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Isotopes and otolith chemistry provide insight into the biogeochemical history of mercury in southern flounder across a salinity gradient†

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Methylmercury (MeHg) continues to pose a significant global health risk to wildlife and humans through fish consumption. Despite numerous advancements in understanding the mercury (Hg) cycle, questions remain about MeHg sources that accumulate in fish, particularly across transitional coastal areas, where harvest is prominent and Hq sources are numerous. Here we used a unique combination of Hq and nutrient isotopes, and otolith chemistry to trace the biogeochemical history of Hg and identify Hg sources that accumulated in an economically important fish species across Mobile Bay, Alabama (USA). Fish tissue Hg in our samples primarily originated from wet deposition within the watershed, and partly reflected legacy industrial Hg. Results also suggest that little Hq was lost through photochemical processes (<10% of fish tissue Hq underwent photochemical processes). Of the small amount that did occur, photodegradation of the organic form, MeHg, was not the dominant process. Biotic transformation processes were estimated to have been a primary driver of Hg fractionation (\sim 93%), with isotope results indicating methylation as the primary biotic fractionation process prior to Hg entering the foodweb. On a finer scale, individual lifetime estuarine habitat use influenced Hg sources that accumulated in fish and fish Hg concentrations, with runoff from terrestrial Hg sources having a larger influence on fish in freshwater regions of the estuary compared to estuarine regions. Overall, results suggest increases in Hg inputs to the Mobile Bay watershed from wet deposition, turnover of legacy sources, and runoff are likely to translate into increased uptake into the foodweb.

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Environmental significance

Despite advancements in understanding the mercury cycle, questions remain about sources of methylmercury that accumulate in fish, particularly across a salinity gradient, where fishery harvest is prominent. Here, isotope and otolith data identified that Hg in an estuarine foodweb primarily originated from local and regional emissions that were deposited in the estuary watershed, and entered through freshwater runoff. These mercury inputs were largely methylated rather than being demethylated or photochemically degraded. Unique isotopic signatures also identified relative sources of Hg in fish collected from different regions across the estuary. In total, this research identified sources and drivers of Hg in a coastal food web and demonstrates the importance of understanding Hg biogeochemical pathways in a changing environment.

Introduction

Mercury (Hg) poses significant global health risks to wildlife and humans through the consumption of fish contaminated with monomethylmercury (MeHg), a neurotoxic and bioavailable form of mercury that bioaccumulates and biomagnifies in foodwebs. ¹⁻³ Despite decades of research, ⁴⁻⁶ questions remain about how Hg cycles through the environment and the sources of Hg that ultimately accumulates in fish tissue. ⁷ This

knowledge is particularly vital for estuarine and coastal species that make-up a large portion of fisheries for human consumption and thus human exposure to MeHg.⁸ Sources of MeHg in estuarine and coastal fish tissues are often uncertain and can come from more than one source, even in relatively small transitional coastal regions (*e.g.*, freshwater run-off, estuarine sediments, marine sediments, anaerobic microcosms surrounding particulate organic matter in the water column, among others).⁹⁻¹² Here we define source as meaning all possible sources of Hg in fish tissue, including but not limited to inorganic and organic Hg entering an ecosystem through deposition, runoff, and turnover of buried Hg, and areas prone to increased methylation and thus increased net MeHg accumulation in foodwebs. Further, many of these estuarine Hg sources are influenced by anthropogenic activities that provide

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opportunities for mitigation (*e.g.*, limiting nutrient run-off, landscape restoration, reducing local emissions, remediation of legacy pollution).¹³ As such, filling these knowledge gaps, which largely stem from uncertainties in Hg sources and pathways (defined as Hg movement and transformations through the biogeochemical Hg cycle), is needed.^{11,14-16}

Specifically, the biogeochemical Hg cycle involves the release of three inorganic forms of Hg into the atmosphere, largely from anthropogenic sources (e.g., combustions of fossil fuels): elemental (Hg⁰), oxidized (Hg(II)), and particulate (PHg).^{7,17} While oxidized and particulate forms of inorganic Hg deposit relatively close to emission sites, the long atmospheric residence time of Hg⁰ allows it to travel to distant sites before transformation to Hg(II) and deposition. 18,19 As such, Hg deposition in an estuarine ecosystem has the potential to include a mixture Hg emitted from distant, local, and regional sources. Direct releases of inorganic Hg to the terrestrial and aquatic environment from past industrial uses (e.g., industrial use of mercury catalysts) have also resulted in pools of legacy Hg in soils and sediments.12,13 Consequently, Hg(II) and PHg enter aquatic systems through rainfall (wet deposition), settlement (dry deposition), runoff, or turnover of previously deposited legacy Hg in sediment and soils, where it can be methylated by bacteria into MeHg.20,21 Net rates of methylation in aquatic systems can vary substantially due primarily to dissolved oxygen concentration, pH, local land practices, dissolved organic matter (DOM) concentrations and type, and microbial composition.²²⁻²⁵ These factors affect net methylation rates by influencing Hg transport, bioavailability, and whether Hg transformation processes will be dominated by MeHg and Hg(II) degradation and reduction (e.g., volatilization of Hg⁰ that can atmosphere) or by methylation the bioaccumulation.25,26

Stable isotope applications developed over the last few decades provide tools to better understand the Hg biogeochemical cycle and identify sources of Hg that accumulate in fish tissue. 11,27-29 Mercury, for instance, has seven naturally occurring stable isotopes that do not fractionate appreciably during trophic transfer. 10,30 Mercury isotope signatures are therefore preserved when incorporated into the foodweb. 10 Odd Hg isotopes undergo mass-independent fractionation (MIF) from photochemical processes (represented by Δ^{199} Hg and Δ^{201} Hg) and can be used to distinguish between the type of photochemical process that dominated Hg fractionation before entering the foodweb (e.g., photodegradation of MeHg to Hg(II) or photoreduction of Hg(II) to Hg⁰). 31,32 Mass dependent fractionation is represented by δ^{202} Hg and results from both abiotic and biotic processes. Because MDF occurs with both abiotic and biotic processes, and MIF of odd isotopes (e.g., Δ^{199} Hg) is associated only with abiotic photochemical processes, the slope of $\Delta^{199} Hg$ to $\delta^{202} Hg$ in relation to previous well-established slopes of this relationship³¹ can provide insight into the extent of MDF that occurred as a result of photochemical processes; with the remaining portion of MDF attributable to microbially-driven biotic processes (i.e., methylation, and microbially-driven reductive and oxidative demethylation). 25,31,32 Mass independent fractionation of even Hg isotope ²⁰⁰Hg $(\Delta^{200}{\rm Hg})$ also provides insight into the relative importance of atmospherically derived Hg, wherein positive values reflect Hg sources from precipitation and negative values reflect gaseous elemental Hg exposed to upper-atmosphere higher-energy UV light during long-distance transport. Using these relationships, Hg isotopes have successfully linked fish tissue Hg to sources such as non-point urban runoff, atmospheric deposition and transport, terrestrial runoff, and industrial releases. 9,11,12,21

Like Hg isotopes ratios, bulk C isotope ratios (δ^{13} C) and source amino acid N isotope ratios (e.g., phenylalanine, $\delta^{15}N_{Phe}$) remain largely unaltered during trophic transfer, preserving isotope signatures at the base of the foodweb. As such, C isotopes can distinguish whether energy supplying a foodweb originated from C3 versus C4 plants³⁴ and whether sources originated from benthic or pelagic algae.35 Nitrogen isotope ratios for source amino acids (e.g., δ¹⁵N_{Phe}) can determine the relative amount of nitrate-rich (e.g., benthic) versus nitrate-poor (e.g., atmospheric) sources at the base of the foodweb. 36,37 Bulk N isotopes (δ^{15} N), which average source and trophic amino acid N isotope ratios, have commonly been used in environmental toxicology and ecology to identify the trophic position of individual organisms, providing insight into contaminant bioaccumulation.35,37 Given that N, C, and Hg accumulate in individual fish through the same dietary items, together these data can refine our understanding of Hg sources that accumulate in an ecosystem foodweb.

An additional limitation to understanding Hg sources to fish tissue across a salinity gradient is knowledge about fish habitat use, which can influence the sources of Hg that an individual fish is exposed to. Otoliths, bony structures in fish ears, accrue calcium carbonate throughout the life of an individual with annual marks (annuli) separating regions deposited in specific years of life. Within each annual region in the otolith, trace elements (e.g., strontium [Sr], barium [Ba]) from the surrounding environment (i.e., water) are incorporated (after passing through multiple membranes and biological barriers in the gills, blood, and endolymph) in approximate proportion to their concentration in the environment by randomly substituting for calcium (Ca) in the calcium carbonate matrix of the otoliths.38,39 Once incorporated into the otolith, the concentration of these elements is permanent as otoliths are inert and are not resorbed during periods of stress, providing a lifetime record of salinity exposure for an individual fish.³⁸⁻⁴⁰ As a result, otolith chemistry (defined here as the ratio of Sr: Ca measured across otolith annuli) has been a useful tool in providing inferences about habitat use and residency patterns of fishes across estuarine salinity gradients. 41,42 In sum, using otolith chemistry in combination with tissue Hg isotopes, total tissue Hg, and nutrient isotopes (defined here as C and N isotope data) can help to refine the spatial scale at which we distinguish sources of Hg in fish tissues across transitional ecosystems, such as an estuarine salinity gradient; where numerous and variable Hg sources may contribute to mercury concentrations in economically important coastal fisheries.^{8,43}

In this study, we targeted southern flounder (*Paralichthys lethostigma*); a species that inhabits a wide variety of estuarine

salinities (i.e., freshwater to hypersaline)41,44,45 in the northern Gulf of Mexico, where they support economically important recreational and commercial fisheries. Juvenile southern flounder inhabit freshwater and estuarine environments to adult sizes before moving offshore to overwinter and spawn.46 Because southern flounder inhabit freshwater, estuarine, and marine environments,47,48 understanding sources of Hg to this species provides insight on how Hg is transferred through foodwebs across these various ecosystems and to humans. As such, our objectives were to quantify the sources and pathways of Hg bioaccumulation for this economically important, and frequently consumed species in the Mobile Bay, Alabama (USA) estuary and to identify finer scale sources of Hg in individual fish across the salinity gradient using Hg isotopes, bulk N isotopes, amino acid compound-specific N isotopes, bulk C isotopes, and otolith chemistry.

Materials and methods

Sample collection

The Mobile-Tensaw River is the primary source of freshwater to Mobile Bay and is considered the fourth largest river system in the continental United States. Three additional smaller sources of freshwater also contribute to the bay: Dog River (237 km² watershed), Fowl River (184 km² watershed), and Week's Bay (521 km²).^{50,51} Samples were collected from ten sites located along a 60+ km seasonal salinity gradient of saltmarshes, bays, tidal creeks, and freshwater ecosystems. Sites were sampled one to two times monthly during May-July of 2018 and March, May-July of 2019. Within the Mobile Bay estuary, three salinity sampling regions (freshwater-oligohaline [<5%], mesohaline [5–18%], polyhaline [18-30%]) were targeted. Sample sites within each region were randomly stratified by available suitable southern flounder habitat types (e.g., mud flats, oyster beds, tidal marsh).

Southern flounder between 270 mm and 547 mm TL (average age in years = 0.9 ± 0.4) were collected using gill nets, electrofishing, and hook-and-line. Gillnets (30 m by 2.4 m with 127 mm stretch mesh) targeted large juveniles and adults (≥100 mm TL) at meso- to polyhaline sites by soaking nets for four hours (two 2 hour sets). Nets were set parallel to shore with a hook towards shore at the downstream end. At freshwater sites, pulsed DC boat electrofishing (Midwest Lake Electrofishing Systems Infinity Box) was used along shorelines. Six 15 minute boom mounted electrofishing transects were conducted during each site visit. Hookand-line sampling was conducted after all standardized sampling was completed or in areas not accessible by the previous methods. At each site, date, time, GPS coordinates, and water depth were recorded at the beginning and end of each sampling transect or gillnet set. All southern flounder collections were conducted according to use guidelines outlined in IACUC protocol #AUP2018-001 at Clemson University.

Chemical analyses of fish tissue and otoliths collected from **Mobile Bay**

Dorsal axial tissue samples (N = 23) were processed using metals-cleaned stainless-steel instruments and freeze-dried

following EPA standard protocols for measuring total Hg (USEPA method 7473).52 Previous research analyzed southern flounder muscle tissue collected at four sites across the freshwater-oligohaline portions of the Mobile-Tensaw River Delta for both total Hg and MeHg and found that MeHg comprised 87% (±20% SD) of total Hg and did not differ across the Mobile-Tensaw River Delta.⁵⁰ We therefore assumed that muscle tissue from the same species in this study were consistent with these results. However, it should be noted that southern flounder collected in this study were on average larger and from additional sites (i.e., Lower Mobile Bay, Weeks Bay; Fig. 1) not sampled by Farmer et al.,50 and thus could have had higher proportions of MeHg to total Hg in muscle tissue.44 Dried muscle tissue samples were ground to a homogenous powder and analyzed for total dry weight Hg concentrations using a Prodigy-7 dual-view ICP optical emission spectrometer. Ti was used as an internal standard for mass bias correction. The procedural blank was 0.03 ppb, equivalent to $\sim 0.5\%$ of the analytical signal. Seven of 23 samples were run in duplicate for total Hg. The mean relative standard deviation of duplicates was $9.98\% \pm 5.5\%$ SD. The mean relative difference for DORM-2 Hg

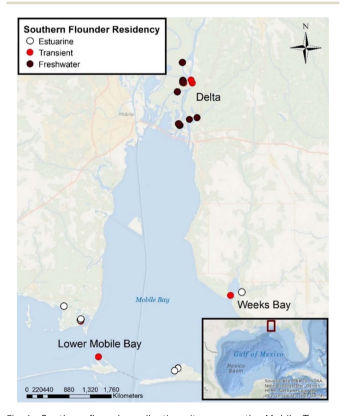


Fig. 1 Southern flounder collection sites across the Mobile-Tensaw River Delta and Mobile Bay, Alabama (USA) for isotope analyses. Sites were sampled in three locations: the Delta (freshwater riverine areas of the Mobile-Tensaw River Delta), Week's Bay (small estuary that drains into lower Mobile Bay and experiences freshwater to polyhaline conditions), and Lower Mobile Bay (sites located within the lower Mobile Bay with salinities ranging from mesohaline-polyhaline). Lifetime salinity exposure classifications are indicated by color on the map and were classified based on otolith chemistry and water chemistry analyses.42

standard reference material (certified value as 4640 ± 260 ng g⁻¹ dry weight) was 1% \pm 1% SD (N=2; values = 4543 and 4640.2 ng g⁻¹ dry weight). Total dry weight Hg concentrations were converted to total wet weight Hg concentrations using the percent moisture measured for each sample. Total wet weight Hg are reported as $\mu g g^{-1}$ ww in this study.

All N and C isotope analyses were conducted by the Biogeochemical Stable Isotope Laboratory in the School of Ocean and Earth Science and Technology at the University of Hawaii, with methods described in Dale et al., 53 Sackett et al., 37 and Wall et al. 54 Briefly, bulk N and C isotopes (δ^{15} N and δ^{13} C) were analyzed using a continuous flow isotope ratio mass spectrometer (Costech ECS 4010 Elemental Combustion System using a Zero Blank Autosampler) coupled with a Thermo Scientific Delta V Advantage through a Thermo Scientific Conflo IV. Isotopic values are reported in δ-notation relative to atmospheric N2 and V-PDB (Vienna Peedee Belemnite), for N and C respectively. A glycine and an in-house white muscle tissue tuna standard, characterized for $\delta^{15}N$ and $\delta^{13}C$ using international reference materials (δ¹⁵N: USGS 32, USGS 34, USGS 35, NIST 3; and δ^{13} C: NBS 18, NBS 19, NIST 1547) and verified by an independent laboratory, were used to quantify and correct isotopic results. Known standard values were 11.4 $\delta^{15}N$ and -36.5 $\delta^{13}C$ for glycine and 14.2 $\delta^{15}N$ and -16.2 $\delta^{13}C$ for tuna. Standards were run in triplicate before samples were analyzed and then with every 7 samples thereafter. All isotope data accuracy was $\leq \pm 0.2\%$. Procedural blanks were 0.06 ppb or 1.3% of the analytical signal.

For Hg isotope analyses, aliquots of homogenous dried muscle tissue samples were digested using 3 mL of aqua regia in a 40 mL amber glass vessel by heating on a hot plate at 120 °C for 4 hours. After digestion, the supernatant was extracted and made-up to 20 mL with 0.002 M BrCl. Digested samples were analyzed for Hg isotopes using a Nu Plasma II MC-ICP-MS in standard wet plasma cones. An ESI ICP Hydride generation system was used to generate Hg⁰ vapor by quantitative reduction of Hg(II) in solution using stannous chloride (3% w/v in 1 M HCl). The Hg⁰ vapor was introduced directly into the ICP. The Tl internal standard was nebulized using a 50 µL min⁻¹ selfaspirating micro-concentric PFA nebulizer and a cyclonic/Scott dual spray chamber. Instrument tuning and gain calibration of the detectors was performed daily. An NIST cinnabar standard (RM 8610) was used to verify data accuracy and agreed with values reported in the literature (N = 5; mean, 1 SD; $\delta Hg^{202} =$ -0.50 ± 0.03 , $\delta Hg^{201} = -0.44 \pm 0.04$, $\delta Hg^{200} = -0.27 \pm 0.05$, $\delta Hg^{199} = -0.16 \pm 0.04, \Delta Hg^{199} = -0.04 \pm 0.03, \Delta Hg^{201} = -0.06$ \pm 0.04).55 Mercury isotope analyses were conducted at the Trent University Water Quality Centre, Peterborough ON Canada. Hg isotope ratios were reported in per mil (%), referenced to the NIST SRM 3133 mercury standard, where:

$$\delta^{xxx}Hg = \{ [(^{xxx}Hg/^{198}Hg)sample/ (^{xxx}Hg/^{198}Hg)_{NIST3133}] - 1 \} \times 1000$$
 (1)

and xxx equals 199, 200, 201, or 202. Mass dependent fractionation (MDF) is represented by δ^{202} Hg and results from both abiotic and biotically-driven Hg transformations. Mass

independent fractionation (MIF) is measured as the difference between the measured δ^{xxx} Hg, where xxx equals 199, 200, 201. Values are predicted based on MDF and the δ^{202} Hg, and are reported as Δ^{xxx} Hg, with calculations as follows:

$$\Delta^{199} Hg = \delta^{199} Hg - (\delta^{202} Hg \times 0.2520)$$
 (2)

$$\Delta^{200} Hg = \delta^{200} Hg - (\delta^{202} Hg \times 0.5024)$$
 (3)

$$\Delta^{201} Hg = \delta^{201} Hg - (\delta^{202} Hg \times 0.7520)$$
 (4)

Odd Hg isotopes undergo MIF from photochemical processes (represented by Δ^{199} Hg and Δ^{201} Hg) and can be used to distinguish between the type of photochemical transformation that dominated Hg fractionation.

Amino acid compound-specific stable isotope (AA-CSIA) analyses were conducted at the Biogeochemical Facility at the University of Hawaii following the same protocols reported in Dale et al.,53 Sackett et al.,37 and Wall et al.54 Briefly, dried tissue samples were subjected to acid hydrolysis, esterification of the carboxyl terminus and trifluoracetylation of the amine group prior to being introduced into a Delta V or MAT 253 mass spectrometer interfaced with a Trace GC gas chromatograph through a GC-C III combustion furnace (980 °C), reduction furnace (650 °C) and liquid N₂ cold trap. Internal reference compounds, L-2-aminoadipic acid (AAA, $\delta^{15}N = -6.2\%$) and L-(+)-norleucine (Nor, $\delta^{15}N = 19.06\%$) of known nitrogen isotope compositions, were co-injected with samples and used to normalize the measured isotope values of unknown amino acids and determine accuracy and precision. Amino acids targeted here included glycine (Gly), phenylalanine (Phe), lysine (Lys), leucine (Leu), and glutamic acid (Glu). All amino acids were analyzed in triplicate and isotope values are reported in δnotation relative to atmospheric N2. The mean relative standard deviations (RSD) for internal reference values L-(+)-norleucine and L-2-aminoadipic acid were 0.99% and 7.4%, respectively. The mean measured values for internal reference compounds Nor and AAA were $19.3\%_0 \pm 0.19\%_0$ SD (difference of $-0.2\%_0$) and -4.8% $\pm 0.35\%$ SD (difference of -1.4%), respectively. Mean RSD for triplicate amino acid results used in this study were low, indicating high precision in analytical methods and study results (glycine = 3.1%, leucine = 1.4%, phenylalanine = 8.6%, glutamic acid = 0.85%).

The elemental composition of fish ear bones (otoliths) and water samples were used to determine the lifetime salinity exposure of each southern flounder collected across the salinity gradient in Mobile Bay as part of a separate study. As such, detailed otolith chemistry and residency pattern classification methods for these samples are described in Chrisp *et al.* Briefly, following methods from Farmer *et al.* In otoliths were ablated from the core to the distal edge of the otolith along a straight transect parallel to the sulcal groove. Otolith chemistry analysis targeted Ca and Sr concentrations every 0.6 seconds using an Agilent 7700x quadrupole inductively coupled plasma mass spectrometer (ICPMS) coupled to a 213 nm Nd:YAG NWR laser. Using Ca (37.69%) as an internal standard, elemental concentrations were converted to molar ratios. Water

samples were processed for elemental concentrations of Ca and Sr with the same ICPMS system in solution mode coupled with an Agilent autosampler. Concentrations of each element measured in water were converted to molar ratios using Ca as a standard to compare with otolith chemistry. 42,56,57 Using these data, southern flounder were classified into one of three lifetime residency patterns (i.e., freshwater, estuarine or transient) based on lifetime salinity exposure (as inferred from Sr to Ca molar ratios: Sr: Ca) across the otolith transect. To accomplish this, nonlinear models were fit to the data to quantify relationships between water Sr: Ca versus ambient sample site salinity. Predicted values from these nonlinear models were then multiplied by a partition coefficient to develop a link between expected otolith Sr: Ca and ambient salinity. 42 Finally, a regime shift algorithm was used to detect any significant shifts in Sr: Ca and thus, salinity, across the lifetime of the individual.^{58,59} Otolith Sr: Ca equivalent to ≤1 psu salinity was classified as a freshwater resident for a given time step, while otolith Sr: Ca equivalent to >1 psu salinity was classified as an estuarine resident for a given time step. If more than 90% of the time steps across the lifetime of an individual fell into one classification (i.e., freshwater or estuarine), residency patterns were assigned to that classification. If neither classification comprised 90% of the transect, then a 'transient' classification was assigned to indicate a fish that either moved between freshwater and estuarine habitats or a fish that resided in an area that experienced seasonal changes in salinity.

Statistical analyses

To examine sources of nitrogen to southern flounder across Mobile Bay, and the relative trophic position of individual fish, we calculated the weighted mean $\delta^{15}N$ value for source amino acids using glycine, lysine, and phenylalanine measured from AA-CSIA and bulk δ¹⁵N data following the same methods as Sackett et al.³⁷ Briefly, we calculated the weighted mean δ^{15} N value for source amino acids glycine, lysine, and phenylalanine using AA-CSIA:

$$\delta^{15} N_{\text{source}} = \frac{\sum \frac{\delta^{15} N_x}{\sqrt{\sigma_x^2}}}{\sum \frac{1}{\sqrt{\sigma_x^2}}}$$
 (5)

where $\delta^{15}N_x$ is the $\delta^{15}N$ value of a specific source amino acid and σ_r is the standard deviation of triplicate isotope analysis of the specific amino acid.40 Given the cost of AA-CSIA analyses, only a portion of our samples were analyzed for these values (16 of 23). We calculated trophic position for those samples where AA-CSIA data were available using glutamic acid and phenylalanine in an equation developed by Chikaraishi et al.:60

$$TP = ((\delta^{15}N_{Glu} - \delta^{15}N_{Phe} - \beta)/TEF) + 1$$
 (6)

where TP is trophic position, $\delta^{15}N_{Glu}$ is the $\delta^{15}N$ value for glutamic acid, and $\delta^{15} N_{\rm Phe}$ is the $\delta^{15} N$ value for phenylalanine with constants developed by Bradley *et al.*⁶¹ ($\beta = 3.6 \pm 0.5$ SD; TEF = 5.7 \pm 0.3 SD). This approach has been used successfully in past research.46 For bulk δ15N values, which were analyzed for all

collected tissue samples, β and TEF are unknown. Consequently, exact trophic position for individuals with only bulk δ¹⁵N data could not be calculated. However, a proxy for trophic position was estimated for all samples by correcting bulk $\delta^{15}N$ values using δ15N_{source} values (eqn (5)) averaged for each lifetime residency pattern (i.e. freshwater, transient, and estuarine). A similar approach was successfully used in previous research.³⁷ Specifically, mean $\delta^{15}N_{\text{source}}$ values (see eqn (5)) for each lifetime residency group were subtracted from bulk δ15N values for each individual classified within that group. This process normalized bulk δ¹⁵N values for individuals from different lifetime residency groups and provided a relative measure of trophic position without making assumptions about the magnitude of TEF for bulk data.37 This approach also allowed for a measure of relative trophic position for all 23 fish samples to be used in model analyses.

Southern flounder tissue Hg isotopes, $\delta^{15}N_{source}$, $\delta^{13}C$, total Hg concentrations, and relative trophic position were used to evaluate differences in Hg sources, dietary sources of N and C, and relative trophic position across sampling regions and lifetime residency classifications of fish across Mobile Bay. The Δ^{199} Hg/ Δ^{201} Hg ratio resulting from the regression relationship between Δ^{199} Hg to Δ^{201} Hg was used to determine whether MIF was primarily driven by photodegradation of MeHg (Δ^{199} Hg/ Δ^{201} Hg ratios of 1.2 to 1.36) or the photoreduction of Hg(II) to Hg^0 ($\Delta^{199}\mathrm{Hg}/\Delta^{201}\mathrm{Hg}$ ratios of \sim 1.0) prior to Hg being ultimately methylated and incorporated into the foodweb. 10,31,32 The Δ^{199} Hg/ δ^{202} Hg ratio was also evaluated in relation to experimentally derived values from Bergquist and Blum³¹ to distinguish between the relative contribution of photochemical versus biotic fractionation processes that occurred prior to Hg entering the foodweb.10 Previous research has also used these Hg isotope ratios (i.e. Δ^{199} Hg/ Δ^{201} Hg ratio and Δ^{199} Hg/ δ^{202} Hg ratio) to provide an understanding of these Hg biogeochemical processes.9-11,62 Also, like previous work,11 we included results from several of these published studies for comparison to the results presented here in Fig. 2. Based on laboratory trials by Bergquist and Blum,³¹ a Δ^{199} Hg/ δ^{202} Hg ratio of 2.43 is representative of Hg undergoing only MIF from photochemical processes, while a ratio of 0 is representative of little to no MIF or MDF from photochemical processes, and instead suggests microbial Hg transformations dominated MDF prior to Hg entering the foodweb. As such, dividing Δ^{199} Hg/ δ^{202} Hg ratio from southern flounder tissue by the slope representative of 100% MIF from photochemical processes (i.e., 2.43; eqn (7)) gives insight into the relative proportion of photochemical processes that fractionated Hg prior to its incorporation into the foodweb. Subtracting this proportion by 1 in-turn provides the remaining proportion of Hg fractionation that was driven by microbial processes (eqn (7)) rather than photochemical processes. It should be noted that while there is evidence that dark abiotic demethylation can also occur under controlled laboratory settings and contribute to MDF, the strong reductants and oxidants used in these experiments are typically absent from the natural environment.25 It is therefore unlikely that dark abiotic demethylation contributed to demethylation processes in this study or Hg MDF values. Additionally, the

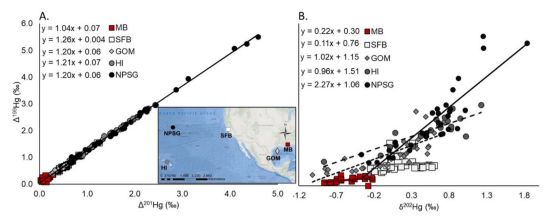


Fig. 2 Regressions of fish tissue Hg isotope values as indicators of (A) the type of photochemical processes that occurred (P < 0.01 for all relationships) and (B) the relative amount of photochemical *versus* microbial processes that occurred to fish tissue Hg prior to its incorporation into the foodweb (P < 0.01 for all relationships except MB and SFB).³¹ These data were collected from Mobile Bay, Alabama (MB, N = 23, (A) $R^2 = 0.64$, (B) P = 0.09), San Francisco Bay (SFB, N = 35 (A) $R^2 > 0.99$, (B) P = 0.57), the Gulf of Mexico (GOM, N = 32, (A) $R^2 > 0.99$, (B) $R^2 = 0.61$), the waters surrounding Hawaii (HI, N = 19, (A) $R^2 > 0.99$, (B) $R^2 = 0.72$), and the North Pacific Subtropical Gyre (NPSG, N = 33, (A) $R^2 > 0.99$, (B) $R^2 = 0.78$).

foundational study on Hg isotopes by Bergquist and Blum³¹ showed no abiotic dark demethylation in experiments, even when Hg concentrations were high. Therefore, using the $\Delta^{199} \text{Hg}/\delta^{202} \text{Hg}$ ratio representative of 100% MIF from photochemical processes $(2.43)^{31}$ in comparison to the measured $\Delta^{199} \text{Hg}/\delta^{202} \text{Hg}$ ratio as described above allows for the calculation of:

% of biotically driven fractionation =
$$(1-([\Delta^{199}Hg/\delta^{202}Hg\ ratio]/2.43))\times 100\quad (7)$$

Using this equation, we estimated the relative percent of microbial *versus* photochemical processes that fractionated Hg in fish tissue prior to entering the foodweb.

Total Hg, fish total length, δ^{13} C, δ^{15} N_{Source}, relative TP, δ^{202} Hg, Δ^{199} Hg, Δ^{200} Hg, and Δ^{201} Hg were compared for significant differences among southern flounder lifetime residency classifications to understand how different habitat uses across the estuary may have influenced Hg sources to fish tissues (Table 1).42 To better understand sources of Hg among southern flounder collected across the salinity gradient in Mobile Bay, Δ^{199} Hg with Δ^{201} Hg, Δ^{199} Hg with δ^{202} Hg, and Δ^{199} Hg with δ^{13} C, were also analyzed with k-means cluster analysis ($\alpha = 0.05$; k = 3clusters). Additionally, a multiple linear regression model (N =23) was used to predict how lifetime salinity exposure as a proxy for lifetime estuarine habitat use, total length, and trophic position impacted the concentration of total Hg in fish tissue (α = 0.05). However, given that fish total length and trophic position can be interactive (e.g., the effect of total length can often depend on trophic position) and are well known primary drivers of total Hg in fish tissue, 22,37 we included an interaction between total length and relative trophic position in this model to ensure these measures did not confound results (total Hg values that accounted for this interaction are summarized as least squares means of total Hg in Table 1). This model was also

re-run using sample collection sites (Delta = freshwater habitats within Mobile-Tensaw River Delta that drains the 114 000 km² watershed, Week's Bay = shallow estuary with 521 km² watershed that drains into lower Mobile Bay with freshwater-polyhaline conditions, and Lower Mobile Bay = lower portion of Mobile Bay with mesohaline-polyhaline salinities; Fig. 1) in place of lifetime residency classifications, to determine if fish capture location was more important in explaining tissue Hg concentrations than lifetime estuarine habitat use based on salinity. Models were compared using Akaike's Information Criterion for small sample size (AIC_c), where the model with the lowest AIC_c value is the model that best explains the variation in the data. Total Hg data were normal and homoscedastic.

Results and discussion

Biogeochemical history of Hg across Mobile Bay

MIF from photochemical processes in Mobile Bay. Odd Hg isotope data from southern flounder collected across the Mobile Bay estuary indicated that photochemical processes had little influence on Hg fractionation prior to Hg entering Mobile Bay foodwebs, and of the little that did occur, it was dominated by the photochemical reduction of Hg(II) to Hg⁰ (Table 1, Fig. 2A). Δ^{199} Hg and Δ^{201} Hg data from southern flounder across Mobile Bay were low and clustered near zero (Table 1; Fig. 2A). The near zero values for odd Hg isotopes suggest little Hg photoreduction or photodegradation occurred in Mobile Bay prior to incorporation into the foodweb. 9,31,62,64 The published literature supports that only photochemical processes involving Hg(II) and MeHg impart large (>0.5%) MIF on odd Hg isotopes, with Δ^{199} Hg/ Δ^{201} Hg ratios ranging from \sim 1.0 to 1.36, indicative of photochemical processes acting on Hg prior to its incorporation into the foodweb.21,31,62,64 A variety of studies using fish tissues have reported a very narrow range of Δ^{199} Hg/ Δ^{201} Hg ratios between 1.2 and 1.36 (Fig. 2).9,11,31,62,65 Based on previous laboratory work by Bergquist and Blum31 using aqueous Hg(II) and

ndividual, or the mean 8¹⁵N_{source} for the lifetime residency group from bulk tissue 8¹⁵N values), and least squares modeled (LSM) Hg concentrations (accounting for total length and relative comparisons of otolith microchemistry and water chemistry from Chrisp et al.⁴² Mean fish total length, relative trophic position (TP; estimated by subtracting the 8¹⁵N_{source} value for an able 1 Mean total Hg concentrations, source nutrient isotopes, and Hg isotopes measured in southern flounder tissue with different lifetime salinity exposure classifications based on rophic position (TP); N = 23; P < 0.01) are also reported for each grouping. Parenthetical values represent standard error

Salinity exposure classifications	N	Total Hg $(\mu g g^{-1} ww)$	LSM total Hg^a ($\mu g g^{-1} ww$)	Total length ^a (mm)		Bulk δ^{13} C ^a (%) δ^{15} N _{source} (%)	Relative TP	$\delta^{202}\mathrm{Hg}^{a}\left(\% ight)$		$\Delta^{199} \mathrm{Hg} \left(\%\right) \Delta^{201} \mathrm{Hg} \left(\%\right)$	Δ ²⁰⁰ Hg (‰)
Freshwater Transient	8 8 8	0.08 (0.01) $0.06 (0.004)$	$0.09 (0.01) \\ 0.06 (0.01)$	315.3 (10.9) 387.4 (17.8)	-28.38 (0.54) -22.73 (1.34)	$0.89 (0.04) \\ 0.82 (0.05^b)$	12.74 (0.26) 11.79 (0.34)	$\begin{array}{c} -0.62 \ (0.04) \\ -0.55 \ (0.08) \end{array}$	0.17 (0.02) 0.20 (0.05)	0.07 (0.01) $0.14 (0.03)$	0.03 (0.01) $0.01 (0.01)$
Estuarine All fish	7 23	0.07 (0.02)	0.06 (0.01) 0.07 (0.005)	397.6 (35.2) 365.39 (14.6)	$-20.93 (0.50) \\ -24.14 (0.84)$	0.71 (0.06) $0.81 (0.04)$	11.93 (0.43) 12.16 (0.21)	-0.37 (0.05) -0.52 (0.04)	0.20 (0.06) 0.19 (0.02)	0.13 (0.05) 0.11 (0.02)	$0.05 (0.01) \\ 0.03 (0.01)$

ANOVA significantly different among salinity exposure classifications, P < 0.05. b Because the $\delta^{15}N_{source}$ sample size (N) here is two the parenthetical value represents the range.

MeHg in the presence of natural dissolved organic matter, values of 1.2 to 1.36 indicate photodegradation of MeHg as the dominant photochemical transformation process. Here the Δ^{199} Hg/ Δ^{201} Hg ratio measured from southern flounder collected across Mobile Bay was 1.04. Based on the results of Bergquist and Blum, ³¹ a Δ^{199} Hg/ Δ^{201} Hg ratio of 1.0 indicates photoreduction of Hg(II) to Hg⁰ was primarily responsible for the MIF that occurred before Hg was ultimately methylated and entered Mobile Bay foodwebs. This result can occur when Hg(II) is photoreduced to Hg⁰, either in the atmosphere or after deposition and volatilization back to the atmosphere, causing MIF before atmospheric transformation back to Hg(II) and deposition in this estuarine ecosystem. These results also indicate that of the Hg in Mobile Bay biotically transformed to MeHg, the vast majority either entered the benthos or foodweb rather than being photochemically degraded back to Hg(II). If photochemical degradation of MeHg was a dominant process fractionating Hg prior to its ultimate incorporation into Mobile Bay foodwebs, this would have resulted in a Δ^{199} Hg/ Δ^{201} Hg ratio closer to 1.2 to 1.36, as has been seen for other ecosystems (Fig. 2A). Additionally, while relationships between odd Hg isotopes (Δ^{199} Hg/ Δ^{201} Hg ratio) during photochemical reduction have recently been shown to be sensitive to complexing ligands in controlled laboratory studies,66 in estuarine and marine fish tissue samples from natural systems, slope values close to 1.0, as seen here, were representative of the Hg(II) being photochemically reduced to Hg⁰. 9,25,52,64

Estimating the relative importance of microbial versus photochemical Hg transformations in Mobile Bay. Southern flounder tissue Δ^{199} Hg values and δ^{202} Hg values (Table 1), along with a Δ^{199} Hg/ δ^{202} Hg ratio (0.22) that was not significantly different than zero (P = 0.09; Fig. 2B) suggested that microbial processes likely dominated the transformations of MeHg that ultimately accumulated in southern flounder tissues across Mobile Bay. These results can be inferred because experimentally derived Δ^{199} Hg/ δ^{202} Hg values of 2.43 were established to represent Hg fractionation from only photochemical processes31 with values lower than 2.43 representing a lesser influence from photochemical fractionation. Indeed, Δ^{199} Hg/ δ²⁰²Hg values close to zero demonstrate little to no photochemical processes influenced Hg fractionation, and instead signify microbial processes dominated Hg transformations.31 For instance, Hg isotope ratios in biota in a remote location of the north Pacific (Δ^{199} Hg/ δ^{202} Hg = 2.27) demonstrated a high proportion of Hg was fractionated though photochemical degradation¹⁰ (Fig. 2B), while Hg isotope data $(\Delta^{199}\text{Hg}/\delta^{202}\text{Hg} =$ 0.11) from biota in San Fransisco Bay9 (USA; Fig. 2B) demonstrated very little photochemical processes contributed to Hg fractionation. Indeed, using the experimentally derived Δ^{199} Hg/ δ²⁰²Hg ratio of 2.43 as representing 100% photochemical fractionation,49 the ratio of 2.27 from fish collected in the north Pacific (Fig. 2B) suggests \sim 93% (2.27/2.43 \times 100) of Hg was photochemically fractionated before ultimately being methylated and incorporated into the north Pacific foodweb. For San Fransisco Bay, using this same approach results in an estimate of \sim 5% (0.11/2.43 \times 100) of photochemical Hg fractionation occurring before Hg entered the foodweb; leaving ~95% of Hg

fractionation attributable to microbial transformation processes (see eqn (7)). Mercury in fish from Mobile Bay were similar to San Fransisco Bay in that photochemical processes only accounted for \sim 9% (0.22/2.43 \times 100) of Hg fractionation, leaving the remaining \sim 91% of Hg fractionation being microbially-driven. Further, microbially-driven Hg transformation processes such as Hg methylation, demethylation, and reduction, can all cause MDF. However, some have demonstrated that MDF during Hg(II) methylation to MeHg preferentially enriches MeHg with lighter isotopes, resulting in lower or more negative δ^{202} Hg values, while demethylation of MeHg to Hg(II) often results in Hg(II) enriched in heavier isotopes and thus higher δ^{202} Hg values. ^{67,68} Thus, if bioavailable Hg(II) enters an ecosystem and undergoes several rounds of Hg methylation and demethylation before ultimately being methylated and entering the foodweb, MeHg in fish tissue would have more positive and higher δ^{202} Hg values (see San Fransisco Bay data depicted in Fig. 2B). However, when bioavailable Hg(II) enters an ecosystem and primarily undergoes methylation and uptake into the foodweb, δ^{202} Hg values in fish tissues are likely to be more negative, as seen here in fish tissue in Mobile Bay (Table 1). However, previous studies have also suggested that industrial sources of Hg result in distinct narrow δ^{202} Hg (-0.50 \pm 0.27 SD) and Δ^{199} Hg (0.00 \pm 0.06 SD) values.¹³ The δ^{202} Hg values measured in fish tissue here fell within this range while Δ^{199} Hg values were slightly higher than those suggested to be a signature for industrial sources (Table 1). The Mobile Bay watershed is home to several sources of industrial legacy pollution. 69 These include long-term industrial and commercial operations, some of which took advantage of the presence of salt domes to provide chloride for mercuric-chloride reduction processes to produce chemical compounds. One area, approximately 40 km upstream of the Mobile Bay-Tensaw River Delta was deemed a Superfund Site for Hg, with sediments capped in the mid-1980s. In an investigation of industrial legacy sources to sediments and biota by Janssen et al.12 in the Saint Louis River estuary in Minnesota (USA), a sample site with similar δ^{202} Hg and Δ^{199} Hg values in biota, as were seen for southern flounder across Mobile Bay, was suggested to be only partially influenced by industrial legacy sources compared to other sites. The authors also suggested having δ^{202} Hg values in-line with industrial legacy signatures but Δ^{199} Hg values that are slightly higher than industrial signatures suggested Hg deposited into environmental conditions with high net Hg methylation and bioaccumulation contributed to these results. Given the similarities in our study, enhanced methylation and bioaccumulation of Hg deposited to the Mobile Bay watershed, along with partial contributions from upstream legacy Hg sources were likely mixed in runoff into the estuary and contributed to the isotope signatures in fish tissue measured here as well.

Hg isotope indicators of deposition. Mass independent fractionation of even Hg isotope $^{200}{\rm Hg}\,(\Delta^{200}{\rm Hg})$ provides insight into the relative importance of atmospherically derived Hg, wherein positive values reflect Hg sources from precipitation and negative values reflect gaseous elemental Hg exposed to upper-atmosphere higher-energy UV light during long-distance

transport. The largely positive Δ^{200} Hg values (19 of 23 individuals had positive Δ^{200} Hg values; Table 1) measured in Mobile Bay fish tissue suggested that Hg from long-range atmospheric transport was not a primary source of Hg entering the foodweb, but instead was indicative of wet deposition within the watershed from local and regional sources. 19,33,70 Data collected by the National Atmospheric Deposition Program have demonstrated that the Gulf Coast, including Mobile Bay, has some of the highest wet deposition rates of Hg in the United States.71 Additionally, Engle et al. 19 suggested that rapid Hg(II) and PHg deposition from local sources are important Hg inputs to this region, while Ren et al. 17 measured average wet deposition rates of 309 \pm 407 ng per m² per week, with results indicating Hg deposition was influenced by nearby local and regional atmospheric Hg sources in coastal Mississippi (adjacent to our study site in Alabama). Other studies have also noted that the northern Gulf of Mexico experiences high atmospheric wet deposition due to unique atmospheric conditions, relatively high Hg concentrations in rain, and high rainfall amounts. 70,72,73

Effect of lifetime estuarine habitat use on sources of southern flounder tissue Hg. Southern flounder classified as having different patterns of lifetime habitat use across Mobile Bay showed differences among sources of tissue Hg, C, and N (Fig. 3). A k-means cluster analysis using Δ^{199} Hg and Δ^{201} Hg demonstrated that estuarine residents were significantly (P =0.05) clustered together with higher Δ^{199} Hg and Δ^{201} Hg values compared to significantly clustered freshwater residents (Fig. 3A). Differences in Δ^{199} Hg and Δ^{201} Hg between lifetime residency groups demonstrate that bioaccumulative Hg in freshwater residents photochemically fractionated to a lesser extent than estuarine residents (Table 1). Similar conclusions were inferred for isotopic results in eastern and western cohorts of fish in Lake Erie (USA), where higher Δ^{199} Hg values in an eastern cohort of fish were attributed to a greater contribution of photochemically fractionated Hg compared to a western cohort of the same species.⁶⁵ It should be noted that Δ^{199} Hg values from this study conducted in the Great Lakes (USA) were much higher (Δ^{199} Hg = 2.27 to 7.16%) than those found here (Table 1). The authors noted that measured Δ^{199} Hg in fish from the Great Lakes were comparable to those in shallow open ocean fish and attributed these results to the low DOC concentrations in offshore regions of the Great Lakes and DOC content from in situ biotic production with low humic content; similar to DOC concentrations and content in shallow open ocean ecosystems. Higher δ^{202} Hg values (0.17 to 2.06) and a Δ^{199} Hg/ Δ^{201} Hg ratio of 1.26 from this Great Lakes study also demonstrated that in situ MeHg photodegradation was the primary driver of MIF. Other studies have also linked the content and concentrations of DOM and DOC to MeHg photodegradation.65,74,75 Specifically, high inputs of DOM and DOC have been shown to absorb and block radiation and complex or scavenge the iron and OH radicals needed for photodegradation, reducing Hg loss from an ecosystem through photochemical processes. High DOM concentrations have been measured across Mobile Bay76 and are likely linked to the low level of photochemical MIF observed for Hg in southern

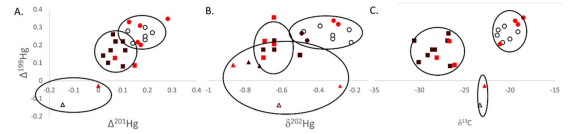


Fig. 3 Isotopic data ((A) Δ^{199} Hq to Δ^{201} Hq; (B) Δ^{199} Hq to δ^{202} Hq; (C) Δ^{199} Hq to δ^{13} C) measured in southern flounder muscle tissue across the salinity gradient in Mobile Bay, Alabama (USA). Lifetime salinity exposure classifications were assigned to an individual based on otolith chemistry and water chemistry analyses and are indicated by color (white = estuarine, bright red = transient, and black = freshwater; colors are consistent with those in Fig. 1). 42 k-means cluster analyses were conducted for each pair of isotope relationships depicted. Hollow ellipses represent the 95% confidence area for each grouping. Data points assigned to distinct clusters are based on shape, with circles representing a cluster, squares representing a second cluster, and triangles a third cluster.

flounder across Mobile Bay. A cluster analysis between Δ^{199} Hg and δ^{202} Hg demonstrated clustered separation between lifetime freshwater and lifetime estuarine residents, with freshwater residents largely having more negative δ²⁰²Hg (Fig. 3B). High influxes of DOM and DOC from freshwater runoff likely contributed to the distinction between freshwater residents in closer proximity to runoff than estuarine residents. Also, as discussed previously, more negative $\delta^{202} \mbox{Hg}$ values are indicative of methylation being the primary transformation pathway and the specific $\delta^{202} Hg$ value also indicates that a portion of the Hgin southern flounder tissue originated from industrial legacy pollution (-0.50 ± 0.27 SD).^{12,13}

Cluster analysis using Δ^{199} Hg and δ^{13} C resulted in very distinctive groupings of tissue data (Fig. 3C). Sources of carbon among lifetime salinity exposure classifications were significantly different (P < 0.01; Table 1) and drove much of the separation in these data (Fig. 3C). Mean δ^{13} C was lowest for freshwater residents at -28.4% (SE = 0.54%). δ^{13} C values between -22% and -30% are indicative of C3 plant carbon sources from terrestrial freshwater runoff.34,35 Consequently, these results show that individuals classified as freshwater residents and those classified as transient individuals that were captured in the same region as freshwater residents (Fig. 3C) relied heavily on foodwebs based on freshwater terrestrial C3 plant carbon sources (e.g., agricultural runoff from prominent Alabama crops such as cotton, peanuts, soybeans, and fruit are C3 plants).77 Those fish classified as estuarine residents occupied more saline habitats throughout their life history and were therefore furthest from freshwater outflows in Mobile Bay. Estuarine residents also had the highest mean δ^{13} C at -20.9%(SE = 0.50%) among lifetime residency classifications (P < 0.01); signifying a lesser influence of terrestrial carbon inputs (-22%to -30% and the mixing of terrestrial carbon sources with marine benthic sources $(-17\%_{00})^{34,35}$ Marine benthic sources of carbon are more enriched in the heavier isotope (C^{13}) with $\delta^{13}C$ values of approximately -17% compared to marine phytoplankton with δ^{13} C values of approximately -22%. This distinction occurs because of the larger boundary layer that surrounds benthic algae (>1 mm) compared to pelagic algae (\sim 10 µm), which limits access to the preferred lighter 12 C isotope for photosynthesis. Southern flounder are a benthic

species, further supporting that the higher δ^{13} C values for estuarine residents were likely from the mixing of freshwater terrestrial (δ^{13} C $\sim -28\%$) and marine benthic algae (δ^{13} C \sim -17% carbon sources. The low level of photochemical degradation indicated by odd Hg isotopes across the entire Mobile Bay estuary supports the notion that freshwater terrestrial inputs high in DOC likely caused more turbid waters and less sunlight penetration for photochemical processes, even to those habitats occupied by estuarine residents in the lower reaches of Mobile Bay.⁷⁸ δ¹⁵N_{source} data further supported these conclusions with significant differences in source N among lifetime salinity exposure classifications (P = 0.047; Table 1). The highest values were found for freshwater residents and lowest for estuarine residents (Table 1). High (or less negative) δ¹⁵N_{source} data suggest a more nitrate-rich source of nitrogen (e.g., agricultural runoff) than lower (or more negative) values, which suggest a more nitrate-poor source (e.g., atmospheric nitrogen), to a foodweb. 36,37 As such, the high positive $\delta^{15}N_{\text{source}}$ values in Mobile Bay would further suggest a terrestrial and thus more nitrate-rich base to the foodwebs across the entire estuary that was more pronounced in freshwater residents that occupied habitats closer to the source of those inputs. Together these data demonstrate that terrestrial freshwater run-off was a primary source of N, C, and Hg to Mobile Bay foodwebs, and that those individuals that resided in the freshwater portion of the estuary through their life history were more influenced by these terrestrial inputs. Those individuals with lifetime residency furthest from freshwater terrestrial inputs were also influenced by terrestrial N, C, and Hg inputs but had mixed terrestrial sources with marine benthic N, C, and Hg sources.

For all cluster analyses (Fig. 3), transient individuals (those that experienced both the freshwater and more saline estuarine conditions throughout their life history) were largely grouped with estuarine and freshwater residents based on their location of capture (Fig. 1). These results suggest that fish captured together likely occupied the same habitat and relied on the same food sources for much of their life history, but that those individuals classified as transient ventured into other habitats more so than their freshwater residency and estuarine residency counterparts. As with most groupings, each could be brokendown to a finer scale, particularly for southern flounder

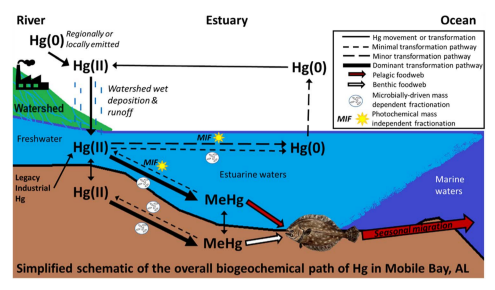


Fig. 4 Mercury (Hg), C, and N isotopes, and otolith chemistry identified methylmercury (MeHg) sources and pathways before accumulating in an estuarine foodweb.

residency, which likely exists on more of a spectrum. For instance, in all clusters, two individuals were consistently and significantly grouped together despite different lifetime salinity exposure classifications (Fig. 3). Both individuals were captured in Week's Bay (Fig. 1), suggesting isotope signatures identifying sources of carbon and tissue Hg were specific to this bay. Previous research has suggested that nutrient-rich, acidic, low salinity riverine inputs to Weeks Bay forms a wedge that floats on top of denser, high salinity, high pH water that flows into the upper portion of this bay.79 The mixing zone of these two layers of water has warm, low dissolved oxygen conditions with low oxidation-reduction potential, all of which are favorable for Hg methylation. 22,23,79 This study also suggested that wet deposition was the primary source of inorganic Hg to the watershed of Weeks Bay, a notion supported by positive tissue Δ^{200} Hg values measured in our study (Δ^{200} Hg = 0.06 for both individuals).⁷⁹ As such these results suggest that Hg in fish from Week's Bay had been deposited in the Week's Bay watershed where it underwent very little to no photochemical processes through terrestrial freshwater run-off with carbon that was depleted in C^{13} ($\delta^{13}C \sim$ -28%₀₀), 35 and efficiently methylated in Week's Bay near the benthos, where benthic carbon sources were more enriched in C^{13} ($\delta^{13}C \sim -17\%$). The mixing of these different sources of carbon and Hg to the Weeks Bay foodweb likely resulted in unique and intermediate δ^{13} C values (mean δ^{13} C = 23.1%) and unique Hg isotope signatures in fish tissue (Fig. 3).

Differences in the biogeochemical histories and sources of tissue Hg among lifetime residency groups also resulted in differences in total tissue Hg concentrations. Results from a multiple linear regression demonstrated that when fish length and relative trophic position were accounted for, southern flounder lifetime residency classifications had significantly different concentrations of tissue Hg (P < 0.01, N = 23, $R^2 = 0.58$ Table 1), with freshwater residents having significantly higher concentrations than estuarine or transient fish (Table 1).

Previous studies have also observed negative relationships between the salinity where fish were captured and tissue Hg concentrations. These results were also significant (P = 0.01, N = 23, $R^2 = 0.47$) when three general sample locations (Delta, Week's Bay, and Lower Mobile Bay; Fig. 1) were used in place of lifetime residency exposure classifications based on salinity; however, lifetime residency classifications had a lower AIC_c value (ΔAIC_c using sampling locations = 5.5) and thus were better at explaining the variation in southern flounder tissue Hg concentrations. Results showed that the more direct biogeochemical path of methylated Hg being incorporated into the foodweb from terrestrial freshwater runoff in the Mobile-Tensaw River Delta (Fig. 1) resulted in higher Hg accumulation in the foodweb in the freshwater portion of the estuary and, specifically, in southern flounder with lifetime freshwater residency patterns.

Conclusions

Southern flounder results across Mobile Bay have important implications for consumptive health risks. Primary sources of Hg in fish tissue in Mobile Bay serve as a proxy for Hg entering and accumulating in Mobile Bay foodwebs. Here results found sources of fish tissue Hg largely stemmed from local and regional atmospheric Hg that entered the estuary through wet deposition within the watershed, and runoff from terrestrial environments, with turn-over and run-off of legacy Hg sources from direct industrial releases at least partly contributing to fish Hg body burdens (for a simplified depiction of these results see Fig. 4). Our results also indicate Mobile Bay conditions are conducive to Hg methylation. As such, additional Hg loading to Mobile Bay or its watershed from wet deposition and runoff, or storm systems that turnover and release legacy Hg from sediment is likely to result in increased net MeHg and uptake into foodwebs.¹³ Of further concern, large storm systems (e.g.,

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hurricane) are increasing in frequency and intensity because of climate change81-84 and can increase each of the fish tissue Hg sources identified in Mobile Bay (local and regional wet deposition and terrestrial runoff, legacy Hg turn-over). Because southern flounder also move offshore to spawn during the fall and winter, freshwater residents and at least some transient and estuarine residents, are expected biological vectors transporting Hg from terrestrial watersheds to marine ecosystems in the Gulf of Mexico. Moreover, southern flounder are fundamental to Alabama's commercial and recreational fishery with the majority of those harvested having transient salinity exposure classifications.42 Because transient individuals in this study often had Hg sources similar to estuarine and freshwater residents collected in similar locations, it is likely that harvested fish contain Hg sources from across the estuarine ecosystem. As such, increases in Hg inputs to this aquatic ecosystem are likely to translate into higher concentrations in Mobile Bay foodwebs, predators that consume prey from this ecosystem, and in human consumers of those fish. Lastly, this study demonstrates the utility of combining N, C, and Hg isotopes with otolith chemistry to better understand the biogeochemical cycle that ultimately accumulates Hg in economically important coastal fisheries.

Conflicts of interest

There are no conflicts to declare.

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