A high-resolution marine mercury model MITgcm-ECCO2-Hg with online biogeochemistry

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Abstract. Mercury (Hg) is a global persistent contaminant. Modeling studies are useful means of synthesizing a current understanding of the Hg cycle. Previous studies mainly use coarse-resolution models, which makes it impossible to analyze the role of turbulence in the Hg cycle and inaccurately describes the transport of kinetic energy. Furthermore, all of them are coupled with offline biogeochemistry, so they cannot respond to short-term variability in oceanic Hg concentration. Here we use a high-resolution (horizontal resolution is 1/5°) ocean model (MITgcm-ECCO2, high-resolution-MITgcm) coupled with the online biogeochemistry of Darwin project to simulate the global Hg cycle. The finer portrayal of surface Hg concentrations in estuarine and coastal areas, strong western boundary flow and upwelling areas, and concentration diffusion as vortex shapes demonstrate the effects of turbulence that are neglected in previous models. Ecological events such as algal blooms can cause a sudden enhancement of phytoplankton biomass and chlorophyll concentrations, which can also result in a dramatic change in particle-bound mercury (Hgₐq) sinking flux at the same time in our simulation. In the global estuary region, the inclusion of riverine Hg input in the high-resolution model allows us to reveal the outward spread of Hg in an eddy shape driven by fine-scale ocean currents. With faster current velocities and diffusion rates, our model captures the transport and mixing of Hg from river discharge in a more accurate and detailed way and improves our understanding of Hg cycle in the ocean.

1 Introduction

Mercury (Hg) is a potent neurotoxin that poses health risks to the global population and causes substantial economic losses regulated by Minamata Convention (Zhang et al., 2021). Anthropogenic emissions of Hg from mining and coal combustion severely disturb the biogeochemical cycling of Hg (Nriagu, 1993; Lacerda, 1997; Pacyna et al., 2010; Streets et al., 2011). Mercury can be transported and recycled in the global ocean, atmosphere, and land until buried in deep-sea sediment (Outridge et al., 2018). The ocean is an important part of the biogeochemical cycle of Hg, evidenced by the large air-sea exchange flux and complex biogeochemical transformation that have direct implications for human exposure to Hg through seafood consumption.
The source of marine Hg contains atmospheric deposition, riverine discharge, benthic sediments, and hydrothermal vents (Mason et al., 2012). The air–sea exchange of Hg is driven by the concentration gradients between the atmospheric and seawater interface (Soerensen et al., 2013), which are influenced by atmospheric deposition of $Hg_{aq}^{II}$ and photochemical and dark reduction/oxidation of $Hg_{aq}^{II}/Hg_{aq}^0$ in the seawater (Costa and Liss, 1999; Gardfeldt et al., 2001; Andersson et al., 2011; Ci et al., 2016). In the mixed layer, the reduction reactions of $Hg_{aq}^{II}$ to $Hg_{aq}^0$ and high vapor pressure of $Hg_{aq}^0$ makes it emitted into the atmosphere again (Mason et al., 1995; Amyot et al., 1997; Rolhus and Fitzgerald, 2001). $Hg_{aq}^{II}$ can be absorbed by suspended organic-rich particulate matter to produce $Hg_{aq}^P$ (Fitzgerald et al., 2007). Then $Hg_{aq}^P$ sinks to the subsurface ocean as part of the biological pump and is released in the dissolved phase by the remineralization of sinking particles (Strode et al., 2010).

Marine Hg models have been developed to improve our understanding of the cycling of oceanic Hg. One of the earliest marine model developments was a 2-D slab ocean model coupled with the atmosphere (GEOS-Chem), which considered the red-ox chemistry and transport of inorganic Hg (Strode et al., 2007; Selin et al., 2008; Soerensen et al., 2010). Three-dimensional marine Hg models were applied to simulate the spatiotemporal dynamics of inorganic Hg species in the ocean (Zhang et al., 2014a; Zhang et al., 2014b; Bieser and Schrum, 2016). Later models were developed with improved ocean biogeochemistry. For example, Zhang et al. (2020) and Rosati et al. (2022) coupled Hg cycling with a biogeochemical and ecological model, considering the uptake to and release of Hg from marine biota. The latest developed biogeochemical multi-compartment model for Hg cycling MERCY v2.0 developed by (Bieser et al., 2022), includes all currently known processes controlling marine Hg cycling.

One limitation of earlier models is their inaccurate representation of eddies due to the relative coarse model resolutions. For example, previous studies that simulated the chemistry and transport using the MIT General Circulation Model (MITgcm) had a horizontal resolution of 1° over the global ocean, with improved resolution only in the Arctic and equator (Zhang et al., 2015; Zhang et al., 2020; Huang and Zhang, 2021). The Nucleus for European Modeling of the Ocean (NEMO) framework version 3.1 was developed by Semeniuk and Dastoor (2017), and it adopted a tripolar, variable resolution grid (called ORCA2), which resolves less than a degree in the tropics, the Mediterranean Sea, and the Red Sea and 2° elsewhere. The dissipation of kinetic energy in the turbulent boundary layers and the ocean interior at much smaller scales than the model grid (Wunsch and Ferrari, 2004), and the turbulent mixing controls their transport and storage (Sarmiento et al., 2004), which cannot be accurately represented by coarse-resolution ocean models. In addition, Hg cycling in coastal waters is highly sensitive to anthropogenic disturbances and climate-induced changes (Jonsson et al., 2017; Obrist et al., 2018). The coarse model resolution also cannot reflect the transport and mixing conditions at estuaries and continental shelves. Therefore, models with higher resolutions or parameterizations of eddies are required to improve the accuracy of the simulation of the Hg cycle in the ocean.

Besides, the chemistry of Hg and biogeochemistry is offline so it couldn’t reflect the short-term changes of biogeochemistry, such as algal/phytoplankton bloom events that can significantly impact phytoplankton and zooplankton structure and aquatic environments (Amorim and Moura, 2021). Phytoplankton is the base of pelagic food webs and has been found to
bioaccumulate 105 times more Hg than successive trophic transfers (Watras et al., 1998; Moye et al., 2002; Pickhardt and Fisher, 2007). Therefore, small changes in phytoplankton may significantly impact overall Hg bioaccumulation in the entire system. In the study of Luengen and Russell Flegal (2009), total Hg concentrations decreased when the bloom decayed, possibly because particles (i.e., phytoplankton) with low Hg concentrations were lost from the water column. Another research showed that degradation and sedimentation of algal debris will increase the organic matter load in the sediment (Macalady et al., 2000), which could impact particle-bound Hg levels.

We develop a new simulation for Hg chemistry, transport, and trophic transfer within the high-resolution MITgcm combined with an ecology and biogeochemistry model (the Darwin project; http://darwinproject.mit.edu). The model has a horizontal resolution of 1/5°, allowing the formation of eddies and narrow currents (Menemenlis et al., 2008). The eddy-permitting and delicate resolutions model allows us to examine the influence of physical dispersal, which is a major way of marine substance transport on the transmission of marine Hg, especially over western boundary currents and coastal upwelling regions. We also investigate the influence of online biogeochemistry on Hg chemistry. Finally, we study the fate of the riverine discharge of Hg and the impact of nutrients over coastal waters near big river mouths on the transport of coastal Hg.

2 Model description

2.1 High-Resolution MITgcm

We use MITgcm to simulate the transport and chemistry of Hg in the global ocean. The physical component of the model uses the ECCO2 physical configuration, which addresses mesoscale features in the tropics and allows eddies in subpolar regions where the deformation radius is comparable to the grid scale. It employs the cube-sphere grid projection, permitting relatively even grid spacing throughout the domain, and avoids polar singularities (Adcroft et al., 2004). Each face of the cube consists of 510 grid cells with an average horizontal grid spacing of 18 km. There are 50 vertical levels ranging from 10 m near the surface to approximately 450 m at a maximum model depth of 6150 m in the model. The ECCO2 project provides the best-possible, global, time-evolving synthesis of the most available ocean and sea-ice data at a resolution permitting ocean eddies (Menemenlis et al., 2008).

Our model includes three Hg tracers: dissolved elemental ($Hg_{aq}^0$), dissolved divalent ($Hg_{aq}^{II}$), and particle-bound mercury ($Hg_{aq}^P$). We take atmospheric deposition flux and atmospheric GEM (gas element mercury) concentrations as the upper boundary conditions for $Hg^{II}$ and $Hg^0$, respectively from Zhang and Zhang (2022) and Horowitz et al. (2017). The model takes the initial conditions of ocean Hg concentration from Zhang et al. (2020), which is linearly interpolated to the high-resolution MITgcm model grid. A 6-month free-running simulation is conducted.
2.2 Ocean Biogeochemistry and Ecology

We use a coupled ocean plankton ecology and biogeochemistry model (the Darwin project) within the high-resolution MITgcm, simulating the production and growth of different plankton species and organic carbon remineralization in the marine water column. The Darwin project includes ocean biogeochemistry and ecological variables (e.g., POC and DOC). $Hg_{aq}$ is scavenged by POC to the deep sea through the biological pump. The biogeochemical and biological tracers interact through organic matter formation, transformation, and remineralization. This model simulates the cycling of nutrients (C, N, P, Si, and Fe), phytoplankton growth, zooplankton grazing, and mortality (Dutkiewicz et al., 2009; Ward et al., 2014; Zhang et al., 2015).

There are 51 plankton types (35 phytoplankton and 16 zooplankton) in the complex plankton community, phytoplankton is divided into six functional groups: prokaryotes, picoeukaryotes, coccolithophores (that calcify), diazotrophs (cyanobacteria), diatoms (that utilize silicic acid), and mixotrophic dinoflagellates. The six representative categories of phytoplankton have different sizes, growth rates, grazing, sinking, and affinity to nutrients and other physiological parameters (Kuhn et al., 2019). The online coupled variables coupled with the Darwin project are used to parameterize the Hg chemistry, including deposition and escape of Hg in the surface ocean and photochemical and biochemical redox of Hg in seawater. This is explained in more detail in the following sections.

2.3 Air-sea exchange

The ocean model is coupled to the atmosphere by receiving the atmospheric deposition of $Hg^{II}$ and $Hg^0$. Atmospheric deposition of $Hg^{II}$ is the main input of Hg to the surface environment (Amos et al., 2012), which is largely affected by the amount and type of precipitation and is also influenced by the wind, through controlling the removal of Hg by sea-salt particles. The air-sea exchange of $Hg^0$ is mainly evasion from the ocean to the atmosphere (Mason et al., 2017), which is driven by the concentration gradient at the interface between the atmosphere and seawater (Soerensen et al., 2013) and the piston velocity is a function of wind speed and temperature parameterized by Nightingale et al. (2000). The air-sea exchange of Hg$^0$ is calculated following Strode et al. (2007):

$$ F_{\text{evasion}} = k_w \times \left( \frac{Hg_{aq}^0}{H} \right) $$

where $k_w$ is the gas exchange velocity (piston velocity), and $H$ is the dimensionless temperature-dependent Henry’s Law constant. $k_w$ is related to the sea ice fraction and the instantaneous wind speed. Through the above, the spatial distribution of the evasion flux is controlled by the supersaturation of $Hg^0$ concentrations, and wind speeds, and is associated with sea ice fraction in some regions. The calculation process variables are online coupled, we could simulate a time-sensitive source and sink of marine Hg.
2.4 Hg Chemistry

In the euphotic layer of the ocean, the photo-chemistry and biological-mediated oxidation and reduction reactions between $Hg_{aq}^{0}$ and $Hg_{aq}^{II}$ in our model are based on (Zhang et al., 2014a), with some associated variables are replaced by the ones coupled online. The photochemical oxidation and photochemical reduction first-order rate constants ($k_1$ and $k_2$ in Table 1) are proportional to short-wave radiation at the sea surface ($RAD$ in Table 1), which is coupled by online variables from the Darwin project:

$$RAD = \int_0^{\lambda_{nlam}} PAR(1 - ice) \times 0.2174 / 0.4,$$

where $PAR$ is each band of short-wave radiation at the sea surface, and $nlam$ is the total band of $PAR$. The unit of $PAR$ is $ein \ m^{-2} \ d^{-1}$, so it needs to transfer to $W \ m^{-2}$ by multiplying a coefficient of 0.2174. And the $PAR$ to total upward shortwave radiation is 40%, so we also need to divide this scale coefficient.

The biological oxidation and reduction processes first-order rate constants ($k_3$ and $k_4$ in Table 1) are proportional to the microbial remineralization of particulate organic carbon ($OCRR$ in Table 1). Biologically mediated Hg redox reactions are attributed to the activities of heterotrophic and chemotrophic microorganisms (Mason et al., 1995; Monperrus et al., 2007). Zhang et al. (2014b) scale the biologically mediated reduction of $Hg_{aq}^{II}$ to $Hg_{aq}^{0}$ ($k_4$) to $OCRR$, which is a measure of the microorganism activity. Here the online variable $OCRR$ is one of the focuses of the high-resolution MITgcm coupled with and Darwin project.

$$OCRR = \left( \text{reminDOC} + \text{reminPOC} \right),$$

where reminDOC and the reminPOC are the remineralization part of DOC and POC, respectively.

Partitioning of $Hg_{aq}^{II}$ onto particulate organic carbon (POC) to form HgP, then HgP sinking to deeper waters as part of the biological pump in the ecology and biogeochemistry model. Assuming that $Hg_{aq}^{II}$ and $Hg_{aq}^{0}$ can exchange to each other reversibly and the ratio of them ($k_d$) is proportional to the local particulate organic carbon (POC) level (Zhang et al., 2014b). The POC pool includes both detritus and living phytoplankton, as partitioning of inorganic $Hg_{aq}^{II}$ to living and dead cells are similar (Pickhardt and Fisher, 2007).

$$F_{Hg}^{a} = \left( Hg_{aq}^{0} \times F_{poc-phy} \right) / \left( \text{sumPOC} + \text{sumPhyto} + \text{sumZoop} \right),$$

the sinking flux of HgP(aq) ($F_{Hg}^{a}$, $mol \ m^{-2} \ s^{-1}$) is assumed to be proportional to the sinking flux of POC and phytoplankton ($F_{poc-phy}$, $mol \ m^{-2} \ s^{-1}$). The sumPOC, the sumPhyto, and the sumZoop are the sum of surface chlorophyll concentrations (unit, $mg \ chl \ a \ m^{-3}$), biomass concentration of phytoplankton (unit, $mmol \ C \ m^{-3}$), and biomass concentration of zooplankton (unit, $mmol \ C \ m^{-3}$) respectively. These parameters are coupled with the Darwin project.
2.5 Riverine Hg

Most riverine Hg transported to the ocean is associated with particles (Sonke et al., 2018), and Zhang et al. (2015) infer most of the river Hg discharged to the ocean is refractory. We use the riverine discharge Hg inventory from Liu et al. (2021), including riverine dissolved and particulate Hg. A separate refractory HgP tracer is used to simulate the particulate Hg from rivers (HgPR), reflecting its strong combination with terrestrial source POC. The refractory HgP is released into the dissolved phase only when the organic carbon is remineralized. Furthermore, we use the nutrient export by river from WaterSheds 2 (NEWS 2) developed by Mayorga et al. (2010), which is a global, spatially explicit, multi-element, and multi-form model of nutrient exports by rivers. Global annual export of total N (TN), P (TP), and organic C (TOC) from rivers is estimated to be 44.9 Tg N, 9.04 Tg P, and 317 Tg C, respectively. The distribution among forms for global exoreic exports varies by the element, with DIN (43.7%) and PN (31.2%) dominating nitrogen exports, PP (76.5%) and DIP (16.7%) dominating phosphorus exports, and DOC (53.9%) dominating organic carbon exports. By incorporating these riverine nutrients, we can assess their impact on the ocean ecosystem, including chlorophyll and marine plankton, and their influence on the cycling and sinking of particle-bound Hg.

Table 1. Description of the Main Processes and Input Fields Used in the MITgcm-Hg Simulation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Reference or Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>$F_{dep}$</td>
<td>Atmospheric wet deposition flux of $Hg^{II}$ to the ocean, $mol \ m^{-2} \ s^{-1}$</td>
<td>Zhang and Zhang (2022)</td>
</tr>
<tr>
<td>$F_{evasion}$</td>
<td>Net evasion flux of $Hg^{0}$ across the air-sea interface, $mol \ m^{-2} \ s^{-1}$</td>
<td>Model results</td>
</tr>
<tr>
<td>$F_{riverine}$</td>
<td>Riverine input of Hg, $mol \ m^{-2} \ s^{-1}$</td>
<td>Liu et al. (2021)</td>
</tr>
<tr>
<td>$k_1$</td>
<td>Photooxidation of $Hg^{0}<em>{aq}$ to $Hg^{II}</em>{aq}$</td>
<td>Soerensen et al. (2010), updated by Zhang et al. (2015)</td>
</tr>
<tr>
<td></td>
<td>$k_1 = k_{photo-ox} \times RAD$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$k_{photo-ox} = 4.7 \times 10^{-6}$</td>
<td></td>
</tr>
<tr>
<td>$k_2$</td>
<td>Photoreduction of $Hg^{II}<em>{aq}$ to $Hg^{0}</em>{aq}$</td>
<td>Soerensen et al. (2010), updated by Zhang et al. (2015)</td>
</tr>
<tr>
<td></td>
<td>$k_2 = k_{photo-re} \times RAD$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$k_{photo-re} = 1.56 \times 10^{-6}$</td>
<td></td>
</tr>
<tr>
<td>$k_3$</td>
<td>Biological oxidation of $Hg^{0}<em>{aq}$ to $Hg^{II}</em>{aq}$</td>
<td>Soerensen et al. (2010), updated by Zhang et al. (2015)</td>
</tr>
<tr>
<td></td>
<td>$k_3 = k_{bio-ox} \times OCRR$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$k_{bio-ox} = 9.9 \times 10^{-7}$</td>
<td></td>
</tr>
</tbody>
</table>
3 Model evaluation

3.1 Global inorganic Hg distribution in surface ocean

The global average daily inorganic Hg ($Hg^0$ and $Hg^2$) distribution in surface ocean derived from our high-resolution-MITgcm/Hg simulation compares with the observed total Hg concentrations in global surface seawater (Figure 1). Our model takes into account the effects of temperature, ocean current motion, turbulent mixing, and nearshore topography, all of which contribute to differences in oceanic Hg distribution. As a result, we obtain a more refined global distribution of Hg at the ocean surface. Delicate simulations of inorganic Hg in the surface ocean reflect regional differences and eddy scale changes of concentration, which are aroused by ocean currents or geographical factors. Our high-resolution model shows more detailed changes and a clearer gradient between continent and ocean due to the narrower, swifter currents and strong shear, which can increase the efficiency of mixing. Overall, the distribution of inorganic Hg in our model is similar to that simulated in the
lower-resolution ECCO v4 MITgcm (Fig. S1). The modeled concentration of Hg (mean and standard deviation: 0.60 ± 1.41 pM) is slightly lower than the observed concentration (0.87 ± 0.46 pM) in seawater. As we use a similar Hg chemistry and exchange scheme as previous models (Table 1), the difference in model results reflects a potential dependence of these parameters on model resolution. Besides, the various parameters of the ecosystem vary greatly between this study and previous ones (e.g., different community structures, etc.).

Figure 1. Global simulation of the daily mean total inorganic Hg (0-10 m depth; color, unit pM($10^{-9}$mol m$^{-3}$)) and observation of total inorganic Hg in the ocean surface (0-10 m depth; scatter, unit pM). Observations are from the North Pacific Ocean (Laurier et al., 2004), the Atlantic (Kuss et al., 2011), the Equator & South Atlantic (Mason and Sullivan, 1999), the North Atlantic (Mason et al., 1998; Mason et al., 2001; Bowman et al., 2015), the Southern Ocean (Cossa et al., 2011), and the Arctic Ocean (Kirk et al., 2008; Chaulk et al., 2011; Lehnherr et al., 2011; Heimbuerger et al., 2015).

The simulated total inorganic Hg shows greater global and regional differences compared to previous models. Variables coupled with online biogeochemistry are involved in the chemical production and consumption of inorganic Hg, resulting in modeled Hg concentrations that better match observations (the $Hg^{II}$ deposition flux and $Hg^0$ evasion flux are shown in Fig. S2). Typically, observations are not continuous, and the values of certain adjacent observations may vary significantly. This variability has been observed in the high-precision cruise data (Figure 2 Scattered points in red box). This indicates that the distribution of Hg in the real ocean is highly variable, but this variation was not apparent due to the resolution limits in the previous marine models. Figure 2 illustrates that high-resolution simulated results are closer to observations than previous lower-resolution models (Zhang et al., 2020). In the red box circled position, Figure 2. a(c) can replicate the significant difference in the observed data at nearby locations at the same time (or in a short time).
Figure 2. MITgcm simulation for total inorganic Hg results of high resolution (1/5°×1/5° horizontal) ECCO2 (H-R) and coarser (1°×1° horizontal) resolution ECCO v4 (LLC90) (0-10 m depth; color, unit pM, (Huang and Zhang, 2021)). The scatter is observations from two single high spatial precision cruises, (a,b) from Kuss et al. (2011) Atlantic Ocean observations in a month, and (c,d) from Laurier et al. (2004) North Pacific observations in a day(0-10 m depth; scatter, unit pM).

Ocean currents and turbulence play an important role in the physical transport of Hg. Strong upwelling occurs where the western boundary stream is strong or where cold and warm currents meet, which facilitates the horizontal and vertical exchange of Hg in the ocean, resulting in a heterogeneous distribution and drastic changes. In addition, the effect of turbulence across the ocean causes the spread of Hg in a vortical shape. The finer portrayal of inorganic Hg in Figure 3 demonstrates the influence of ocean currents and turbulence. The Kuroshio in the western Pacific Ocean and the Gulf Stream in the North Atlantic Ocean are the most powerful Western Boundary Currents in the world, and the western boundary flow region is the area with very active vortices and a fast flow rate. This region shows more significant concentration changes and clear eddy structures in the distribution of inorganic Hg compared to the surrounding ocean (Figure3.a and Figure3.b). Other areas in the distribution of Hg with significant changes are the Pacific and Atlantic equatorial regions, which are possibly influenced by the westward
Equatorial Warm Currents. Moreover, the transport and mixing of Hg are stronger and faster where cold and warm currents meet (Figure 3.b and Figure 3.d). It was also found that cold currents have a milder effect on Hg transport and dispersion as compared to warm currents (Figure 3.c and Figure 3.d).

Figure 3. Total inorganic Hg Simulation of regions with ocean currents (0-10 m depth; color, unit pM). Zoom for (a) the Northwest Pacific Ocean, (b) the Northwest Atlantic Ocean, (c) the North Indian Ocean, and (d) the South Atlantic Ocean.

3.2 Riverine Hg discharge

Rivers are important sources of coastal Hg. We model Hg from rivers as a refractory tracer following Zhang et al. (2015). Driven by the riverine input (1000 Mg a⁻¹, 893–1,224) from Liu et al. (2021), after transferring and mixing for six months in our simulation (Figure 4), most of the Hg (93.7%) is buried in estuarine sediments (ocean depth < 55 m), while 6% is buried...
in shelves (ocean depth between 55 m and 180 m), and less than 0.01% export to the open ocean (ocean depth >180 m). We model a higher percentage of refractory particulate oxidized Hg (HgPR) buried in estuarine sediments and less is buried in shelves and exports to the open ocean, compared with previous studies. For example, 72-73% buried in estuarine sediments, and 5-6.4% transported to the open ocean by Zhang et al. (2015) and Liu et al. (2021). Indeed, with a higher resolution, coastal currents can be resolved with river discharge being transferred along the seashores. In contrast, the coarse resolution model assumes even mixing at the first grid point for a long distance, leading to an overestimate of the transmission to distant oceans.

Figure 4. Global simulations of the fate of refractory particulate oxidized Hg (HgPR) discharged by rivers. Estuaries, shelves, and open oceans are defined as ocean regions with depth < 55 m, 55–185 m, and >185 m, respectively. Coastal ocean refers to estuaries and the shelf. (a)-(c) is the simulation results of the last day of 4-6 months, respectively.

The spatial distribution of riverine input refractory particulate oxidized Hg across the global ocean is driven by both variabilities in HgPR concentrations and suspended sediment discharges (Figure 5). Major rivers and highly contaminated systems are visible on the global chart (e.g., Yangtze, Amazon, Ganges). The high Hg discharges from Mexican rivers into the Pacific Ocean reflect the high total suspended sediment flux values, as previously described by (Liu et al., 2021). To gain a clearer understanding of the fate of HgPR discharge, we zoom in on the major river mouth regions for further investigation. The rivers selected for the analysis include the Rio Grande River in North America, the Yellow River, the Yangtze River, the Amur River, the Pearl River, and the Zigzag River in Asia, the Orinoco River and the Amazon River in South America, and the Congo River in Africa. We find that riverine inputs of HgPR to the coastal ocean have a limited impact on Hg concentrations in surface seawater beyond the shelf region. HgPR is transported to the left or right along the coast by the Coriolis force, the influence of currents and eddies leads to dispersion slightly further away from the coast.
In the case of the Amazon River, which is the largest exporter of freshwater and particulate matter to marine waters globally (Amos et al., 2014), generating up to a $1.3 \times 10^6 \text{ km}^2$ plume, and extensive muddy bottoms in the equatorial margin of South America (Moura et al., 2016). Our model results show that Hg discharges from the river are transported northwestward along the Brazilian continental shelf without significant offshore transport before intersecting the Equatorial Counter Current (Figure 5c). This is consistent with the model simulation by Zhang et al. (2015), but the plume we simulate is much finer and narrower due to the higher resolution of the model. This is also consistent with the observation (Mason and Sullivan, 1999), which shows that there is an apparent absence of a riverine Hg plume and low total Hg concentrations in the open Equatorial Atlantic Ocean near the Amazon River mouth.

Hg exports from rivers are important to specific coastal areas, Liu et al. (2021) have raised further concerns about riverine Hg and a deeper understanding of the role of rivers in the global Hg cycle. The discharge of HgPR from rivers greatly impacts environmental pollution in coastal and shelf regions. Our model results show that HgPR is transported from the coast to the ocean in an outwardly extending eddy shape, indicating the influence of turbulence mixing and the transmission of ocean eddy energy (Wyrtki et al., 1976). Mesoscale eddies are widely present in the global ocean and dominate the ocean’s kinetic energy. They are also involved in energy cascade at different scales (Shang et al., 2013). Clayton et al. (2013) proposed that increasing the resolution of physical models can enhance dispersal rates by narrowing and accelerating boundary currents, resolving swift transports associated with eddy stirring. To better illustrate the dynamic process of the transport and mixing of HgPR, we create global and regional videos using data from the entire simulation period (Fig. S6 and Fig. S7). The results show that HgPR spreads outward in an eddy shape, driven by HgPR concentrations and influenced by the kinetic energy transmission.
by ocean eddies and the mixing of turbulence. This further emphasizes the important role of eddy-driven processes and turbulence mixing, in the transport and distribution of riverine Hg in coastal and shelf regions.

3.3 Impact of Biological Pump

The spatial distribution of modeled $Hg_{aq}^P$ sinking flux (Figure 6c) follows the distribution of POC out of the euphotic zone (Fig. S3). Globally, the $Hg_{aq}^P$ sinking flux from the surface (0-10m) is $1.12 \text{ mol m}^{-2} \text{a}^{-1}$. The POC pool encompasses both detritus and living phytoplankton in aquatic food webs. Phytoplankton is known to accumulate Hg from their surrounding aqueous environment and is considered to be the primary entry point for Hg into marine food chains (Pickhardt and Fisher, 2007). The modeled distributions of all phytoplankton and zooplankton biomass in the euphotic zone are shown in Figures 6a and 6d, respectively. And the modeled distributions of phytoplankton and zooplankton are consistent with previous model results and observations (Kuhn et al., 2019; Wu et al., 2020). The distribution of surface chlorophyll concentrations (Figure 6b) also aligns well with the phytoplankton distribution. The model simulates elevated sinking flux of HgP in high-latitude oceans of the northern hemisphere due to enhanced nutrient input from terrestrial erosion, in the equatorial region of the Southern Ocean, and along the west coasts of continents with strong upwelling.

Figure 6. Global distribution of the daily mean ecological variables (0-10m). (a) the sum of 35 kinds of biomass (in carbon content) concentration of phytoplankton, mmol C m$^{-3}$, the areas highlighted by boxes are possible algal regions, and the green box area is the final selected algal region; (b) the sum of 35 kinds of surface chlorophyll concentrations (color, unit mg chl a m$^{-3}$); (c) HgP(aq) sinking flux (color, unit fM m$^{-2}$ s$^{-1}$); (d) the sum of 16 kinds of biomass (in carbon content) concentration of zooplankton, unit mmol C m$^{-3}$.)
Online biogeochemistry makes Hg more sensitive to environmental changes at shorter time scales. In our study, we try to establish a correlation between plankton and the sinking of particle-bound Hg. To achieve this, we need to observe and compare the changes in the HgP sinking flux during periods when plankton undergo significant fluctuations. One such process is algal bloom caused by eutrophication. Eutrophication is the process by which the primary production in a water body increases, in severe cases can lead to large blooms of algal. Algal bloom mostly occurs in coastal areas, Dai et al. (2023) found that algal blooms occurred in 126 out of the 153 coastal countries examined. These blooms can significantly affect the biogeochemistry in the water column, including the distribution and cycling of nutrients, organic matter, and other elements, including Hg. The decomposition of algal debris can increase the flux of particulate organic matter (Macalady et al., 2000) and associated Hg to the sediments. Furthermore, the uptake of Hg by phytoplankton can also increase during algal blooms, which can lead to higher Hg concentrations in the sinking organic matter. Investigating the changes in the HgP sinking flux during algal blooms can provide valuable insights into the biogeochemical processes controlling the fate and transport of Hg in aquatic systems.

We identify algal bloom events on a global scale for the simulated period. The ICES meeting (1984) concluded that a bloom is a deviation from the normal phytoplankton mass. Therefore, we can design such an algorithm to detect algal bloom events: during the simulated period, an algal bloom event is if it has the largest deviation from the average in a certain range of time. In this way, we capture 5 possible bloom regions, located in the Red Sea (detected region’s center position:34.4°E, 26.2°N), the Northwest Pacific (155.5°E, 33.4°N), the Bering Strait (171.2°W, 67.5°N), the west coast of the United States (124.5°W, 42.2°N), and the southern coast of the Arabian Peninsula (52.2°W, 15.4°N), respectively (Figure 6a). Upon analyzing the time series data, we observe significant fluctuations and peaks in the phytoplankton biomass at two regions, Point A in the northwestern Red Sea and Point B in the northwest Pacific Ocean (Figure 7). To further confirm that these fluctuations are indeed bloom events, we examine the nutrient levels of nitrogen and phosphorus in the water at the two regions (Fig. S4). As elevated nitrogen and phosphorus elements are known to be associated with eutrophication and algal blooms, we find that the nutrient levels at Point A are above the average during the peak days, suggesting that the occurrence of algal blooms here. It is possibly attributed to higher sea surface temperature or increased use of nitrogen fertilizers in neighboring countries such as Egypt, and Saudi Arabia (Ritchie et al., 2022), both of which can contribute to the blooms (Dai et al., 2023). Previous research has also reported algal bloom events in the Red Sea (Mohamed, 2018). The intense fluctuations in phytoplankton biomass at Point A during the bloom period (possibly between mid-March and early May) are indicative of a bloom event, as shown by the abrupt peaks in plankton biomass, chlorophyll concentration, and Hg_sinking flux, all of which are significantly higher than the average (Figure 7).
We determine the approximate extent of blooms by generating a global distribution of plankton, chlorophyll, and \( Hg_{aq} \) sinking flux on the peak day (Fig. S5). Our results indicate that the Point A bloom spanned two latitudes and two longitudes, possibly influenced by coastal topography. On the peak day of Point A bloom, the \( Hg_{aq} \) sinking flux is 2.7 times greater than the overall average and 5.2 times greater than the non-blooming mean. This suggests that the sinking of particle-bound Hg is influenced by ecological changes in seawater. During algal blooms, the increased phytoplankton biomass creates more particulate matter
to bind with Hg, leading to accelerated organic particle scavenging and remineralization in the water column. Thus, in this way, increase the sinking flux of particle-bound Hg after a short time. Our discovery was facilitated by integrating our model with online biogeochemistry, which promptly reflected the abrupt fluctuations in phytoplankton and enabled us to quickly observe changes in the sinking of Hg particles.

### 4 Conclusions

In this study, we estimate the global biogeochemical cycling of Hg in a state-of-the-art physical-ecosystem ocean model (High-resolution-MITgcm/Hg), realizing Hg transport in eddy-scale. One of the biggest advantages of our model is the promoted resolution so that the small-scale changes of marine Hg can be captured. We use the online biogeochemistry fields to participate in the ocean transformation and reaction processes of Hg. Online biogeochemistry is involved in the reaction between elemental Hg and divalent, and the process of particle-Hg forming as well as the remineralization processes. The coupling with the Darwin project further enhances the specificity of the Hg biogeochemistry in the MITgcm model. Overall, this approach provides a comprehensive understanding of the global biogeochemical cycling of Hg and its interactions with the marine ecosystem.

The horizontal resolution of 18km allows our model to simulate the turbulence, which is omitted in previous global models. Turbulent mixing and shear effects affect the transport of materials (including Hg) in the ocean, so only a high enough resolution can refine this physical process. With this advanced physical configuration, we introduce the latest river discharge inventory from Liu et al. (2021) and 12 nutrients exported by river from WaterSheds 2 (Mayorga et al., 2010). Our high-resolution model provides a more detailed depiction of the fate of riverine Hg discharged to the global oceans. The ocean plays a critical role in the global Hg cycle, and the large size of the ocean is a major site for atmospheric mercury transport, transformation, and deposition. Global ocean models of Hg require a better understanding of biogeochemical controls on Hg speciation. We qualitatively analyzed the biogeochemistry effect on the particle scavenging process of inorganic Hg. The high-resolution MITgcm can help us better predict the transport and fate of Hg in the ocean and its impact on the global Hg cycle. This provides us with more advanced tools and methods to address natural and human-disturbing global Hg e Data availability. The data supporting the findings of this study are available within the article and its Supplement.

Code availability. The MITgcm model code is available at https://github.com/MITgcm/MITgcm.git. The code of MITgcm-ECCO2-Hg in this paper is permanently archived on Zenodo at https://doi.org/10.5281/zenodo.7932859 (Zhu and Zhang, 2023).

Data availability. The data supporting the findings of this study are available within the article and its Supplement.
Author contribution. Zhang Y and Zhu S conceived the idea and designed the model experiments. Zhang Y and Jahn O developed high-resolution MITgcm model. Zhu S and Zhang Y modified the code of the MITgcm-ECCO2-Hg model. Wu P and Zhang S improved part of the code. Zhu S performed the simulations, conducted the analysis, and wrote the paper. Zhang Y and Zhu S edited the paper.

Competing interests. The contact author has declared that neither they nor their co-author has any competing interests.

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References


