Mercury in the Black Sea: New Insights From Measurements and Numerical Modeling

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Abstract Redox conditions and organic matter control marine methylmercury (MeHg) production. The Black Sea is the world’s largest and deepest anoxic basin and is thus ideal to study Hg species along the extended redox gradient. Here we present new dissolved Hg and MeHg data from the 2013 GEOTRACES MEDBlack cruise (GN04_leg2) that we integrated into a numerical 1-D model, to track the fate and dynamics of Hg and MeHg. Contrary to a previous study, our new data show highest MeHg concentrations in the permanently anoxic waters. Observed MeHg/Hg percentage (range 9–57%) in the anoxic waters is comparable to other subsurface maxima in oxic open-ocean waters. With the modeling we tested for various Hg methylation and demethylation scenarios along the redox gradient. The results show that Hg methylation must occur in the anoxic waters. The model was then used to simulate the time evolution (1850–2050) of Hg species in the Black Sea. Our findings quantify (1) inputs and outputs of HgT (~31 and ~28 kmol yr\(^{-1}\)) and MeHgT (~5 and ~4 kmol yr\(^{-1}\)) to the basin, (2) the extent of net demethylation occurring in oxic (~1 kmol yr\(^{-1}\)) and suboxic water (~6 kmol yr\(^{-1}\)), (3) and the net Hg methylation in the anoxic waters of the Black Sea (~11 kmol yr\(^{-1}\)). The model was also used to estimate the amount of anthropogenic Hg (85–93%) in the Black Sea.

1. Introduction

The ubiquitous presence of mercury (Hg) in the environment is a matter of concern due to the transformation of inorganic oxidized Hg (Hg\(^{\text{II}}\)) into methylmercury (MeHg), which is a biomagnifying neurotoxin (Clarkson & Magos, 2006; Cossa et al., 2012; Fitzgerald et al., 2007; Hammerschmidt & Fitzgerald, 2006). As humans are exposed to MeHg through fish consumption (Oken et al., 2012) understanding the mechanisms underlying MeHg production, degradation, transport, and accumulation in the marine environment is crucial.

Hg methylation is thought to be a primarily microbially driven process that occurs in marine sediments (Bouchet et al., 2013; Hammerschmidt et al., 2004; Hollweg et al., 2009, 2010; Monperrus, Tessier, Point, et al., 2007; Schartup et al., 2013) and the marine water column (Cossa et al., 2009; Heimbürger et al., 2010; Mason & Fitzgerald, 1990; Monperrus, Tessier, Point, et al., 2007; Monperrus, Tessier, Amouroux, et al., 2007; Sunderland et al., 2009) during remineralization of natural organic matter (NOM). NOM, which includes dissolved and particulate organic matter (i.e., DOM and POM, respectively), affects Hg methylation both acting as an electron donor for microbial activity and controlling Hg partitioning (Drott et al., 2007). Indeed, in the marine environment, Hg is effectively scavenged and transported by POM and also by authigenic Fe hydroxide and Mn oxide minerals (hereafter Fe/Mn oxides) and lithogenic particles (Lamborg et al., 2016). While sinking, particles undergo degradation, which prompts the release of dissolved Hg\(^{\text{II}}\) making it bioavailable to microbes (Cossa et al., 2009; Heimbürger et al., 2010; Muresan et al., 2007; Sunderland et al., 2009). Biogeochemical controls on Hg methylation rates are thus related both to the composition and the activity of the microbial community and to the speciation and partitioning of Hg\(^{\text{II}}\) (Cossa et al., 2014; Merritt & Amirbahman, 2009), with a critical role played by redox conditions, NOM availability, and quality (Bravo et al., 2017; Cossa et al., 2014; Schartup, Ndu, et al., 2015). Previous studies have used chemical equilibrium...
modeling to explain the observed oxic or suboxic MeHg maxima in sediment and stratified water. These studies, which include a previous assessment of Hg species in the Black Sea water column, have concluded that methylation is hindered above sulfide concentrations of ~10 μM due to a shift of Hg speciation from neutral to charged species that makes it unavailable to microbes (Benoit, Gilmour, et al., 1999; Benoit, Mason, & Gilmour, 1999; Benoit et al., 2001; Lamborg et al., 2008). However, there is a lack of consensus on the relevance of the methylation process under different redox conditions. Hg methylation is known to be mediated by anaerobic bacteria (Gilmour et al., 2013); however, methylation in euxinic (i.e., anoxic and sulfidic) environments is not a well-understood process (Hsu-Kim et al., 2013; Merritt & Amirbahman, 2009), which has been shown to occur in sediments and lake water (Drott et al., 2007; Eckley & Hintelmann, 2006; Hollweg et al., 2009; King et al., 2000). The occurrence of Hg methylation in oxic waters was not discussed by Lamborg et al. (2008) for the Black Sea but has been shown to occur in other basins (Lehnerr et al., 2011; Monperrus, Tessier, Amouroux, et al., 2007; Schartup, Balcom, et al., 2015; Sharif et al., 2014). MeHg demethylation seems to occur at less variable rates among various environments, and the capacity of bacteria to demethylate seems to be more widespread (Heyes et al., 2006).

The Black Sea is ideal for investigating biogeochemical cycles under changing redox conditions because its waters encompass oxic, suboxic, and anoxic conditions (Schijf et al., 1991), with gradients stretched over several tens of meters. The Black Sea is a deep, semienclosed basin (Figure 1a, maps are plotted with Ocean Data View (Schlitzer, 2016), with permanently stratified water owing to the hydrological balance between the inflow of water from the Mediterranean Sea via the Marmara Sea-Bosporus system and freshwater inputs (Figure 1b and Text S1 in the supporting information) (Özsoy & Ünlüata, 1997). The surface circulation of the Black Sea is driven by two large cyclonic gyres in the eastern and western basin, which are connected to each other by the Rim Current that homogenizes surface waters (Murray et al., 2007). Vertical mixing is weak, and when oxygen depletion occurs, bacterial remineralization of NOM is driven by a sequence of alternative electron acceptors according to a sequence that is fixed across systems ($O_2 > N_2 > MnO_x > FeO_x > SO_4$) (Froelich et al., 1979). Hence, the redox progression along the Black Sea water column is similar to the one in sedimentary environments and hypoxic basins worldwide, with the advantage that the redox layers of the Black Sea are quite stable throughout the year and are stretched over a scale of several tens of meters; these are thus easier to discern.
The sequence of redox reactions driving NOM remineralization in the water column is mirrored by the concentration profiles of the electron acceptors (Figure 2). The upper part of the oxic layer (OL) is euphotic (~50 m depth) with high O₂ concentrations (~300 μM), while below, in the aphotic waters, O₂ is gradually consumed by nitrifying bacteria (oxycline), as revealed by increasing NO₃⁻ concentrations (Oguz et al., 2000). The suboxic layer (SOL) at a depth of approximately 100 m is where concentrations of oxygen and sulfides are extremely low (O₂ < 20 μM, HS⁻ < 1 μM). The upper boundary of the anoxic layer (AOL) occurs at a depth of approximately 140 m, and sulfide levels increase within the AOL down to the seabed (2,200 m depth) (Murray et al., 2007). The boundaries between the redox layers computed from the cruise data set are shown in each panel (OL = oxic layer; SOL = suboxic layer; AOL = anoxic layer).

This paper aims to understand Hg speciation along a redox gradient by focusing on the pathways of MeHg production and transport. We integrated novel marine Hg and MeHg observations from the 2013 GEOTRACES MEDBlack cruise with a fate and transport 1-D model for Hg species (Melaku Canu et al., 2015) adapted from the
WASP7 Hg model /MERC7 sub-model (Wool et al., 2001) and already successfully applied in shallow waters (Canu & Rosati, 2017; Melaku Canu et al., 2015). One-dimensional (1-D) modeling has widely been used to investigate the biogeochemical and ecological features of the basin (Grégoire & Soetaert, 2010; Konovalov et al., 2004; Lamborg et al., 2008; Oguz et al., 2000; Yakushev et al., 2007). The implementation of the WASP7 model requires a careful compilation of Hg fluxes within the basin, which includes exchanges with the Marmara Sea, rivers and the atmosphere, and water-sediment exchanges and burial (Figure 3). We computed the budget of Hg species and used the model to assess the occurrence of Hg methylation and MeHg demethylation along the extended redox gradient of the Black Sea water column. Furthermore, we ran additional simulations to test the hypothesis that Hg methylation occurs exclusively in suboxic waters and is inhibited in the anoxic waters, as proposed by Lamborg et al. (2008). The authors detected a MeHg maximum in the SOL and very low levels in the AOL during a previous assessment of Hg species dynamics in the Black Sea. The model was also used to estimate the amount of anthropogenic Hg in the Black Sea water.

2. Materials and Methods

2.1. Sampling and Laboratory Analysis

Samples were collected during the second leg of the 2013 GEOTRACES MEDBlack (GA04-leg2) cruise in the Black Sea. The research vessel Pelagia occupied 12 full-depth stations in the Black Sea along an east-west transect between 13 and 25 July 2013. High-resolution vertical profiles, including one ultrahigh-resolution...

Figure 3. Overview of Hg species and processes simulated in the WASP7 MERC model. HgT is total Hg ($\text{Hg}^{\text{II}} + \text{Hg}^0 + \text{MeHg}$). $\text{Hg}_P$ and $\text{MeHg}_P$ are $\text{Hg}^0$ and MeHg bound to particulate organic matter or silt, while $\text{Hg}_D$ and $\text{MeHg}_D$ are dissolved forms of $\text{Hg}^0$ and MeHg (ionic and dissolved organic carbon complexed). Transformations (dashed arrows) include photochemical ($\text{Hg}^{\text{II}}$ photoreduction, $\text{Hg}^0$ photooxidation, and MeHg photodemethylation) and biological (methylation and demethylation) processes. Transport processes include atmospheric deposition and evasion, input from rivers, exchange driven by advection and diffusion, and transport of $\text{Hg}_P$ and $\text{MeHg}_P$ (settling, deposition, and burial).
profile (5 m steps), were sampled using a titanium ultraclean conductivity-temperature-depth frame (De Baar et al., 2008) equipped with 24 × 24 L polyvinylidene fluoride samplers. Samples were filtered (0.2 μm, Sartoroban 300), drawn directly into individual precleaned 250 mL perfluoroalkoxy Teflon bottles (Savillex Purillex™), and acidified to 0.4% (v/v) with double-distilled HCl. We also sampled one station (40.833°N, 27.5667°E, Figure 1a) in the Marmara Sea (Mediterranean Sea) during the first leg of the 2013 GEOTRACES MEDBlack cruise (GA04-leg1) to obtain HgD and MeHgD data that we used to parametrize the inflow and outflow to the Black Sea.

Total HgD was measured in a 35 mL aliquot following the USEPA 1631 method, which was modified for ultralow-level seawater measurements (Heimbürger et al., 2015). Potassium bromide (Sigma-Aldrich, USA) and potassium bromate (Sigma-Aldrich, USA) were heated for 4 h at 250°C to remove Hg traces before preparing a BrCl solution with freshly double-distilled HCl. We used a custom-made semi-automatic single gold trap setup coupled to a cold vapor atomic fluorescence spectrometer (Brooks Rand Model III, USA), modified with a mirrored quartz cuvette (Hellma Optics, Germany).

We measured dissolved methylmercury (MeHgD) as the sum of monomethylmercury and dimethylmercury (MMHg + DMHg). MeHgD was analyzed via isotope dilution (ID) using a high-sensitivity coupled gas chromatograph-sector field ICP-MS (GC-SF-ICP-MS) (Heimbürger et al., 2015). Briefly, enriched spikes of [199]Hg and [201]MeHg (JSC Science, Spain) were added to a 115 mL aliquot of the seawater samples. After 24 h equilibration, the pH was adjusted to 3.9 with NH3 (ULTREX® II Ultrapure Reagent, J.T. Baker, USA) and a buffer solution of acetic acid (glacial, ULTREX® II Ultrapure Reagent, J.T. Baker, USA)/sodium acetate (J.T. Baker, USA). Sodium tetra propyl borate (1 mL, 1%, v:v; Merseburger Spezialchemikalien, Germany) was then added to 200 μL hexane (Sigma-Aldrich, USA). The glass bottles were hermetically sealed with Teflon-lined caps and vigorously shaken for 15 min. The organic phase was recovered, and 2 μL was injected into the GC (Thermo Trace Ultra) coupled to a SF-ICP-MS (Thermo Element XR). Initial results indicated undetectable amounts of MeHg in the anoxic waters. However, we also did not recover the added isotopic spikes, indicating that our method was not appropriate for anoxic samples. Another acidified aliquot was oxygenated for 15 min after the spike addition and equilibration but prior to the derivatization step. This led to full spike recovery and significant amounts of MeHg detected in all samples. The detection limit was 0.025 and 0.001 pM for HgD and MeHgD, respectively.

We took GEOTRACES intercalibration samples on 15 July 2013 in the western gyre (31.402°E; 42.521°N), in the oxic waters near the chlorophyll maximum (45 m depth) and sent these out to 25 participating laboratories. Our results compare well to consensus values: HgT = 0.92 ± 0.36 pM, n = 17, MeHgT = 0.57 ± 0.36 pM, n = 9. We measured the GEOTRACES intercalibration sample three times for HgT and obtained 0.98, 0.95, and 0.96 pM, on 10 September 2013, 15 December 2013, and 30 October 2014, respectively. We determined 0.091 and 0.063 pM of MeHg on 10 September 2013 and 15 December 2013, respectively.

### 2.2. Vertical Discretization of the Data Set

We analyzed the vertical distribution of physical properties, nutrients, and metals at 10 deep stations of the 2013 GEOTRACES MEDBlack cruise (Gerringa et al., 2016; Margolin et al., 2016), excluding station 12, which is in the Bosporus Strait, and station 11, which is shallow and influenced by the Mediterranean inflow. Many authors (e.g., Konovalov & Murray, 2001) have already observed that, due to the distribution of water masses along isopycnals, the profiles of several variables show a strong similarity when plotted against density but differ when depth is used (Figures 2, S1, and S2). As depth increases, the adiabatic compression on water molecules causes a temperature increase that leads to an apparent decrease in density ($\rho(S, T, p)$, kg m⁻³). To compare the density of two parcels of water from different depths, is recommended the use of potential density anomaly ($\sigma_0 = \sigma(S, 0, 0) - 1000$, kg m⁻³), which is computed from potential temperature (θ) defined as “the temperature that a water parcel would have if moved adiabatically to another pressure” (Talley et al., 2011). To exclude possible outliers, we first analyzed HgD and MeHgD profiles (up to 2,160 m depth; $\sigma_0 = 17.25$) by discretizing the water column using fine resolution (Table S1). This partitioning is appropriate to handle outliers and define the average concentration profile of metals and nutrients, but it is too detailed for our modeling purposes. A vertical discretization model (Table 1 and Figure 1b) was used in order to decompose the water column into its redox layers (OL, SOL, and AOL), further dividing some layers to properly represent the water balance of the system and simulate the evolution of the target variables (Hg, MeHg, and POM). Thus, the OL and AOL were divided into four sublayers each (namely, from O1 to O4, and from A1 to A4), and the sediments were divided into two sublayers (SED1 and SED2).
### Table 1

**Physicochemical Properties of Water and Sediment Layers and Sublayers**

| Redox layer | Model layer | Depth (m) | $\sigma_\theta$ (kg m$^{-3}$) | $E_z$ (m$^2$ s$^{-1}$) | $O_2$ (µM) | Hg (pM) | MeHg (pM) | MeHg/Hg (%) |
|-------------|-------------|-----------|-----------------|---------------------|-------|---------|-----------|-------------|
| OL (oxic layer) | O1 | 0–20 | 10.50–12.05 | $5.9 \times 10^{12}$ | $1.1 \times 10^{-5}$ | 246 | 1.86 | 0.12 | 8.5 |
| | O2 | 20–40 | 12.05–14.25 | $5.9 \times 10^{12}$ | $1.1 \times 10^{-5}$ | 315 | 2.13 | 0.14 | 8.6 |
| | O3 | 40–55 | 14.25–15.18 | $4.5 \times 10^{12}$ | $9.5 \times 10^{-6}$ | 165 | 1.77 | 0.29 | 19.9 |
| | O4 | 75–77 | 15.18–15.64 | $5.9 \times 10^{12}$ | $9.5 \times 10^{-6}$ | 313 | 1.90 | 0.31 | 15.7 |
| SOL (suboxic layer) | S | 75–100 | 15.64–16.24 | $7.4 \times 10^{12}$ | $9.5 \times 10^{-6}$ | 3.5 | 0.1 | 2.06 | 1.6 |
| AOL (anoxic layer) | A1 | 100–280 | 16.2–16.66 | $5.3 \times 10^{13}$ | $1.0 \times 10^{-5}$ | 18 | 3.16 | 0.80 | 28.6 |
| | A2 | 280–460 | 16.6–17.04 | $5.3 \times 10^{13}$ | $1.0 \times 10^{-5}$ | 97 | 2.81 | 0.55 | 21.8 |
| | A3 | 460–1460 | 17.04–17.24 | $2.9 \times 10^{14}$ | $5.5 \times 10^{-6}$ | 317 | 3.71 | 0.75 | 21.0 |
| | A4 | 1460–1810 | 17.24–17.25 | $1.0 \times 10^{14}$ | $8.7 \times 10^{-6}$ | 421 | 3.71 | 0.74 | 20.6 |
| Sediment | SED1 | 8.8 | $10^6$ | — | — | — | — | — |
| | SED2 | 2.0 | $10^{10}$ | — | — | — | — | — |

*Note: For each model layer are given: the depth and density range, the volume, the vertical eddy diffusivity ($E_z$) and the observed average concentrations of oxygen, sulfide, dissolved Hg and MeHg, and the MeHg%.

$^aE_z$: vertical eddy diffusivity from Konovalov et al. (2004).*

### 2.3. Model Structure

The WASP7 Hg model/MERC7 submodel (Wool et al., 2001) is a dynamic process-based model designed to simulate the Hg cycle within a system of well-mixed discrete spatial units (layers) of water and sediment as well as the exchanges of the system with its boundaries. This model has been already adapted and successfully applied to a shallow environment in a 2D configuration (Canu & Rosati, 2017; Melaku Canu et al., 2015), and is here used in a 1D configuration to explore Hg dynamics along an extended redox gradient.

Figure 3 shows the conceptual model of the processes that affect the evolution of the state variables (Table S2). Modeled Hg species (Table S3) include dissolved and particulate compounds of oxidized Hg ($Hg^+$), elemental mercury ($Hg^0$), and methylmercury (MeHg), whereas dimethylmercury (DMHg) is not explicitly considered. Dissolved species include $Hg^0$, dissolved organic carbon (DOC)-complexed species of $Hg^+$ and MeHg (i.e., $Hg_{DOC}$ and MeHg$_{DOC}$), and their ionic species ($HgCl^+$, $HgCl_2^+$, and MeHgCl) in marine environments (Morel et al., 1998). Particulate species involve only $Hg^+$ and MMHg, which in the model can be adsorbed to POM (Hg$_{POM}$ and MMHg$_{POM}$) or silt (Hg$_{silt}$ and MMHg$_{silt}$); silt represents inorganic particles, and POM is used to represent both organic particles and Fe/Mn oxides that are subjected to precipitation and dissolution. POM is modeled as a state variable that can be produced and decomposed according to the formulas given in Table S4, while DOC is modeled as constant in time, with different concentrations in each box according to Ducklow et al. (2007) and Margolin et al. (2016).

Hereafter, we refer to total Hg ($Hg_T$) and total MeHg (MeHg$_T$) as the sum of all Hg and MeHg species in the dissolved and particulate phases. We refer to dissolved Hg ($Hg_D$) to indicate the sum of all dissolved species ($Hg^0$, $Hg_{DOC}$, MeHg$_{DOC}$, $HgCl^+$, $HgCl_2^+$, and MeHgCl), while dissolved $Hg^+$ and MeHg (MeHg$_D$) will be used to indicate, respectively, the sum of $Hg_{DOC} – Hg_{N}$ and MeHg$_{DOC} – MeHg_{N}$. We will refer to particulate Hg ($Hg_P$) as the sum of inorganic Hg particle-bounded species (Hg$_{POM}$ and Hg$_{silt}$) and refer to particulate MeHg (MeHg$_P$) as the sum of MeHg$_{POM}$ and MeHg$_{silt}$.

The main transport (i.e., diffusion, advection, volatilization, deposition, settling, and burial) and transformation (i.e., biotic methylation and demethylation, photoreduction, photooxidation, and photodemethylation) processes related to the Hg cycle are considered in the model (Tables S3–S6). Resuspension does not occur in the deep water of the Black Sea due to low turbulence (Stanev & Kandilarov, 2012), and thus, it is not included in our model (Table S7). Transport processes (Tables S5) are described through advective water fluxes, diffusion coefficients, and transport mediated by particles. Hg transformation processes (Tables S6) are modeled using first-order kinetics, through a reaction constant ($k_x$) modulated based on environmental forcing such as temperature and light intensity.

### 2.4. Model Physical and Biological Setting

We represent the Black Sea as a 1-D vertical system with nine layers of water and two layers of sediment (Table 1). Since we aimed to simulate the dynamics in the offshore basin of the Black Sea, the shelf area
was excluded from analysis (~30% of surface area) (Panin & Jipa, 2002), constraining the model domain to represent 70% of the volume of the upper water and 100% of the deep water, overall accounting for 99% of the total volume. The steady state balance of water masses (Text S1 and Figure 1b) was arranged according to Ozsoy and Unlüıata (1997), and turbulent vertical diffusion ($E_z$; Table 1) was set according to Konovalov et al. (2004). Monthly averaged data for solar irradiation, day length, and sea surface temperature for the Black Sea in the year 2013 (Gencarelli et al., 2015; Grell et al., 2005) were used as forcing for photoreactions, biological reactions, and volatilization. To satisfy the sediment balance of the Black Sea (Text S2, Figure S3, and Tables S4 and S8), we considered silt and POM exchanges with the Marmara Sea (Mediterranean Sea) (Çoban-Yildız et al., 2000, 2006), loadings from rivers and the Azov Sea (Agapov et al., 2012; Berlinksy et al., 2006; Chasovnikov et al., 2012; Jaoshvili, 2002; Ovsienko et al., 2012; Panin & Jipa, 2002; Teschari et al., 2006) and internal POM production and degradation, including both primary production (Agirbas et al., 2014) and chemosynthesis (Nyffeler et al., 2007; Yılmaz et al., 2006). POM degradation rates were calibrated in agreement with the carbon budget (Deuser, 1971; Grégoire & Soetaert, 2010; Margolin et al., 2016), and 80% of suspended particles in the model is POM, in agreement with Stanev and Kandilarov (2012). Total suspended particles were calibrated to match concentrations (Figure S4) estimated from the beam attenuation coefficients (Karageorgis et al., 2008, 2009) measured during the cruise, which are in good agreement with those given in Stanev and Kandilarov (2012). For the sediment compartment, two layers were modeled: a 3 cm deep fluffy layer with porosity of ~0.96 (Yücel et al., 2012) on top of a 7 cm deep layer of more consolidated sediment.

2.5. Mercury Species Data and Time Dynamic Model Implementation

To simulate the historical evolution of Hg dynamics in the Black Sea in our 1-D model, we took into account the inputs and outputs of the system, transport mediated by water and sediment, and transformation processes (Tables S2–S8). We chose to reconstruct past inputs into the Black Sea because the residence time of the deep water has been estimated to be between 330 and 1,500 years (Murray et al., 2007), and thus, present-day concentrations in the deep water must reflect processes that have occurred over long periods. As anthropogenic Hg emissions increased dramatically with the onset of the industrial revolution (Amos et al., 2015), we estimated the temporal evolution of Hg inputs beginning in 1850, assuming that prior inputs are negligible for the Hg dynamics in the Black Sea. We gathered information about Hg species loads, boundary concentrations, and fluxes for the year 2013 (Tables S7–S9) and rescaled riverine and atmospheric Hg input according to historical anthropogenic enrichment factors estimated for global emissions and the deposition of Hg (Amos et al., 2015; Horowitz et al., 2014). The model was initialized at a pristine level, with 0.14 pmol of Hg\textsuperscript{0}, 0.02 pmol of Hg\textsuperscript{II} in surface waters, and 0.3 pmol of Hg\textsuperscript{II} in the waters below 100 m depth (Zhang Jaeglé, & Thompson, 2014); water MeHg was assumed to be 5% of Hg\textsuperscript{II}. Hg sediment concentration was set to 235 pmol g\textsuperscript{-1} (47 ng g\textsuperscript{-1}) based on the background Hg concentrations of sediment cores taken during the 2013 GEOTRACES MEDBlack cruise. The background Hg levels are 3.4 times lower than present-day surface level and do not show any influence of preindustrial Hg enrichments, supporting the idea that anthropogenic Hg inputs to the Black Sea prior 1850 are negligible. The simulation was projected to the year 2050, assuming that future Hg input will remain constant at 2013 levels.

Atmospheric concentrations of Hg\textsuperscript{II} and Hg\textsuperscript{0} and fluxes of Hg\textsuperscript{II} deposition (Table S7) over the Black Sea (year 2013) were computed with the Weather Research and Forecasting Model with Chemistry-Hg atmospheric model (Gencarelli et al., 2014, 2015, 2017). The atmospheric model was forced with the chemical initial and boundary conditions for Hg species taken from the ECHMERIT model (De Simone et al., 2014, 2016; Jung et al., 2009), and the model domain was adapted to include the entire Black Sea surface area; further information is given in Text S3 (Arctic Monitoring and Assessment Programme/United Nations Environmental Programme, 2013; Emmons et al., 2010; Gårdfeldt et al., 2003; Grell et al., 2005; Lin et al., 2006; Neu & Prather, 2012; Salzmann & Lawrence, 2006; Sandu & Sander, 2006; Wanninkhof, 1992; Wesely, 1989). Marine boundary conditions for Hg and MeHg were characterized based on measured concentrations in the Marmara Sea (Table S7) and hydrodynamics (Figure 1b and Text S1); Hg\textsuperscript{0} concentrations were assumed to be 10% of Hg\textsuperscript{II} (Fantozzi et al., 2013; Gårdfeldt et al., 2003; Kotnik et al., 2015; Soerensen et al., 2013). Photoreaction rates (Table S9) are internally adjusted for the light conditions following Soerensen, Schartup, et al. (2016). These processes are relatively well constrained; others are much less: (1) the partitioning of Hg, which is parameterized through the partition coefficient ($k_D$, l/kg) for which a wide range is reported in the literature (Han, Lehman, et al., 2007; Han, Obraztsova, et al., 2007; Hollweg et al., 2010; Lamborg et al., 2016; Liu et al. (2016)).
et al., 2015; Muresan et al., 2007; Schartup et al., 2014, Schartup, Balcom, et al., 2015); (2) the Hg river inputs (Tables S7 and S8) (Amos et al., 2014; Soerensen, Schartup, et al., 2016; Woitke et al., 2003; Zhang et al., 2015), with the fraction of river MeHg being generally low (~5% of HgT) (Balogh et al., 2003; Mason et al., 2012; Paller et al., 2004; Sharif et al., 2014; Soerensen, Jacob, et al., 2016, Soerensen, Schartup, et al., 2016); and (3) Hg methylation and MeHg demethylation (Hsu-Kim et al., 2013; Semeniuk & Dastoor, 2017).

A separated model setup was implemented to simulate the preanthropogenic cycle of Hg in the Black Sea by running two additional simulations forced only with natural Hg inputs (upper and lower estimates). Natural river inputs were assumed to be 0.4%–16% of present-day inputs (corresponding to HgP = 0.01–0.1 nmol g⁻¹ that is in the lower range of the assessment by Amos et al. (2014)) and natural Hg atmospheric deposition to be 13–20% of present-day deposition (Zhang, Jaeglé, Thompson, & Streets, 2014). Results of these simulations were used to estimate the anthropogenic Hg in the Black Sea relative to the base simulation.

2.6. Model Calibration and Analysis of Uncertainties in Parameters and Model Output

Monte Carlo analysis (i.e., analysis of an ensemble of numerical simulations) can be useful to estimate uncertain model parameters while maintaining a reasonable fit to observations (Beck, 1987; Cossarini & Solidoro, 2008; Jørgensen, 1994; Pianosi et al., 2016; Soetaert & Herman, 2009). Here we constrained the most uncertain processes/parameters to observed concentration of HgD and MeHgD in the water column and HgT in sediment, by using a two steps procedure involving an ensemble of simulations. In the first step, we derived the most likely estimates of river loads and partition coefficient (kD) against water HgD and sediment HgT observations. In the second step, the model was refined and calibrated to observational data of water MeHgD and HgD, by tuning the Hg methylation and demethylation rates (km and kdm). The agreement between modeled (mod) and observed (obs) concentrations in waters and sediment for the reference year (2013) was evaluated for each sensitivity simulation by computing the Pearson’s correlation coefficients (r) and the root-mean-square errors (RMSE, equation (1)).

\[
\text{RMSE} = \sqrt{\frac{\sum_{i=1}^{n} (\text{obs}_i - \text{mod}_i)^2}{n}}
\] (1)

As a first step, we run an ensemble of simulations excluding biotic production and degradation of MeHg within the system. Each simulation was performed by using the same values for all parameters but for kD and the time-variable river load (Lr(t)), which were varied within a large interval, namely, kD = 5 × 10⁵–6.2 × 10⁶ l/kg, and Lr(t) = 15–40 kmol/yr (Lr(2003) were derived from Lr(2003) by backward scaling, according to Horowitz et al. (2014), to simulate the historical evolution). Then we compared each simulation against observations and rejected all the simulations that have \( r < 0.9 \), RMSE for water HgD > 0.8 pM, and RMSE for sediment HgT > 300 pmol g⁻¹. The subset of remaining simulations (the ones with a good fit to observations) was retained, and the corresponding parameters values were used to estimate the distribution (given as mean ± standard error (SE)) of: Lr(t) = 25 ± 2.5 kmol yr⁻¹ and kD = 5.6 × 10⁵ ± 9.3 × 10⁴ l/kg. The simulation from this subset that reproduces the average water HgD and sediment HgT concentrations is hereafter referred to as reference simulation. The other simulations with a good fit to observations were used to compute the SE of modeled concentrations, which is related to the uncertainties in kD and Lr(t).

In the second step, we performed a second ensemble of simulations, starting from the reference simulation obtained in step 1 and adding in Hg methylation and MeHg demethylation rates (km and kdm) in the different water layers. In this case we identified the variability interval by selecting km and kdm values for different redox conditions within literature ranges (Table S10) (Hollweg et al., 2009; Lehn herr et al., 2011; Monperrus, Tessier, Amouroux, et al., 2007; Rodríguez Martín-Doimeadios et al., 2004; Whalin et al., 2007) and by varying the selected rates by ±50% (Table 2). To gain insights into the significance of these processes in the redox layers of the Black Sea (OL, SOL and AOL), we set up three different scenarios (Table 2) based on the following hypotheses:

1. ubiquitous Hg methylation and MeHg demethylation;
2. Hg methylation is limited to the SOL, ubiquitous MeHg demethylation;
3. Hg methylation and MeHg demethylation are both limited to the SOL.

As before, we rejected the simulations that did not have a sufficient fit with the data and used the subset of retained simulation (those having \( r \) for water MeHgD and HgD > 0.9 as well as RMSE for water
Table 2
Parameterization of Hg Methylation (km) and MeHg Demethylation (kdm) in the Simulations: a) Ubiquitous km and kdm; b) km Limited to the SOL and Ubiquitous kdm; c) km and kdm Limited to the SOL

| Layer | km (d⁻¹)          | kdm (d⁻¹)          | km (d⁻¹)          | kdm (d⁻¹)          | km (d⁻¹)          | kdm (d⁻¹)          |
|-------|-------------------|--------------------|-------------------|--------------------|-------------------|--------------------|
| OL    | 4 x 10⁻³ [±50%]   | 1.0 x 10⁻¹ [±50%]  | 0                 | 1.7 x 10⁻² [±50%]  | 0                 | 0                  |
| SOL   | 1.6 x 10⁻² [±50%] | 2.1 x 10⁻¹ [±50%]  | 1.6 x 10⁻² [±50%] | 2.1 x 10⁻¹ [±50%]  | 1.6 x 10⁻² [±50%] | 2.1 x 10⁻¹ [±50%]  |
| AOL   | 3.3 x 10⁻³ [±50%] | 1.5 x 10⁻² [±50%]  | 0                 | 1.5 x 10⁻² [±50%]  | 0                 | 0                  |

Note: See Table S10 for literature ranges of km (d) and kdm (d) under different redox conditions. OL = oxic layer; SOL = suboxic layer; AOL = anoxic layer.

MeHg/D < 0.2 pM and RMSE for water Hg/D < 0.35 pM). Obviously, to take into consideration the uncertainties in km and kdm, values produces some variability in model output. However, from the analysis of the second ensemble of simulations it turned out that such variability is significantly smaller than the one related to the parameters variation in the first calibration step (not shown) and that the tuning of km and kdm induced minor variations in the fit against water Hg/D and sediment Hg/T. Finally, we selected the model ensemble giving the best fit to data and use that to define the fluxes and related uncertainties in a budget of Hg species in the Black Sea.

3. Results and Discussion

3.1. Observations

Hg/D and MeHg/D vertical profiles show similar behavior across the 10 deep stations sampled in the Black Sea (Figures 2, S1, and S2), suggesting that all stations can be described in terms of a similar dynamic along the vertical dimension. Thus, a 1-D representation of the system seems suitable to capture the most important dynamics of Hg in the Black Sea. Concentrations in the OL and SOL are lowest (Hg/D = 1.9 ± 0.6 pM; MeHg/D = 0.13 ± 0.05 pM) and do not differ from each other (Wilcoxon signed-rank test; p value <0.05), while concentrations in the AOL (Hg/D = 3.4 ± 0.6 pM; MeHg/D = 0.71 ± 0.17 pM) are significantly higher (Wilcoxon signed-rank test; p value <0.05). Both Hg/D and MeHg/D concentrations exhibit two maxima in the AOL: the first one is just below the SOL-AOL interface and the second one in the deep part of the AOL. While the Hg/D and MeHg/D profiles mimic each other, the relative MeHg increase from SOL to AOL (+340%) is more important than the Hg/D increase (+62%), suggesting that production of MeHg occurs in the AOL. Figure 2 shows the vertical distribution of Hg/D, MeHg/D, and other biogeochemical variables against σθ (Figure 5). We found that the Hg/D and MeHg/D maxima are broadly coincident with the Fe/D and/or Mn/D (Figure 5) in the Black Sea water column, as previously observed for Hg/D and Mn/D/Fe/D (Lamborg et al., 2008) as well as for Hg/T and Fe/D/Mn/D (Cossa & Coquery, 2005). The association between Hg species and Mn/Fe redox cycles is usually explained by the “shuttle theory” that postulates a transportation process based on Mn/Fe oxides (Neretin et al., 2003). According to this theory, Hg is scavenged from surface layers by POM and Mn/Fe oxides and settles toward the bottom of the SOL, where oxide reduction drives NOM remineralization, causing Hg/D release. Reduced Mn/Fe diffuses upward and is oxidized at the top of the SOL; likewise, Hg/D released in the ALO diffuses upward and is scavenged by oxides. Therefore, Hg species are trapped in this cycle between the SOL and ALO. This process has been invoked to explain the distribution of phosphates in the Black Sea (Dellwig et al., 2010; Pahmova & Yakushev, 2013; Shaffer, 1986) and the distribution of Hg species in stratified water bodies with anoxic bottom water (Cossa & Coquery, 2005; Han, Lehman, et al., 2007; Mason et al., 1993). The partition coefficients (KD) of Hg to oxides have been shown to be higher than the range generally observed in sediments; Lamborg et al. (2016) estimated a K_D,Mn = 10⁸⁻⁴, and K_D,Fe = 10⁷⁻⁶ for Hg/T in the ocean. In the Black Sea, Mn is much more abundant than Fe (Figures 2 and 5) and likely...
plays a major role. As shown in Figure 5, concentrations of dissolved Mn (MnD) in the Black Sea rise in the SOL, peak below the SOL-AOL interface ($\sigma_\theta \sim 16.3$) and remain elevated and constant until the seabed. Fe oxide reduction follows Mn oxide reduction in the sequence of electron acceptors used by bacteria for NOM remineralization (Froelich et al., 1979); indeed, the increase in dissolved Fe concentrations (FeD) is deeper than that of MnD, and FeD concentrations are maximal at $\sigma_\theta \sim 16.5$. Below $\sigma_\theta \sim 16.5$, FeD decreases due to precipitation as FeS$_{(s)}$ or FeS$_2$$_{(s)}$; at this depth, there is a local minimum of HgD and MeHgD concentrations (Figures 2, S1 and S2), which might be due to their partial pyritization (coprecipitation) in the formation of FeS$_2$ (Huerta-Diaz & Morse, 1992; Morse & Luther, 1999). This process has also been proposed also for the

Figure 4. Distribution of MeHgD (pM) along 2013 GEOTRACES MEDBlack cruise transect. The numbers of the sampling stations (Figure 1) are given on the top of the figure. Black dots indicates sampling points. The contours of the oxic layer (OL), suboxic layer (SOL), and anoxic layer (AOL) are indicated by white lines.

Figure 5. High-resolution profiles of dissolved Hg (HgD), and MeHg (MeHgD), oxygen (O$_2$), hydrogen sulfide (HS$^-$), nitrates (NO$_3^-$), phosphates (PO$_4^{3-}$), manganese (MnD), and iron (FeD) at station 5 (Figure 1). OL = oxic layer; SOL = suboxic layer; AOL = anoxic layer.
cycling of Hg species in the Pettaquamscutt Estuary (Mason et al., 1993), while in Offatts Bayou (USA), it has been suggested to affect the water profile of Hg\(^{2+}\) but not that of MeHg; this is likely because higher POM concentrations in the shallow bay outcompete Fe\(\text{S}_2\) for binding sites (Han, Lehman, et al., 2007).

Our high-resolution profiles (Figure 5), with 5 m sampling intervals covering the extended redox gradient from oxic waters (40 m depth) to anoxic waters (160 m depth) at station 5, allowed us to further resolve Hg distributions. The high-resolution Hg\(_{\text{D}}\) maximum (3.9 pM at 130 m depth) is shallower at the interface of the SOL and AOL, just below the PO\(_4^{3-}\) peak. The high-resolution MeHg\(_{\text{D}}\) maximum (0.83 pM at 145 m depth) is 15 m deeper, coinciding with the Mn\(_2\) peak, well within the AOL. Mercury methylation may be highest here because of the continuous supply of Hg\(_{\text{D}}\) during the dissolution of Mn oxides. The MeHg/Hg percentage increases further with depth in the AOL, reaching 39% at a 160 m depth. It is remarkable that the MeHg/Hg percentages are comparable to other open-ocean subsurface maxima in oxic waters (Bowman et al., 2015, 2016; Cossa et al., 2009, 2011; Ganachaud et al., 2017; Hammerschmidt & Bowman, 2012; Heimbürger et al., 2015; Kim et al., 2017; Munson et al., 2015; Sunderland et al., 2009; Wang et al., 2012).

In the deeper part of the AOL (\(\sigma_\theta > 17.04\)), Hg\(_{\text{D}}\) and MeHg\(_{\text{D}}\) concentrations increase to a second maximum (Figures 2, S1 and S2), which might be due to the enhanced solubility of Hg species in the presence of sulfides and DOM (Han, Lehman, et al., 2007; Mason et al., 1993; Ravichandran et al., 1998, 1999; Skyllberg, 2008; Slowey, 2010; Sunderland et al., 2006; Waples et al., 2005) and to net in situ methylation, which is favored by DOM (Graham et al., 2012; Moreau et al., 2015; Schaefer & Morel, 2009; Schaefer et al., 2011). It has been shown that in sulfidic systems, some DOM components (thiols, humic acids, and cysteine) delay or inhibit Hg\(_{\text{S}_\text{II}}\) precipitation and favor its dissolution, leading to the formation of nanoparticles smaller than 0.2 \(\mu\)m (Deonarine & Hsu-Kim, 2009; Gerbig et al., 2011; Slowey, 2010): these nanoparticles can be methylated (Jonsson et al., 2012; Kucharzyk et al., 2015; Zhang, Kucharzyk, et al., 2014; Zhang et al., 2012). These processes appear to be favored by low Hg/DOM ratios and may provide a mechanistic explanation for the second maximum of MeHg\(_{\text{D}}\) that we observed in the AOL, where the ratio Hg/DOM is as low as ~0.2 nmol Hg/mg DOM. Recent research has also shown that MeHg adsorbed to reduced sulfur groups on minerals (Fe\(\text{S}_\text{m}\), Hg\(_{\text{S}_\text{II}}\) and Cd\(_{\text{S}_\text{II}}\)) and organic (thiols) surfaces can be methylated to dimethylmercury (DMHg) (Jonsson et al., 2016). We measured DMHg at stations 2 and 5. The results (not shown) were near the detection limit, and we therefore conclude that the DMHg is only a minor fraction of the MeHg pool in the Black Sea, in agreement with Lamborg et al. (2008), who found DMHg to be less than 10% of MeHg. Therefore, we only considered the dominant MMHg species in the numerical model (labeled as MeHg).

While our observations of two deep maxima of Hg\(_{\text{D}}\) are consistent with previous findings for both filtered and unfiltered Hg in the Black Sea (Cossa & Coquery, 2005; Lamborg et al., 2008), the distribution of MeHg\(_{\text{D}}\) differs from that observed in the western gyre by Lamborg et al. (2008), who detected only a shallow MeHg\(_{\text{D}}\) maximum at \(\sigma_\theta \sim 16.2\) and decreasing concentrations toward the seabed. We believe that this discrepancy is due to different analytical methodologies. We applied an ID method that can assess recovery of the analytes (section 2.1), while Lamborg et al. (2008) did not use ID. Their method might not have been efficient for the analysis of anoxic Black Sea waters. On the other hand, our observations are consistent with the distribution of MeHg along the stratified water column of the Gotland Sea (Baltic Sea) (Soerensen, Schartup, et al., 2016) and the seasonally anoxic waters of Offatts Bayou (USA) (Han, Lehman, et al., 2007), where concentrations are minimal above the halocline and maximal in anoxic water. Sulfide levels in these two systems are lower but comparable (150–200 \(\mu\)M) to those of the Black Sea (Han, Lehman, et al., 2007; Neretin et al., 2003).

3.2. Modeled Concentrations and Fluxes

The three ensembles of simulations referring to scenarios a, b, and c were all derived adding Hg methylation and MeHg demethylation in the same reference simulation, which was obtained from the first step of calibration against concentrations of Hg\(_{\text{D}}\) in water (Figure 6a) and Hg\(_{\text{S}_\text{II}}\) in the sediment (Figure 6b). The three scenarios a, b, and c represent different hypotheses for Hg methylation and MeHg demethylation occurrence along the redox gradient (Table 2), which were explored during the second step of calibration by varying \(k_m\) and \(k_{\text{dm}}\) values.

Results show that the ensemble of simulations (a), representing ubiquitous Hg methylation and MeHg demethylation (Figure 7a), reproduces best the MeHg\(_{\text{D}}\) profiles observed during the cruise (\(r = 0.98 \pm 0.01\); RMSE = 0.15 ± 0.07 pM). When Hg methylation is considered to be limited to the SOL (Figures 7b and 7c),
Figure 6. (a) HgD profiles in the water column observed during the cruise (black circles and bars: average and standard deviation of the 10 deep stations) and modeled (gray dashed lines: profiles from all simulations; yellow area: range estimated from the subset of simulations retained from the Monte Carlo ensemble (see section 2.6); blue triangles: reference simulation) by varying the river load and kD. The potential density anomaly ($\sigma_\theta = \sigma(S, \theta, 0) - 1000, \text{kg m}^{-3}$) is used as the vertical coordinate, and the corresponding depths are shown on the left y axes; (b) HgD concentrations in surface sediment (0–3 cm) observed during the cruise (gray box plot) and modeled (yellow box plot); each box plot indicates the median (thick bar), the interquartile range (box height), and the extremes of the distribution (whiskers). OL = oxic layer; SOL = suboxic layer; AOL = anoxic layer.

Figure 7. Distribution of MeHgD in the water column observed during the cruise (black dots, black circles and bars: data points, means, and standard deviation of the 10 deep stations) and range of modeled concentration (shaded areas) profiles from different simulations (gray dashed lines) run assuming: (a) ubiquitous $k_m$ and $k_{dm}$ (green area), (b) $k_m$ limited to the suboxic layer (SOL) and ubiquitous $k_{dm}$ (blue area), (c) $k_m$ and $k_{dm}$ limited to the suboxic layer (pink area), (d) all of the previous (Table 2). The potential density anomaly ($\sigma_\theta = \sigma(S, \theta, 0) - 1000, \text{kg m}^{-3}$) is used as the vertical coordinate, and the corresponding depths are shown on the left y axes. OL = oxic layer; SOL = suboxic layer; AOL = anoxic layer.
the model cannot be tuned to reproduce the observed vertical MeHg distribution. The ensemble of simulations (b) have \( r = 0.5 \pm 0.08 \) and \( \text{RMSE} = 0.47 \pm 0.01 \) pM, while simulations (c) have a better fit \( (r = 0.93 \) and \( \text{RMSE} = 0.34 \pm 0.06 \) pM) but still significantly worse than (a). We conclude that observations and modeling falsify the hypotheses (b) and (c) and corroborate the hypothesis (a), that is, ubiquitous methylation and demethylation. Therefore, the setup (a) defines the most likely model given the available data.

We used the best model setup (scenario a), to compute concentrations and fluxes of Hg species (values are given as mean ± SE) in water and sediment of the Black Sea for the 2013 (Figure 8) and for the whole simulation period (1850–2050; Figure 9).

In its present state, Hg\(_T\) inputs to the water column \( (31 \pm 2.5 \text{ kmol yr}^{-1}) \) are estimated to exceed the outputs \( (28 \pm 2.8 \text{ kmol yr}^{-1}) \). The system is not at steady state, the \( \Delta \text{Hg}_{\text{T}}(2014-2013) \) (difference of Hg pool between 2014 and 2013) is negative in the OL \( (-0.3 \text{ kmol yr}^{-1}) \) and positive in the SOL and AOL \( (+0.04 \text{ and } +0.52 \text{ kmol yr}^{-1}) \). The most relevant input is riverine load \( (77\%, 24.5 \pm 2.5 \text{ kmol yr}^{-1}) \), followed by atmospheric deposition \( (17\%, 5.3 \text{ kmol yr}^{-1}) \). Little Hg\(_T\) comes from sediment via pore water diffusion \( (4\%, 1.4 \pm 0.2 \text{ kmol yr}^{-1}) \) from the Marmara Sea inflow \( (3\%, \sim 0.9 \text{ kmol yr}^{-1}) \) and from the Azov Sea \( (1\%, \sim 0.4 \text{ kmol yr}^{-1}, \text{not shown}) \). The main loss of Hg from the water \( (74\%) \) is the settling and deposition of \( 20.9 \pm 1.2 \text{ kmol yr}^{-1} \) of Hg\(_P\) and MeHg\(_P\). Evasion of Hg\(_0\) \( (23\%, 5.4 \pm 0.5 \text{ kmol yr}^{-1}) \) roughly compensates Hg\(_{\text{II}}\) deposition, and the outflow to the Marmara Sea accounts for \( 6\% \) \( (1.4 \pm 0.1 \text{ kmol yr}^{-1}) \). The 26% of Hg that deposits to the sediment is buried below a 10 cm depth \( (4.1 \pm 0.3 \text{ kmol yr}^{-1}) \), while 8% moves back to the water through pore water diffusion; the remainder constitutes the sediment reservoir. The modeled reservoir of
HgD in the water column is 1675 ± 320 kmol (~320 ± 64 t), 23% of which (358 ± 49 kmol; ~77 ± 10 t) is present as MeHgD; the mismatch between modeled and measured concentrations for the mean modeled concentrations amounts to ~143 kmol for HgD and ~6.8 kmol for MeHgD.

Our budget of HgT in the Black Sea (Figures 8 and 9a), built on biogeochemical modeling and measurements along 10 stations, is here compared to the budget of HgT estimated by Lamborg et al. (2008) based on water column measurements at two sites in the Black Sea. The highest mismatch between the two budgets is in the estimate of Mediterranean inflow-outflow and Hg0 evasion. Our estimate of inflow (~0.9 kmol yr⁻¹ of HgT) and outflow (1.5 ± 0.1 kmol yr⁻¹), based on water column measurements on both sides, implies a small net export of Hg species (~0.6 kmol yr⁻¹ of HgT) to the Mediterranean Sea, while in the budget of Lamborg et al. (2008) a net input of 5 kmol yr⁻¹ is estimated, assuming high concentrations in the Marmara inflow (24 pM) that are not supported by observations. Our estimate of Hg⁰ evasion from the water (5.4 ± 0.5 kmol yr⁻¹) is much lower than the 93 kmol yr⁻¹ estimated by Lamborg et al. (2008). The concentrations of Hg⁰ measured by Lamborg et al. (2008) in surface water (0.3 pM) are slightly higher than the 0.2 pM modeled in our simulation. The evasion rate estimated by Lamborg et al. (2008) and computed by our model (respectively, ~600 pmol m⁻² d⁻¹ and ~50 pmol m⁻² d⁻¹) lies in the upper and lower range of the rates observed in coastal, shelf, and ocean stations (Andersson et al., 2007; Gärdfeldt et al., 2003; Soerensen et al., 2013). The atmospheric Hg deposition to the Black Sea (5.3 kmol yr⁻¹ over the surface area of our model) modeled through the Weather Research and Forecasting Model with Chemistry model is close to the 8 kmol yr⁻¹ estimated by Lamborg et al. (2008). The riverine load (24.5 ± 2.5 kmol yr⁻¹) is higher than the 8 kmol yr⁻¹ estimated by Lamborg et al. (2008), who acknowledged that the flux was likely higher. The scarcity of observations from the Black Sea tributaries calls for a more thorough investigation. HgD diffusion from pore water has a small overall impact on our budget.
(1.4 ± 0.1 kmol yr⁻¹) and is lower than, but consistent with, the 4 kmol yr⁻¹ predicted by Lamborg et al. (2008), who also included the contribution of shelf sediments.

Our model also simulates the time evolution of MeHg fluxes and concentrations (Figures 9b and 9d); thus, here we provide the first attempt—to the best of our knowledge—to quantify the cycle of this important compound in the Black Sea. For the 2013 MeHg budget (Figure 8), the estimated inputs to the water (5.2 ± 0.6 kmol yr⁻¹) are larger than the outputs (4.5 ± 0.5 kmol yr⁻¹), and the bulk of MeHg (3.8 ± 1.3 kmol yr⁻¹) comes from in situ methylation in the AOL. Net MeHg demethylation occurs in the OL (1.1 kmol yr⁻¹) and SOL (6.2 kmol yr⁻¹) and net Hg methylation in the AOL (11.1 kmol yr⁻¹). In the OL, 0.1–0.3% of the Hg³⁺ pool is methylated and 3–8% of the MeHg pool is demethylated on a daily basis; MeHg has a turnover time of 11–33 d⁻¹. In the SOL, both processes are most efficient: every day 0.7–1.4% of the Hg³⁺ pool is demethylated and 9–27% of the MeHg pool is methylated; MeHg has a turnover time of 4–11 d⁻¹. In the AOL, Hg methylation involve 0.1–0.3% of the Hg³⁺ pool, and MeHg demethylation 0.4–1% of the MeHg pool; the turnover time of MeHg is 100–240 d⁻¹.

Rivers account for 1.1 kmol yr⁻¹ of MeHg (20% of total inputs), while minimal contributions come from the Marmara and Azov Seas, which together account for 1.6% (0.06 and 0.04 kmol yr⁻¹). The diffusion of MeHgD₂ is predicted to be a sink for the water rather than a source, transferring 0.1 ± 0.01 kmol yr⁻¹ to the sediment (3% of outputs). Approximately 4.4 ± 0.6 kmol yr⁻¹ of MeHg leaves the water column after settling and deposition, while 0.04 ± 0.01 kmol yr⁻¹ moves to the Marmara Sea, transported by the Mediterranean outflow. According to our budget, methylation in the AOL is the most relevant source of MeHg to the basin. Thus, this process needs to be further investigated in the field. Based on the Fe peak in the upper AOL, methylation may be driven by Fe-reducing bacteria, which have been reported to prompt Hg desorption and methylation by using Fe oxides as electron acceptors (Hellal et al., 2015; Kerin et al., 2006; Schaefer et al., 2011; Si et al., 2015), whereas sulfate-reducing bacteria (Benoit et al., 2001; King et al., 2000; Moreau et al., 2015; Schaefer et al., 2014) are likely to contribute significantly to Hg methylation in the lower part of the AOL, where HS⁻ > 100 μM and increases with depth. However, they could also play a role in Hg methylation within aggregates settling from the SOL to the AOL (section 3.1). Large aggregates of precipitates, organic, and inorganic matter that form in the SOL and sink also operate also as carriers of bacteria (Fuchsmann et al., 2011). While Hg methylation and MeHg demethylation are thought to involve mainly dissolved Hg species, they have been detected in particles and marine snow aggregates and at particle surfaces under laboratory conditions (Jonsson et al., 2016; Ortiz et al., 2015). It has been proposed that in the Black Sea, all S cycling may occur within marine snow aggregates formed in the SOL that settle toward the AOL as well as at the surface of FeS and HgS minerals that precipitate in the AOL. Based on the recent identification of new bacterial strains able to methylate Hg (Gilmour et al., 2013), there are other potential candidates for Hg methylation in the AOL of the Black Sea. These are methanogens and the species Desulfitobacterium metallireducens (genus Firmicutes), which can use either Mn or Fe oxides as electron acceptors (Villemur et al., 2006). Both methanogens and Mn-reducing bacteria are abundant in the water of the lower SOL and in the upper part of the AOL (Fuchsmann et al., 2011; Pimenov & Neretin, 2006).

The idea that Hg methylation occurs in the eutinic waters of the Black Sea is in contrast with previous studies, which suggested that methylation is hindered at HS⁻ > 10 μM (Benoit et al., 2001; Benoit, Gilmour, et al., 1999; Benoit, Mason, & Gilmour, 1999; Lamborg et al., 2008). Such inhibition would be due to a shift in inorganic Hg speciation from a neutral complex (HgS⁰) to charged species (e.g., HgHS⁻) that cannot diffuse through microbial cell membranes. This implies that bacterial Hg uptake depends on passive diffusion through the cell, while it has been shown that it can be an energy-dependent transport mediated by carrier proteins and favored by some thiols such as cysteine (Schaefer et al., 2011, 2014; Schaefer & Morel, 2009). Many authors have noted that binding and solubility constants used to model the interactions among Hg/sulfide and Hg/DOM are highly uncertain (Drott et al., 2007; Hsu-Kim et al., 2013; Merritt & Amirbahman, 2009; Skyllberg, 2008). Furthermore, as discussed in section 3.1, Hg speciation in the presence of sulfides is affected by DOM composition, which can delay or inhibit the precipitation of HgS₃ and enhance its dissolution. HgS nanoparticles form under these conditions, which are very reactive particles at the surfaces of which Hg methylation occurs (Deonarine & Hsu-Kim, 2009; Gerbig et al., 2011; Hsu-Kim et al., 2013; Jonsson et al., 2012; Kucharzyk et al., 2015; Ravichandran et al., 1998, 1999; Skyllberg, 2008; Slowey, 2010; Waples et al., 2005; Zhang, Kucharzyk, et al., 2014; Zhang
et al., 2012). Beyond the methylation of HgS nanoparticles observed in laboratory experiments, there is evidence that adds weight to the argument for methylation under sulfidic conditions. Coquery et al. (2003) observed maximum levels of MeHg\(\frac{M}{M}\) in the anoxic water of a tropical reservoir and attributed these to in situ methylation by sulfate-reducing bacteria. Eckley and Hintelmann (2006) detected methylation only under hypoxic-anoxic conditions in a group of seasonally stratified lakes, with maximal rates just below the oxycline that remained elevated to the bottom of the lake (HS\(^{-}\) up to 30 \(\mu\)M). Hollweg et al. (2009) detected Hg methylation in the sediments of the Chesapeake Bay at sulfide levels as high as 100–1,000 \(\mu\)M, though rates were maximal at HS\(^{-}\) – 1 \(\mu\)M. Merritt and Amirtha-Raman (2008) reported a MeHg maximum with HS\(^{-}\) ranging from 0.5 to 30 \(\mu\)M in the pore water of the Penobscot River Estuary. In the sediment of Passamaquoddy Bay, Sunderland et al. (2006) observed the highest MeHg/Hg ratio in the zone with sulfide levels between 1,300 and 6,000 \(\mu\)M. Moreover, Hg methylation at millimolar levels of sulfides has been detected in sediment samples and in pure cultures of bacteria (Drott et al., 2007; King et al., 2000; Langer et al., 2001).

Some authors have pointed out that the decrease in Hg methylation with depth, frequently observed in pore water and sediment profiles, might be attributable to the limitation of available NOM substrates (electron donors) rather than to increasing sulfide levels (Drott et al., 2007; Merritt & Amirtha-Raman, 2009). This could explain why profiles of MeHg in the Black Sea water column differ from profiles often observed in pore waters and sediments that exhibit surface or subsurface maxima. In the upper part of the AOL, microbial activity is high due to the NOM produced at the lower boundary of the SOL by chemosynthetic bacteria (Pimenov & Neretin, 2006; Yilmaz et al., 2006).

The temporal evolution of Hg\(_T\) concentrations (Figure 9c) in the OL and SOL shows a fast response to changes in atmospheric and riverine loadings and follows the seasonal signal of atmospheric Hg deposition. In the AOL, Hg\(_T\) concentrations increase more slowly due to the longer times needed for Hg transport to deep water but are almost linear with time, as the net Hg\(_T\) flux from the SOL to AOL is larger than the flux from the AOL to sediment (Figure 8). Similarly to Hg\(_T\), MeHg\(_T\) tends to increase with time in the AOL (Figure 9d), where it exhibits maximal concentrations. However, the MeHg\(_T\) distribution in the water column differs from Hg\(_T\), and throughout the simulation, concentrations are lower in the OL and SOL and highest in the AOL. This is because MeHg\(\frac{M}{M}\) is much less affected by external loadings and is rather controlled by in situ production and degradation in the water column. The analysis of time evolution highlights that, in spite of a reduction in anthropogenic Hg input since the 1970s (Figures 9a and 9b), the model predicts the retention and accumulation of Hg species in the AOL, likely due to the “Mn shuttle” transportation process described in section 3.1. Briefly, Mn/Fe oxides in the SOL act as a barrier that limits upward diffusion and favors downward transport of Hg species associated with settling oxides, which are reduced and dissolved in the AOL releasing Hg species. In our model, oxide precipitation and dissolution are represented in a simplified fashion by parametrizing the production of POM within the SOL and its degradation in the first AOL sublayer (Text S2 and Figure S3).

The modeled pool of Hg from natural sources in the Black Sea is between 122 and 268 kmol. By difference with our base simulation, anthropogenic Hg is 1,517–1,663 kmol and accounts for 85%–93% of the water Hg pool.

### 4. Conclusions

During the 2013 GOETRACES MEDBlack cruise, we measured Hg and MeHg concentrations along the stretched redox gradient of the Black Sea water column. The new measurements were integrated into a numerical model to better understand MeHg production, Hg species cycling in the Black Sea, and their temporal evolution. A comparison between observational data and ensembles of simulations was performed to calibrate the model estimating the most likely value for the uncertain model parameters and their errors, as well as the uncertainty in model output due to parameters uncertainty. Such a comparison also allowed us to reject the hypothesis that Hg methylation is limited to the suboxic water of the Black Sea, and rather, it supports the idea that both Hg methylation and MeHg demethylation occur ubiquitously.

Contrary to a previous assessment (Lamborg et al., 2008), we found the highest MeHg concentrations in the permanently anoxic waters of the Black Sea. These new observations, backed with numerical modeling and mass balance calculations, indicate that Hg methylation in anoxic water is likely to be the primary MeHg source to the basin. Our findings offer additional theoretical and empirical support to increasing evidence that Hg can be methylated under sulfidic conditions, when the NOM pool of the system is bioavailable to
bacteria and contains functional groups that may inhibit the precipitation of HgS. MeHg percentages are highest in anoxic waters (up to 57%) and comparable to other open-ocean subsurface maxima inoxic waters.

According to our budget, inputs of Hg species to the Black Sea water column exceed outputs. Inflows and outflows to the Mediterranean Sea, as well as flux from sediment, are slight. The main source of HgII is river load followed by atmospheric deposition, while MeHg mostly comes from in situ methylation in anoxic waters. Most of the Hg in the Black Sea water column (85–93%) is of anthropogenic origin. During the time span of the simulation (1850–2050), modeled concentrations of Hg and MeHg increase in the deepest layers of the Black Sea (AOL), owing to recycling between suboxic and anoxic water that limits upward diffusion.

Overall, concentrations of Hg species in the Black Sea water column appear to depend on the balance between (1) the amount of HgII (mostly Hg0) that enters surface layers, is scavenged, and is transported to the SOL; (2) the recycling of Hg and MeHg between the SOL and AOL, driven by precipitation and dissolution of Mn, which causes increased Hg and MeHg with time; (3) the net production of MeHg in the AOL; and (4) the transport toward the sediment of the particulate fraction, which is the only removal pathway from deep waters and concerns only a small fraction of the total pool.

Results suggest that also in other water bodies, the hypoxic-anoxic conditions induced at the bottom of the water column by summer stratification could promote methylation over areas much larger than previously thought, generating a significant amount of MeHg that would then be advected to the rest of the water column during the subsequent mixing period.

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