⁻¹ at 70 m depth. Particulate phosphorus (P_P) increased steadily from negligible values above 54 m (*i.e.*, above the oxycline), up to 0.3 mmol L⁻¹ at the lake bottom (Fig. 3B).

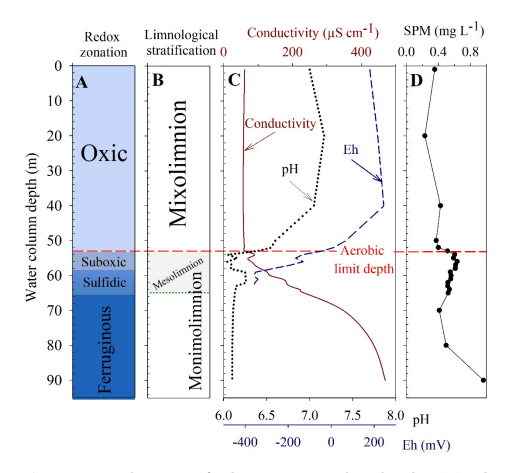


Figure 2. Vertical zonation of Lake Pavin water column based on (A) redox domains, (B) water limnological stratification, (C) electrical conductivity, pH, redox potential (Eh), and (D) suspended particulate matter (SPM) concentrations measured in July 2018.

Consistent with the definitions by Froelich et al. (1979), Poultron and Canfield (2011), and Busigny et al. (2016), four redox zones were identified in the water column of Lake Pavin (Fig. 2A): (i) the oxic zone where molecular oxygen is still present (0-53 m); (ii) the suboxic zone without molecular oxygen, but where other electron acceptors, such as Fe^{III}, Mn^{IV}, or S^{VI}, are present (54-59 m); (iii) the sulfidic zone where the Σ H₂S is abundant (60-65 m); and (iv) the ferruginous layer where Fe^{II} prevails (65-90 m). Zone (i) belongs to the mixolimnion, zones (ii)

and (iii) to the mesolimnion, and zone (iv) to the monimolimnion. Peaks of Fe_P and Mn_P are encountered above and at the interface between the oxic and suboxic layer, respectively, presumably because of precipitation of Fe and Mn diffusing upwards. There, neoformed oxy(hydr)oxides may have adsorbed dissolved organic matter and contribute to the POC peak. In the suboxic zone, Mn_P and Fe_P (Fig. 3C and D) started to decrease successively from 52 and 54 m depth, respectively. Consistently, dissolved Mn and Fe concentrations showed a mirrored increase in the form of reduced species as all dissolved Fe is found in its reduced form (Fe^{II}). The reduction of SO₄²⁻ started below 54 m, Σ H₂S appeared at 58 m and peaked up to 11 μ mol L⁻¹ around 62 m (Fig. 3E). It is worth mentioning that the sulfidic zone (zone with Σ H₂S) was identified between 58 and 65 m, i.e., overlapping the onset of the ferruginous zone. These phase changes are also consistent with the conventional succession of redox biogeochemical reactions along the redox gradient, which trigger reductive dissolution of the

These phase changes are also consistent with the conventional succession of redox biogeochemical reactions along the redox gradient, which trigger reductive dissolution of the oxy(hydr)oxides (Froelich et al., 1979), as already noticed in Lake Pavin by Viollier et al. (1997). Because of the limited amount of sulfur in the lake, ferruginous conditions set in the monimolimnion confers the singular characteristic to Lake Pavin (Bura-Nakic et al., 2009).

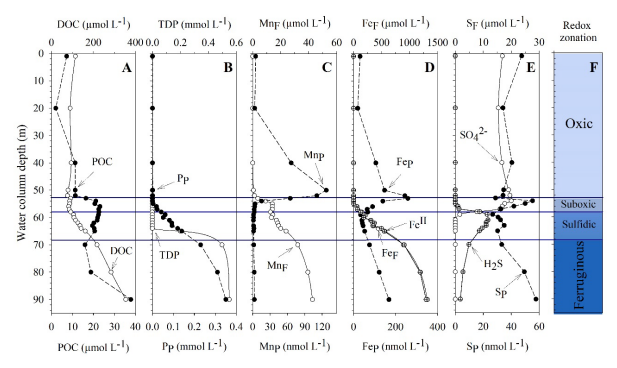


Figure 3. Depth concentration profiles of dissolved and particulate (A) organic carbon, (B) phosphorus, (C) manganese, (D) iron, and (E) sulfur (sulfate, and sulfides) in the water column of Lake Pavin. Subscript F refers to filtered samples and subscript P to the particulate phase. The light blue band indicates the oxic zone, the light-dark blue band the suboxic zone, the blue one the sulfidic zone, and the dark blue band the Ferruginous zone. Fe^{II} is the reduced iron part of aqueous $Fe(Fe_F)$.

This ferruginous zone consists of anoxic conditions where Fe is mobilized as ferrous species (Fe^{II}), and was shown to be subsequently precipitated mainly as siderite and vivianite (Busigny

et al., 2016; Cosmidis et al., 2014; Poulton and Canfield, 2011; Viollier et al., 1997). Near the bottom, the nepheloid layer comprises strong elemental concentration gradients that reflect the upward diffusion of dissolved ions from the surficial sediment (Busigny et al., 2014).

3.2. Mercury speciation profiles

Total Hg concentrations in Lake Pavin waters are in the picomolar range, one order of magnitude higher than the ultra-trace oceanic concentration range (Bowman et al., 2020), but similar to other pseudo-pristine lakes, *i.e.*, lakes without local anthropogenic Hg inputs (Guédron et al., 2017; Leermakers et al., 1996; Regnell et al., 1997). THg_F ranged from 0.4 to 8.8 pmol L⁻¹ and THg_P from 0.4 to 4.6 pmol L⁻¹. Both THg_P and THg_F showed the lowest concentrations in the oxic mixolimnion and the highest ones in the mesolimnion (Fig. 4A).

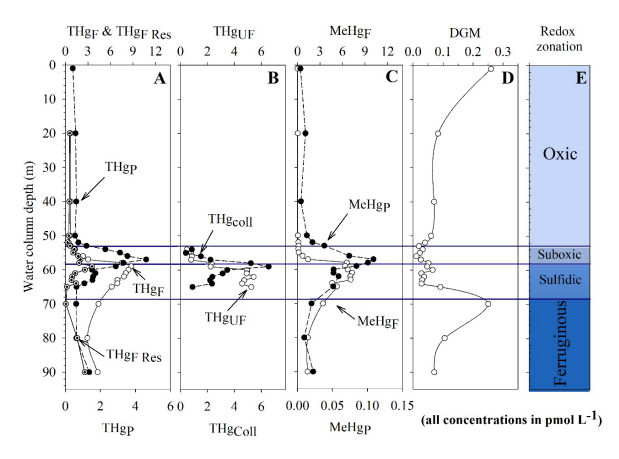


Figure 4. Depth concentration profiles of filtered (upper X-axis) and particulate (bottom X-axis) fractions (A) total Hg filtrated (THg_F), particulate (THg_P) and residual Hg ($THg_{FRes} = THg_F - MeHg_F$), (B) ultra-filtered (THg_{UF}) and colloidal Hg (THg_{coll}), (C) dissolved gaseous Hg (DGM), and (D) methylmercury (MeHg). Corresponding redox zonation is also reported.

THg_P concentrations reached their maxima at 57 m, whereas THg_F maximum was found below, at 59 m. Then, they both show a progressive downward decrease in the sulfidic zone of the mesolimnion (Fig. 4A). In the mixolimnion, THg_F represented $\sim 50\%$ of THg (i.e., THg_P + THg_F) and decreased abruptly to reach 27% at 55 m in the upper part of the suboxic zone of the

mesolimnion, followed by a gradual increase up to 75% at the bottom of this zone. Lateral 244 advection of Hg from underground water inputs (few liters per second) present at mid-depth in 245 the lake are negligible since a few measurements (data not shown) performed in the Goyon 246 spring (45°30'34.7" N; 2°54'15.6" E) waters indicated THg and MeHg concentrations at the 247 picomolar level. Below, in the sulfidic part of the mesolimnion, the proportion of the THg_F 248 increased gradually to reach an average percentage of 85% of THg. Within the filtered fraction, 249 the "truly dissolved" THg (THg_{UF}: 0.5-5.4 pmol L⁻¹) and colloidal part of the THg_F (THg_{coll} = 250 0.4-6.5 pmol L⁻¹) were quantitatively comparable. In the suboxic zone, THg_{coll} presented a sharp 251 peak at 59 m which dominated (i.e., $63 \pm 15\%$) the THg_F fraction, and decreased in the sulfidic 252 zone (32 \pm 9% of THg_F) to the benefit of THg_{UF} (Fig. 4B). Noteworthy is that the maxima of 253 the various THg fractions follow each other within a few meters between 57 and 60 m 254 deepening from particulate, colloidal, <0.45 µm, and truly dissolved. This suggests a 255 256 progressive dissolution of Hg downward. MeHg_F ranged from 0.03 to 6.8 pmol L⁻¹, whereas MeHg_P concentrations were two orders of 257 258 magnitude lower. Its distribution resembles the one of THg, with low concentration in the oxic mixolimnion and higher ones in the anoxic monimolimnion, where it reaches up to 7.1 pmol L⁻ 259 260 ¹ in the sulfidic zone of the lake (Fig. 4C). MeHg_P peaked at 57 m whereas MeHg_F showed a broader maximum between 58 and 63 m (Fig. 4C). MeHgF represented a minor fraction of THgF 261 in the oxic mixolimnion (< 20%), and this proportion increased dramatically in the suboxic and 262 anoxic monimolimnion, reaching 90% between 62 and 70 m, and then decreased below 40% at 263 the bottom of the lake. Overall, MeHg_F, including the colloidal phase, is clearly the dominant 264 Hg species in the sulfidic zone. In contrast, MeHgP never exceeded 10 % of the THgP with the 265 highest proportions at 62 m. 266 Finally, DGM was between one and two order of magnitude lower than THg with the highest 267 concentrations found at the surface (0.26 pmol L⁻¹) and in the middle of the monimolimnion 268

3.3 Crossed modeling using WHAM 7 and PHREEQC 2.0

 $(0.25 \text{ pmol L}^{-1} \text{ at } 70 \text{ m}, \text{Fig. 4C}).$

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According to the thermodynamic calculations with WHAM 7 and PHREEQC 2.0, in the mixolimnion, Hg and MeHg speciation is dominated by complexes with organic ligands (i.e., 98% for Hg-FA and up to 87% MeHg-FA of the total MeHg) (Fig. 5 A and B). However, when dissolved sulfide is present, in the sulfidic zone of the lake, complexes with sulfur ligands dominate (i.e., > 65% of HgS and about 100% of MeHgSH). Calculated saturation indexes indicated the saturation of iron oxides, mostly hematite (Fe₂O₃) and goethite (FeOOH), in the

suboxic part of the lake (Fig. 5C), whereas mineral phases of Fe, S, and P, including crystallized FeS (i.e., mackinawite) and vivianite (Fe₃(PO₄)₂.8H₂0), are predicted to be saturated to oversaturated from the top of the sulfidic zone to the lake bottom (Fig. 5D).

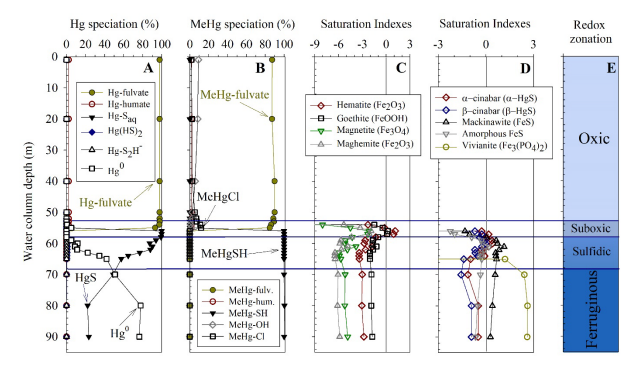


Figure 5. Modeled (A) Hg speciation (= THg-MeHg) including Hg^{II} and Hg^0 species, (B) MeHg speciation, and saturation indexes for (C) iron oxy(hydr)oxides and (D) sulfide and vivianite minerals (WHAM 7.0 and PHREEQC 2.0). Pyrrhotite saturation index was below -100 and is not presented in the figure. Corresponding redox zonation is also reported.

These results are consistent with previously published data (Bura-Nakic et al., 2009; Busigny et al., 2016; Jézéquel et al., 2016; Viollier et al., 1997). Interestingly, cinnabar (alpha and beta) minerals were found to be saturated in the upper part of the sulfidic zone (Fig. 5D), while all other Hg mineral species tested (Tab. SI.2), were found to be under-saturated.

4. DISCUSSION

4.1. Mercury carrier-phase exchanges along the redox gradient

In oxic waters, potential solid Hg carrier phases are limited due to the low mineralogical diversity of SPM in the upper part of Lake Pavin (Miot *et al.*, 2016). In surface waters, the biogenic particulate matter, mainly composed of diatoms and bacteria (Amblard and Bourdier, 1990; Miot et al., 2016), constitutes the sink for atmospheric Hg deposited onto the surface of the lake. Known as the "biological pump" (Hain et al., 2014), this process is common for removing trace elements (including Hg) from surface waters of oceans and lakes, especially when it contains diatoms (Zaferani et al., 2018). This interpretation is supported by the positive relationship between THg_P and POC in the mixolimnion ($R^2 = 0.75$, Fig 6A). THg_P is also found

correlated with Fe_P ($R^2 = 0.90$, Fig 6A) which suggest a possible implication of the Fe oxides in the Hg-particulate organic matter (POM) consortiums. THg_F in the mixolimnion is also correlated with DOC (Fig. 6B), consistently with the predictions of our modeling, which shows that Hg-FA complexes dominate Hg species (98%) in the filtered phase (Fig. 5). The Hg-DOC association in natural waters is often thought to be the result of the high affinity of Hg for thiol functional groups of the DOC (Bouchet et al., 2018; Haitzer et al., 2002; Skyllberg et al., 2003).

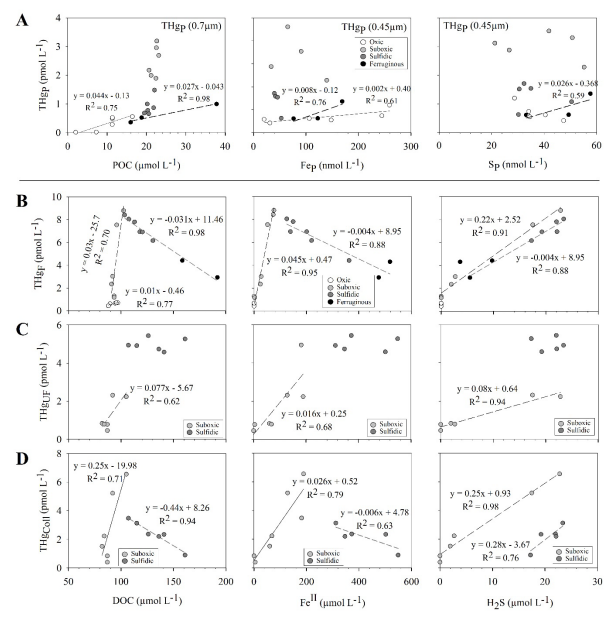


Figure 6. Total Mercury concentration in the A) the particulate (THg_P) , B) the filtered $(THg_F < 0.45 \mu m)$, C) the truly dissolved (THg_{UF}) , and D) the colloidal (THg_{coll}) fractions versus organic carbon, iron and sulfide in the particulate (A) and filtered (B to D) fractions. Left panel of plot A reports particulate mercury (THg_P) and organic carbon (POC) obtained from the digestion of 0.7 μ m (GF/F) filters. Central and right panels report particulate mercury (THg_P) , iron (Fe_P) and sulfur (S_P) concentrations obtained from the digestion of 0.45 μ m (PTFE) filters. All plotted regressions are significant and have p values below 0.05.

The vertical downward POM flux is slowed down when reaching the density gradient in the 313 314 mesolimnion and accumulated as indicated by the increase of THg, POC and SPM. There, POM feeds the heterotrophic microorganisms, resulting in dissolved O₂ consumption as previously 315 described by Michard et al. (1994). 316 317 In the mesolimnion, no significant correlation was found between THg_P and POC, nor with concentration of particulate Fe or S. This likely reflects the recycling of Hg onto successive 318 neoformed phases along the redox gradient (i.e., POC mineralization, reductive dissolution of 319 Mn and Fe oxy(hydr)oxides, Cf section 3.1). Changes in Hg carrier phases is particularly 320 321 evident in the filtered phase, as peaks in THg_F and MeHg_F occur below the onset of the decay of both THg_P and MeHg_P, i.e., from the boundary between suboxic and sulfidic zone (Fig. 4). 322 323 Consistently with the recycling of POC and Mn or Fe oxides, both the truly dissolved Hg (THg_{UF}) and the colloidal Hg (THg_{coll}) are found positively correlated with DOC, Fe^{II} and Σ H₂S 324 in the suboxic zone (Fig. 6 C and D). This confirms that the mineralization of POM and the 325 reductive dissolution of Fe-oxy(hydr)oxides below the oxycline have resulted in the release of 326 dissolved Hg and Fe^{II} (Hellal et al., 2015), and their redistribution in the filtered fraction, with 327 a dominance of Hg species bound to OM and/or FeS (i.e., mackinawite, Fig. 5D) colloids 328 $(THg_{coll} = 63 \pm 15\% \text{ of } THg_F)$. Below, in the sulfidic mesolimnion, our modeling predicted the 329 330 formation of both α- and β-cinnabar and FeS mineral phases (i.e., mackinawite and other amorphous FeS phases, Fig. 5D) consistently with previous studies (Bura-Nakic et al., 2009). 331 In the filtered phase, truly dissolved THg (THg_{UF} = $68 \pm 9\%$ of THg_F) becomes dominant, and 332 is found independent of DOC, Fe^{II} and ΣH_2S (Fig. 6C). In contrast, THg_F and THg_{coll} are 333 inversely correlated with DOC and Fe^{II} , but positively correlated with ΣH_2S (Fig. 6 B and D). 334 This most probably illustrates the complex formation of Hg with sulfides in solution (i.e., HgS 335 nanoparticles and/or dissolved Hg-polysulfides) at the expense of OM complexation as 336 confirmed by our model (Figs. 5A, 5B), and consistently with colloidal HgS identified by 337 Energy dispersive X-ray spectrometry in deep-water of the Lake Pavin (Miot et al., 2016). 338 Although the thermodynamic modeling does not allow modeling colloidal HgS, the model 339 predicts the precipitation of HgS particles at the top of the sulfidic mesolimnion (Fig. 5D). 340 Hence, during the degradation of the POM in suboxic and anoxic conditions, the nature of the 341 342 particulate Hg-OM binding sites change, favoring the preferential binding of Hg to stronger dissolved ligands, such as thiol groups of the FA (Bouchet et al., 2018; Feyte et al., 2012; Liang 343 et al., 2019) and the formation of FeS and HgS colloids in the dissolved phase, which is at 344 equilibrium with HgS minerals. 345

In the ferruginous monimolimnion, although the number of observations is limited (n = 3), 346 THg_P rise with rising POC, Fe_P and S_P concentrations (Fig. 6A). This is consistent with an 347 association of Hg with both iron sulfides, and/or sulfidized OM, consistent with the predictions 348 of the model. It is however important to recall that this area is the most devoid of particulate 349 Hg. In this ferruginous zone, Fe_P and P_P covary ($R^2 = 0.95$, n = 3, p < 0.10), with 1.5 mole of Fe 350 per mole of P, supporting the presence of vivianite as already identified in deep waters of Lake 351 Pavin (Busigny et al., 2016; Cosmidis et al., 2014) and confirmed by our thermodynamic 352 modeling (Fig. 5D). It is thus probable that such Fe-phosphates precipitates together with other 353 FeS minerals, in association with bacteria (Miot et al., 2016) or diatoms (i.e., THg_P vs Si_P, R²= 354 0.64, n = 6, p < 0.05), could favor the scavenging and export of Hg to the sediment of Lake 355 356 Pavin. In brief, the Hg-OM partition appears to be involved in the Fe-S redox and partition changes known as the "iron-wheel" of the Lake Pavin (Busigny et al., 2016; Viollier et al., 357 1997). 358

4.2. Mercury methylation/demethylation

359 MeHg in Lake Pavin is in similar proportions as in other meromictic lakes or artificial reservoirs 360 in various lakes around the world (Peretyazhko et al., 2006b; Watras and Bloom, 1994). In the 361 Great Salt Lake, the Petit-Saut reservoir, and Wisconsin or Canadian lakes, all stratified with 362 an anoxic hypolimnion, comparable MeHg distribution patterns have been observed resulting 363 from increased net Hg methylation in the lake bottom waters (Achá et al., 2012; Eckley et al., 364 2005; Eckley and Hintelmann, 2006; Muresan et al., 2008; Porcella, 1994; Regnell et al., 1997; 365 Regnell and Tunlid, 1991; Watras and Bloom, 1994; Yang et al., 2020). The same is true for 366 367 marine environments where meromixis occurs (e.g., Black Sea and the Baltic Sea), resulting in high MeHg in their anoxic waters (Pempkowiak et al., 1998; Rosati et al., 2018; Soerensen et 368 al., 2018). 369 In Lake Pavin, MeHg is strongly correlated with THg both in the filtered phase ($R^2 = 0.92$, n =370 21, p < 0.01), and in the particulate phase ($R^2 = 0.85$, n = 21, p < 0.01). In the monimolimnion, 371 MeHg constitutes the main fraction of the THg (i.e., 77 ± 22 % of the THg_F, and 71 ± 24 % of 372 the THg_P). MeHg_F is highly correlated with Σ H₂S (R² = 0.96, n = 20, p<0.01) along the entire 373 water column. The onset of MeHg accumulation occurs in the suboxic zone, and reaches a 374 plateau in the sulfidic layer where ΣH_2S and Fe^{II} are the most intensively produced. The steepest 375 MeHg_F gradient is located between 57 and 58 m (Fig. 4D), at the same depth as the steepest 376 SO₄/ΣH₂S drops (Figs. 3D and 3E). This feature indicates that MeHg is dominantly produced 377 378 by methylating-SRB, a group of strains reported owning the methylating hgc gene (Paranjape

and Hall, 2017; Regnell and Watras, 2018). SRB account for a large part of the bacteria in the 379 entire monimolimnion of Lake Pavin (Berg et al., 2019; Lehours et al., 2016). The presence of 380 high MeHg_F and THg_{UF} ($R^2 = 0.84$, n = 12, p < 0.01) concentrations in the sulfidic zone (Fig. 381 4B, 4D) strongly suggests that HgS complexes, neutral Hg polysulfides, and possibly Hg-382 thiolates (Fig. 5A, 5B), are abundant enough to constitute substrates for Hg methylation by SRB 383 as suggested in several experiments or modeling (Benoit et al., 1999; Schaefer and Morel, 2009; 384 Skyllberg, 2008). On the other hand, several strains can demethylate MeHg under anoxic 385 conditions including SRB and methanogens (Bridou et al., 2011; Oremland et al., 1991). The 386 387 decrease in MeHg_F concentrations in the ferruginous zone is coincident with the increase in DGM, which supports the hypothesis of reductive demethylation in the Lake bottom. However, 388 389 the amount of produced DGM could only explain a small fraction (i.e. <5-10%) of the decrease of MeHgF observed, suggesting that significant proportion of MeHgF is also likely exported to 390 391 the sediment in association neoformed FeS minerals (see paragraph 5.1.).

4.3. Mercury reduction

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The DGM concentration profile in the Lake Pavin waters column exhibits high levels in surface waters and at 70 m in the anoxic monimolimnion (Fig. 4C). No trace of dimethylmercury was found in the samples, consistently with previous results (Cossa et al., 1994), suggesting that DGM consists entirely of Hg⁰. The high DGM measured in surface waters suggests that Hg⁰ is formed via Hg^{II} photoreduction in the euphotic zone, and is partially reinjected in the atmosphere as already shown in numerous lakes (Amyot et al., 1994; Guédron et al., 2020a; Saiz-Lopez et al., 2018). In contrast, the presence of DGM peaks in the aphotic monimolimnion supports non-photochemical processes for DGM production, such as Hg^{II} microbial reduction (Lamborg et al., 2013; Mason et al., 1995; Rolfhus and Fitzgerald, 2004) and/or natural OMmediated reduction of Hg^{II} (Allard and Arsenie, 1991; Zheng et al., 2011). Hg⁰ is consistently predicted by the model, increasing from the sulfidic mesolimnion to the ferruginous monimolimnion (Fig. 5A). The large proportion of truly dissolved Hg (i.e., 59 and 81% of THg_F) possibly bound to labile organic molecules could favor its reduction by natural OM (Gu et al., 2011; Zheng et al., 2011). Alternatively, the abiotic reduction of Hg^{II} to Hg⁰ could also be induced by Fe^{II} in the presence of Fe-oxy(hydr)oxides (Charlet et al., 2002; Peretyazhko et al., 2006a), such as hematite reported from 70 m down to 86 m in the Pavin Lake (Cosmidis et al., 2014). Hg reduction in the presence of nanoparticulate vivianite has also recently been demonstrated in anoxic circumneutral pH water (Etique et al., 2021). Also, the reduction of HgII-S-II complexes by FeS to Hg⁰ is thermodynamically possible (Bone et al., 2014) at Eh values in the range of Lake Pavin waters below 59 m. However, the occurrence of DGM peaks in the monimolimnion in phase with decreasing MeHg concentrations (Fig. 4) also supports the hypothesis of reductive MeHg demethylation with the production of Hg⁰ and CH₄ (Aeschbach-Hertig et al., 1999). Reductive demethylation is the fact of *Geobacter* sp. (Lu et al., 2016), some of them identified in Lake Pavin waters (Berg et al., 2019). In summary, under the current observations of Pavin waters, it is not possible to sort out the actual reactions involved in the reducing process that produce the DGM in the monimolimnion. In the present state, we cannot explain the difference between the measured and model-derived DGM concentrations. An indepth study with higher resolution lake bottom profiles together with experimental works are required to solve this issue.

5. SYNTHESIS AND CONCLUSIONS

The high-resolution observation and modeling of Hg speciation and partition at the redox interfaces of Lake Pavin allow the proposal of a conceptual biogeochemical behavior of Hg which can be typical of other meromictic ferruginous lakes (Fig. 7). Because it has a very small catchment area that minimizes Hg inputs from the drainage basin, Lake Pavin receives Hg mainly from the atmosphere. Once deposited onto surface waters, a part of Hg is photo-reduced and reinjected in the atmosphere, whereas another part binds to DOC and sorbs onto biogenic particulate matter produced in the euphotic zone and conveyed at depth by settling POC, in a process known as "biological pumping".

The Hg associated with POM accumulates where the conductivity gradient generates a strong

The Hg associated with POM accumulates where the conductivity gradient generates a strong stratification. There, the mineralization of OM occurs, consuming dissolved oxygen and generating a suboxic zone, causing the mobilization of a fraction of Hg. Mercury diffuses up and down from this Hg-enriched zone. On its way up, Hg adsorbs onto Fe-oxy(hydr)oxides (associated with POM) that precipitated on the top of the oxycline; on its way down, Hg diffuses as truly-dissolved and colloidal HgS species and partly precipitate as HgS minerals. Simultaneously, in this suboxic zone, available Hg^{II} is methylated by SRB. Deeper in the anoxic monimolimnion, sulfidation favors the formation of dissolved Hg fraction and possibly the reductive demethylation of MeHg with Hg⁰ production. A part of the Hg and MeHg produced in the suboxic and sulfidic zone is also likely exported to the sediment with settling particles. In brief, the "biological pump", the "iron-sulfur wheel", through associated bacteria consortia, control the Hg cycling in the Pavin waters. This behavior is in line with already existing biogeochemical models and allows us to observe the already identified homogenous and heterogeneous reactions and processes along a single water column covering the entire redox

range of natural waters. More detailed observations on the Hg speciation in the ferruginous monimolimnion are needed to evaluate the possible effect of Hg input from hydrothermal and diffusive inputs at the bottom of the lake in addition to quantitative export fluxes and reaction rates within the lake.

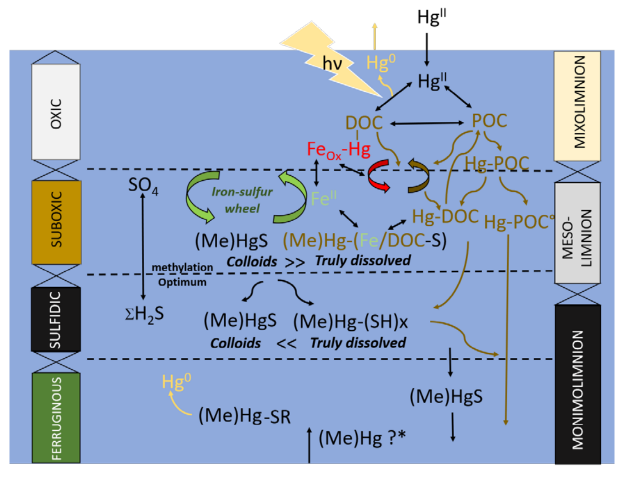


Figure 7. Conceptual model for the Hg and MeHg cycle in Lake Pavin with the recycling of organic matter, sulfur, and iron. (Me)Hg refers to both MeHg and Hg species, ticks (-) indicate the binding with particulate carrier phases (i.e., iron oxides (Fe_{Ox}), particulate organic carbon (POC) and iron sulfides (FeS) or binding with aqueous ligands (i.e., dissolved organic carbon (DOC), free or DOC polysulfides (SR). Hg-POC° considers Hg bound to both particles and microbial cells. (Me)Hg* refers to possible upward release of MeHg and THg from the sediment.

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- 471 Terminology of mercury chemical species:
- 472 THg: Total Hg (all Hg species);
- 473 MMHg: Monomethyl Hg;
- 474 DMHg: Dimethyl Hg;
- 475 MeHg: Methylated Hg (MeHg = MMHg + DMHg);
- 476 Hg^0 : Elemental Hg;
- 477 Hg^{II}: Divalent Hg;
- 478 DGM : Dissolved Gaseous Hg (DGM = $Hg^0 + DMHg$);
- 479 THg_F: Total Hg in filtrated (0.45 μm);
- 480 THg_P: Total particulate Hg (>0.45μm);
- 481 MeHg_F: Methylated Hg in filtrated (0.45 μm);
- 482 MeHg_P: Methylated particulate Hg (>0.45μm);
- 483 THg_{UF}: Truly dissolved Hg, *i.e.*, ultrafiltered samples (< 3 kDa);
- 484 THg_{coll}: Colloidal Hg, *i.e.*, fraction > 3 kDa and $< 0.45 \mu m$;
- 485 THg_{FRes} : THg_{F} $MeHg_{F}$;
- 486 $THg_{P Res}$: $THg_{P} MeHg_{P}$.

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