
Stable isotope analyses revealed the influence of foraging habitat on mercury accumulation in tropical coastal marine fish

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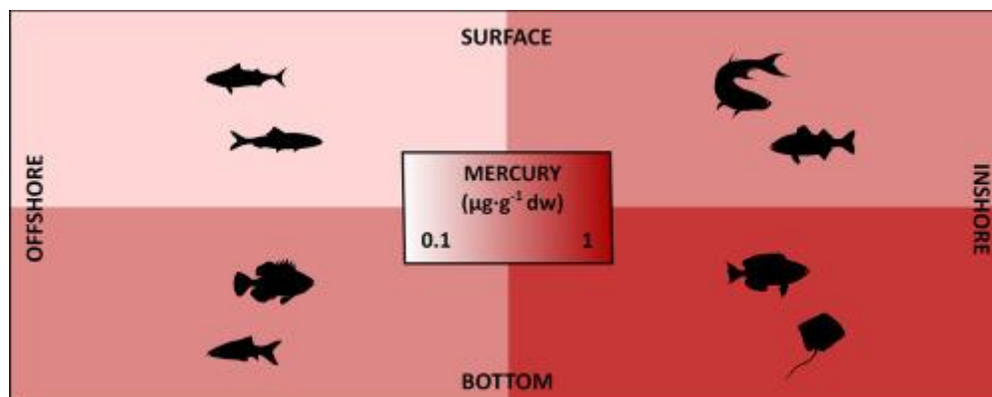
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Abstract :

Bioaccumulation of toxic metal elements including mercury (Hg) can be highly variable in marine fish species. Metal concentration is influenced by various species-specific physiological and ecological traits, including individual diet composition and foraging habitat. The impact of trophic ecology and habitat preference on Hg accumulation was analyzed through total Hg concentration and stable isotope ratios of carbon ($\delta^{13}\text{C}$) and nitrogen ($\delta^{15}\text{N}$) analyses in the muscle of 132 fish belonging to 23 different species from the Senegalese coast (West Africa), where the marine ecosystem is submitted to nutrients inputs from various sources such as upwelling or rivers. Species-specific ecological traits were first investigated and results showed that vertical (i.e. water column distribution) and horizontal habitat (i.e. distance from the coast) led to differential Hg accumulation among species. Coastal and demersal fish were more contaminated than offshore and pelagic species. Individual characteristics therefore revealed an increase of Hg concentration in muscle that paralleled trophic level for some locations. Considering all individuals, the main carbon source was significantly correlated with Hg concentration, again revealing a higher accumulation for fish foraging in nearshore and benthic habitats. The large intraspecific variability observed in stable isotope signatures highlights the need to conduct ecotoxicological studies at the individual level to ensure a thorough understanding of mechanisms driving metal accumulation in marine fish. For individuals from a same species and site, Hg variation was mainly explained by fish length, in accordance with the bioaccumulation of Hg over time. Finally, Hg concentrations in fish muscle are discussed regarding their human health impact. No individual exceeded the current maximum acceptable limit for seafood consumption set by both the European Union and the Food and Agriculture Organization of the United Nations. However, overconsumption of

some coastal demersal species analyzed here could be of concern regarding human exposure to mercury.

Graphical abstract



Highlights

► THg, $\delta^{13}\text{C}$, $\delta^{15}\text{N}$ were determined in 132 marine fish from a tropical coastal region. ► Coastal and demersal species were more contaminated than oceanic and pelagic ones. ► SIA suggest that Hg exposure derived from bottom feeding and/or in estuarine waters. ► Conspecific fish fed on different coastal food webs and Hg increased with fish size. ► Hg levels in fish from Senegal were not of concern for human consumption.

Keywords : Methylmercury sources, Trophic ecology, Feeding habitat, Metal bioaccumulation, Chemical tracers, Isotopic niche

INTRODUCTION

Coastal areas provide many ecosystem services such as fish production, ensuring the supply of marine products to local populations and supporting the economies of coastal countries, particularly through the export (Interwies and Görlitz, 2013). Among the many disturbances to which these ecosystems are subjected, mercury (Hg) contamination is a major environmental concern, due to its deleterious effects on marine organisms and human health (Streets et al., 2011). Hg originates from both natural and anthropogenic processes, including geological weathering, gold mining, combustion and industrial discharges. Although Hg is volatile and can be transported through the atmosphere over long distance away from emission sources, coastal environments are particularly subjected to Hg exposure since they are located at the interface between three main Hg sources: atmosphere, rivers and ocean (Cossa et al., 1996).

After being released into seawater, Hg can be methylated by microorganisms such as sulfate and iron-reducing bacteria or methanogens (Correia and Guimarães, 2017; Gilmour et al., 2013; Paranjape and Hall, 2017; Podar et al., 2015) in coastal and shelf sediments as well as in sub-thermocline oceanic waters, where oxygen concentration is low and organic matter is intensively remineralized (Blum et al., 2013; Cossa et al., 2017; Fitzgerald et al., 2007). Once in the organic methylmercury form (MeHg), this metal is easily taken up by phytoplankton and biomagnifies along trophic chains. Biomagnification (*i.e.* increase in MeHg content with trophic level) is responsible for high Hg concentration in the tissues of top predators such as

tunas and sharks (McKinney et al., 2016). As trophic level is closely correlated with nitrogen stable isotope ratios ($\delta^{15}\text{N}$) (Post, 2002), this marker has been extensively used to explain quantities of Hg in biota (Kiszka et al., 2015; McMeans et al., 2010; Pethybridge et al., 2012). Species with similar trophic levels can, however, display different patterns of Hg accumulation (Bank et al., 2007; Le Croizier et al., 2016) and other factors have been linked to Hg concentrations. First, like other metal elements, Hg strongly binds to sulfhydryl groups of proteins in marine fish leading to a very slow excretion of the bioaccumulated Hg over time, particularly in muscle tissues (Kidd and Batchelar, 2011; Peng et al., 2016). As a consequence, Hg concentration has been extensively shown to increase with fish age, being positively correlated with size or weight (Bosch et al., 2016; Chauvelon et al., 2014; Pethybridge et al., 2010). Second, Hg concentrations in fish tissues are also known to increase with depth of occurrence in the water column, corresponding to concentrations of dissolved organic mercury in seawater and/or proximity to sediments (Chauvelon et al., 2012; Choy et al., 2009). Third, ecosystem characteristics such as human activities, density and composition of phytoplankton communities can influence Hg bioaccumulation and subsequent biomagnification (Chen and Folt, 2005; Chauvelon et al., 2018; Conдини et al., 2017; Heimbürger et al., 2010).

Finally, some studies focusing on Hg sources found differences in Hg concentrations between offshore and nearshore species or between pelagic and benthic species, depending on whether Hg methylation takes place in pelagic deep waters or in coastal sediments (Cresson et al., 2014; Sackett et al., 2015; Senn et al., 2010). Carbon stable isotope values ($\delta^{13}\text{C}$) have commonly been used to discriminate oceanic/pelagic from coastal/demersal habits in marine fish in order to investigate the role of foraging habitat in Hg bioaccumulation (Cresson et al., 2014; Goutte et al., 2015; Pethybridge et al., 2012; Signa et al., 2017). Coastal fish species have

a wide range of prey and habitat use, however, which leads to large intraspecific variability in metal accumulation (Bird et al., 2018; Le Croizier et al., 2016; Sackett et al., 2015). It therefore seems essential to conduct ecotoxicological studies at the individual level to better determine the origin of the accumulated Hg in fish tissues.

In this context, the general goal of the present study was to investigate the trophic origin of mercury exposure to coastal fish from the Canary Current Large Marine Ecosystem (CCLME) in Western Africa. This ecosystem is of particular interest since it is one of the world's major cold-water upwelling currents and ranks first in the world in terms of primary productivity (Messié and Chavez, 2015). It supports one of the largest fisheries among African large marine ecosystems and provides food to local populations but also to foreign countries through the attribution of fishing licenses and export of marine products. This marine ecosystem is prone to metal contamination due to urban effluents and industrial activities (Auger et al., 2015; Diop et al., 2015) and Hg has been recently identified as one of the toxic elements reported at significant concentrations in sediments and marine organisms from this coastal zone (Bodin et al., 2013; Diop and Amara, 2016; Net et al., 2015). Hence, as Hg can cause serious pathologies including neurological impairments (Bosch et al., 2016) and fish consumption represents the major pathway for human exposure to Hg (Driscoll et al., 2013), a proper evaluation of Hg sources and concentrations was needed in fish from this area.

To fulfill this objective, total mercury concentrations were determined in a wide range of fish belonging to 23 different species. Total mercury was used as a proxy of methylmercury, since MeHg generally represents more than 80% of the total mercury found in fish muscle, regardless of their ecological characteristics (Andersen and Depledge, 1997; Bloom, 1992; Magalhães et al., 2007; Storelli et al., 2003). The main origin for Hg contamination was investigated through the reported habitat distribution of the species. This method allowed to

discriminate Hg exposure between oceanic *versus* coastal and pelagic *versus* demersal sources, but did not take into account the individual specialization in terms of foraging habitat. Thus, a second complementary approach was undertaken using stable isotope analyses, particularly $\delta^{13}\text{C}$ that gives information at the individual level but only discriminate between oceanic/pelagic *versus* coastal/demersal source. Finally, the potential health impact associated with fish consumption in the area was discussed in relation to the tolerable Hg intake limit established by the European Union and the Food and Agriculture Organization of the United Nations.

MATERIALS AND METHODS

Sampling

A total of 132 fish belonging to 23 different species were sampled at four different locations along the Senegalese coast in West Africa (from north to south): Saint-Louis, Dakar, Saloum and Casamance (Figure 1). Saint-Louis, Saloum and Casamance are located off estuaries, while Dakar is located along a rocky coast. Samples used in this study were collected during a scientific cruise (doi, 10.17600/14001400) of the BMBF - IRD AWA Project "Ecosystem Approach to the management of fisheries and the marine environment in West African waters" (01DG12073E) in March 2014 aboard the RV Thalassa. Fish were caught with a bottom trawl net, packed in plastic bags and frozen on board at -20°C until further analyses.

Chemical analyses

Once at the laboratory, the fish were weighed (wet weight) and measured (total length) (Table 1). They were then dissected with ceramic tools to avoid metal contamination, and a piece of

dorsal muscle (a standardized cut from just behind the head) was collected. Muscle samples for the analysis of stable isotopes were freeze-dried and ground into a fine homogeneous powder. Approximately 350 µg of powder was then weighed in tin capsules for isotopic analysis. The samples were analyzed by continuous flow on a Thermo Scientific Flash EA 2000 elemental analyzer coupled to a Delta V Plus mass spectrometer at the Pôle de Spectrométrie Océan (Plouzané, France). Results are expressed in standard δ notation based on international standards (Vienna Pee Dee Belemnite for $\delta^{13}\text{C}$ and atmospheric nitrogen for $\delta^{15}\text{N}$) following the equation $\delta^{13}\text{C}$ or $\delta^{15}\text{N} = [(R_{\text{sample}}/R_{\text{standard}}) - 1] \times 10^3$ (in ‰), where R is $^{13}\text{C}/^{12}\text{C}$ or $^{15}\text{N}/^{14}\text{N}$. Samples were not delipidated because muscle generally contains a lower level of lipids than other organs in fish (Njinkoué et al., 2002). However, since lipid content can impact $\delta^{13}\text{C}$, C/N ratios were measured. These values exceeded 3.5 for a few samples (i.e. 18 individuals among 132 in total, see Supplementary Material), so a mathematical normalization was applied to all samples, following the equation $\delta^{13}\text{C}_{\text{normalized}} = \delta^{13}\text{C}_{\text{untreated}} - 3.32 + 0.99 \cdot \text{C/N}$ (Le Croizier et al., 2016; Post et al., 2007). International isotopic standards of known $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ were used: IAEA-600 caffeine, IAEA-CH-6 sucrose, IAEA-N-1 and IAEA-N-2 ammonium sulfate. The certified values of the standards were reproduced within the confidence limits: IAEA-600 (certified values: -27.77 ± 0.04 ‰ for $\delta^{13}\text{C}$ and 1.00 ± 0.20 ‰ for $\delta^{15}\text{N}$; measured value: -27.79 ± 0.10 ‰ for $\delta^{13}\text{C}$ and 1.09 ± 0.06 ‰ for $\delta^{15}\text{N}$), IAEA-CH-6 (certified values: -10.45 ± 0.08 ‰ for $\delta^{13}\text{C}$; measured value: -10.43 ± 0.08 ‰ for $\delta^{13}\text{C}$), IAEA-N-1 (certified values: 0.4 ± 0.2 ‰ for $\delta^{15}\text{N}$; measured value: 0.45 ± 0.10 ‰ for $\delta^{15}\text{N}$) and IAEA-N-2 (certified values: 20.3 ± 0.2 ‰ for $\delta^{15}\text{N}$; measured value: 20.24 ± 0.12 ‰ for $\delta^{15}\text{N}$). A home standard (Thermo acetanilide) was analyzed every 6 samples for experimental precision (based on the standard deviation of the replicates of the internal standard) and indicated an analytical precision of ± 0.12 ‰ for $\delta^{13}\text{C}$ and ± 0.06 ‰ for $\delta^{15}\text{N}$.

Total Hg determination was carried out on an aliquot section (around 50 mg) of the dried muscle by combustion, gold trapping and atomic absorption spectrophotometry detection using a DMA80 analyzer (Milestone, USA). Mercury concentrations in the muscle samples are expressed on a dry weight ($\mu\text{g}\cdot\text{g}^{-1}$ dw), and wet weight ($\mu\text{g}\cdot\text{g}^{-1}$ ww) basis when needed (i.e. for comparison with current maximum acceptable limits for seafood consumption that are expressed on a wet weight basis), applying a moisture correction factor determined for each individual sample by using the mass ratio before and after freeze drying. The accuracy and reproducibility of the method were established using two freeze-dried certified biological materials; a tuna fish flesh homogenate reference material (IAEA 436, IRMM) and a Lobster Hepatopancreas reference material (TORT3, NRCC). The certified values for IAEA 436 ($4.19 \pm 0.36 \mu\text{g}\cdot\text{g}^{-1}$ dw) were reproduced (measured value: $4.28 \pm 0.14 \mu\text{g}\cdot\text{g}^{-1}$ dw) within the confidence limits. The certified values for TORT3 ($0.292 \pm 0.022 \mu\text{g}\cdot\text{g}^{-1}$ dw) were also reproduced (measured value: $0.280 \pm 0.005 \mu\text{g}\cdot\text{g}^{-1}$ dw) within the confidence limits. Repeatability varied from 1.9 to 6.7% depending on the concentration of the sample and its heterogeneity. The detection limit was $0.005 \mu\text{g}\cdot\text{g}^{-1}$ dw.

Ecological data of species

The ecological distribution of each species was defined on both the “horizontal” axis (i.e. the offshore-inshore gradient) and the “vertical” axis (i.e. water column occupation). On the horizontal axis, “Offshore” includes bathydemersal and oceanodromous species while “Inshore” includes coastal, brackish and estuarine species. On the vertical axis, the different fish species were grouped according to their known habitat (i.e. pelagic, benthopelagic and demersal). For both axes, species were classified following general published literature for most species in the area (Caverivière, 1993; Diouf, 1996; Froese and Pauly, 2018; Séret, 2011)

and specific studies for fish species in particular: *Boops boops* (Derbal and Hichem Kara, 2008), *Caranx rhonchus* (Boëly et al., 1973; Sley et al., 2008), *Chloroscombrus chrysurus* and *Galeoides decadactylus* (Faye et al., 2011), *Diplodus bellottii* (Horta et al., 2004), *Scomber japonicus* (Castro, 1993), *Sphyraena guachancho* (Akadje et al., 2013), *Trachurus trecae* (Boëly et al., 1973).

Statistical analyses

All data tested statistically were first checked for normality (Shapiro–Wilks tests) (Shapiro and Wilk, 1965) and homogeneity of variances (Bartlett tests) (Bartlett, 1937). When these conditions were met, raw data (or log-transformed data, if necessary) were used and one-way ANOVAs, followed by Tukey's HSD tests, were performed to test for differences between groups (i.e. Hg variation between sites or groups of species based on their ecological distribution, stable isotope variations between groups of individuals based on their isotopic values). Otherwise, non-parametric analogues were used, i.e. Kruskal-Wallis tests (KW), followed by Conover-Iman multiple comparison tests with Bonferroni's adjustment when more than two groups were compared (Conover and Iman, 1979), to test for differences in $\delta^{15}\text{N}$ values between sites or Hg concentration between groups of individuals based on their $\delta^{13}\text{C}$ values.

To assess the relationships between individual Hg bioaccumulation and feeding habitat and trophic level, Hg concentrations were modeled using linear regressions with isotopic values ($\delta^{13}\text{C}$, $\delta^{15}\text{N}$) as explanatory variables. To evaluate the intraspecific variability in foraging habitat and its effect on mercury concentration, individual fish were classified in two groups based on $\delta^{13}\text{C}$ values, using a K-means cluster analysis (Hartigan and Wong, 1979). This clustering method delineates groups in the data by minimizing the sum of the within-groups

sums of squared-distances, based on Euclidean distance. The number of groups for the partition was defined using the Caliński-Harabasz criterion (Caliński and Harabasz, 1974). The influence of the horizontal and vertical habitat at the species level as well as individual characteristics (i.e. $\delta^{13}\text{C}$, $\delta^{15}\text{N}$, length and weight) on muscle Hg concentrations was tested using linear mixed models (LMMs) with species and/or sampling site as a random effect on the whole data set ($n=132$), using the R package “lme4” (Bates et al., 2014). The models were constructed with a gamma distribution and a log link function based on residual analysis (Zuur et al., 2009). To identify the most important factors explaining Hg concentrations, the different models were ranked based on Akaike's Information Criteria adjusted for small sample sizes (AICc) and compared using ΔAICc and Akaike weights (w) using the R package “wiqid” (Meredith, 2017). Models with AICc values differing by less than two have a similar level of support in the data (Burnham and Anderson, 2002). To assess the explanative power of these models, marginal R^2 was obtained following Nakagawa Shinichi et al., 2012 using the R package “r2glmm” (Jaeger, 2017). All of the statistical analyses were performed using the open source software R (version 3.4.3, R Core Team, 2017).

RESULTS AND DISCUSSION

Hg exposure under estuarine and upwelling influence

Concerning Hg spatial variation, fish sampled off the Saloum estuary (all species combined: $n = 44$) displayed higher Hg concentrations than those from Saint-Louis (all species combined: $n = 24$) or Dakar (all species combined: $n = 29$) (ANOVA followed by Tukey's HSD test; $p < 0.01$) (Figure 2). This variation may be first related to the sampling of different species between

sites. Indeed, the different sites showed very few common species (only *Caranx rhonchus* in Dakar and Casamance, *Pseudupeneus prayensis* in Dakar and Saloum, *Trachurus trecae* in Dakar and Saloum; Table 1). In addition, species-specific characteristics such as ingestion rate, assimilation efficiency, growth rate, organ distribution, subcellular handling and excretion rate, constitute physiological factors that influence Hg concentration in muscle tissue (Peng et al., 2016; Wang and Wang, 2018, 2015; Wang and Wong, 2003). This diversity of species may thus have participated to the Hg variation between locations. Other possible explanations relate to hydrodynamic processes. First, sources of Hg in coastal waters include inputs from upwellings, rivers, groundwater, atmospheric deposition and waste water point sources (Cossa et al., 2017). The river-watershed contributes greatly to Hg inputs to coastal waters (Balcom et al., 2015; Choe and Gill, 2003) and is estimated to be the highest total mercury source in ocean margins, mainly in association with organic particles (Cossa et al., 1996). The higher Hg contamination in fish from the Saloum station may thus be the result of their feeding in the estuarine zone that is influenced by riverine inputs, as suggested by the high $\delta^{15}\text{N}$ baseline. Moreover, the Saloum site is located near the mouth of the Gambia River, that is impacted by the use of Hg for gold amalgamation in small artisanal mining (Niane et al., 2015). It is thus hypothesized that the Gambia River estuary could be a source of anthropogenic Hg for fish feeding in this coastal area, including those sampled at the Saloum site. Second, Hg bioavailability depends largely on water column stratification as MeHg production peaks at the thermohalocline where density gradient is maximum and allows organic matter accumulation as well as the associated bacterial activity (Cossa et al., 2012). In areas under riverine influence such as estuaries, freshwater inputs might generate a thermohalocline near the surface, together with organic matter flocs that could enhance MeHg formation. This higher Hg bioavailability would thus be partially responsible for the higher Hg concentrations

in fish from the Saloum region, that supposedly feed in the Saloum and Gambia river estuaries. Third, upwellings constitute a major Hg source from the ocean to the neritic zone. Indeed, MeHg is mainly formed near the shelf edge, where organic carbon degradation is intense. Upwelled waters from this depth can account for more than half of MeHg accumulated by fish in ocean margins (Cossa et al., 2017, 1996). In Senegal, upwelling occurs mainly in winter and is stronger from February to April. It produces a coastal jet confined nearshore that flows southward alongside the coast (Auger et al., 2015). Recently, coastal waters of southern Senegal were shown to originate from an upwelling area in the north (Ndoye et al., 2017). Because the fish were collected during the active upwelling season when the coastal current is the strongest, transport of MeHg by upwelled waters might contribute to the greater Hg concentrations at the most southern locations (*i.e.* Saloum and Casamance; Figure 2C). This pattern was also previously observed for other metals such as cadmium (Le Croizier et al., 2016).

Influence of foraging habitat on Hg accumulation at the species level

The different fish species were classified in two distinct groups according to horizontal distribution (*i.e.* the offshore-inshore gradient). “Offshore” includes bathydemersal and oceanodromous species while “Inshore” includes coastal, brackish and estuarine species. Hg concentration in the muscle was higher in inshore fish than in offshore ones (ANOVA; $p < 0.001$) (Figure 3A), showing that proximity to the coast implies higher Hg exposure. In the present case study, the higher contamination of inshore species can first be explained by the estuarine influence that may bring substantial inorganic particulate Hg to the coastal marine environment where MeHg production occurs (Figure 4), as observed in other areas like the northern Gulf of Mexico and the Mediterranean Sea (Cossa and Coquery, 2005; Senn et al.,

2010). Then, this contamination pattern can also be explained by the higher methylation rate observed in coastal sediments subjected to significant inputs of organic material (OM) from catchment (Lambertsson and Nilsson, 2006; Merritt and Amirbahman, 2009). In a previous study, higher concentrations of Hg were found in coastal species from Senegal during the wet season (Diop and Amara, 2016), that increases runoff from land and rivers, compared to the dry season where the upwelling intensity is higher (Ndoye, 2016). This observation strengthens the hypothesis of the input of Hg, and/or OM that enhances methylation, to the coastal environment by rivers rather than upwelling.

In order to investigate the influence of vertical distribution (*i.e.* water column occupation), the different fish species were grouped according to their known habitat (*i.e.* pelagic, benthopelagic and demersal). Hg concentration increased from the surface to the bottom, demersal species exhibiting higher Hg concentrations relative to pelagic fish, while benthopelagic species showed intermediate Hg values (ANOVA followed by Tukey's HSD test; $p < 0.01$) (Figure 3B). Trophic level is one of the main contributors to Hg variation when different fish species are compared (Kiszka et al., 2015; McMeans et al., 2010; Pethybridge et al., 2012). However, in the present study, TL derived from the literature (Table 1) did not significantly differ between groups: TL pelagic = 3.8; TL benthopelagic = 3.7; TL demersal = 3.5 (ANOVA; $p > 0.5$). On the other hand, microbial MeHg production and subsequent mobilization from sediments is known to be a major source of Hg accumulation for coastal marine organisms including edible fish (Hammerschmidt and Fitzgerald, 2006), implying a higher Hg bioavailability for benthic organisms (Fitzgerald et al., 2007). Moreover, photodemethylation by ultraviolet radiation acts to remove MeHg from surface water and reduces Hg bioavailability for pelagic species that feed near the surface (Lehnherr et al., 2011) (Figure 4). In the absence of trophic level influence, these two mechanisms may explain the Hg gradient

found along the water column distribution of the species sampled in the present study. These results are in accordance with a previous study on a wide range of species ($n = 120$) from the Bay of Biscay (North-East Atlantic), in which pelagic organisms showed lower Hg concentrations than benthic and benthopelagic ones (Chouvelon et al., 2012). Moreover, in this previous study, water column distribution was the factor that contributed the most to explain the variability in muscle Hg concentrations, highlighting the importance of this parameter in understanding marine fish species exposure to Hg.

Influence of foraging habitat on Hg accumulation at the individual level

Hg accumulation was previously shown to be linked to species-specific characteristics in the horizontal and vertical distribution (Figure 3). Some fish species can however display a high intraspecific variability in terms of diet composition and foraging habitat, which can be reflected in metal accumulation (Das et al., 2000; Le Croizier et al., 2016). Intraspecific variability in horizontal distribution can lead to differences in Hg accumulation in the dusky grouper *Epinephelus marginatus* from the Southern Brazilian coast, where individuals sampled in an estuarine zone were more contaminated than those from an offshore bank due to anthropogenic Hg discharges into the drainage basin of the estuary (Concini et al., 2017). Vertical distribution can also contribute to variation in Hg among individuals of the same species. It can occur through an ontogenetic shift in foraging habitat, as for the bluefish *Pomatomus saltatrix*, that presents higher Hg accumulation during its early life stage when feeding on benthic preys compared to feeding on pelagic preys in later years (Szczebak and Taylor, 2011). It can also result from depth of occurrence, as shown for the European hake *Merluccius merluccius* in the northwestern Mediterranean, where the Hg concentrations in muscle followed the MeHg distribution in the water column (Cossa et al., 2012). These authors

concluded that Hg bioavailability was higher for hake inhabiting the continental slope than the continental shelf. Working at the species level might not, therefore, be the most relevant method to study metal bioaccumulation. Following this postulate, individual stable isotope values were used to investigate the role of trophic position and foraging habitat in Hg contamination, through $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$, respectively.

Numerous studies found that Hg concentrations in marine fish are mainly explained by $\delta^{15}\text{N}$, that is a proxy of trophic level (Sackett et al., 2015; Senn et al., 2010). MeHg is biomagnified (*i.e.* to increase with trophic level) within marine food webs (McMeans et al., 2010; Signa et al., 2017), making $\delta^{15}\text{N}$ a primary indicator for Hg accumulation. In the present study, $\delta^{15}\text{N}$ varied among locations (see Supplementary Material, Figure S1; KW followed by Conover-Iman test, $p < 0.001$) suggesting differences in $\delta^{15}\text{N}$ baselines between ecosystems, so that the link between $\delta^{15}\text{N}$ and Hg was only investigated at the local level. Hg concentration in fish muscle was found to be positively correlated with $\delta^{15}\text{N}$ in Saint-Louis (Figure 5A, $p < 0.001$) and Dakar (Figure 5B, $p < 0.001$) while no correlation was observed in Saloum (Figure 5C) or Casamance (Figure 5D). In the first two sites, Hg accumulation followed trophic position, as frequently observed in fish communities. In Saloum and Casamance, the lack of observed biomagnification can be attributed to the limited $\delta^{15}\text{N}$ gradient within these fish communities compared to those from northern sites. Indeed, a previous study on bioaccumulation of other metal elements found three $\delta^{15}\text{N}$ -based trophic levels in fish from Dakar, while those from Casamance displayed only two trophic levels (Le Croizier et al., 2016). These findings suggest that $\delta^{15}\text{N}$ would not be relevant to explain Hg accumulation variability among fish that were too close in trophic level. Furthermore, the northern ecosystems (*i.e.* Saint-Louis and Dakar) may support only one main nitrogen source of marine origin, while southern sites (*i.e.* Saloum and Casamance) may display two main nitrogen sources, one of marine origin and the other

one of estuarine or upwelling origin. Multiple nitrogen sources may thus obscure the classical correlation between Hg and $\delta^{15}\text{N}$ in locations influenced by estuarine inputs. Because both vertical and horizontal distribution is reflected in $\delta^{13}\text{C}$, this marker can give confusing information concerning water column distribution and proximity to the coast. However, it allows to globally discriminate pelagic/offshore from demersal/inshore habits of the fish. $\delta^{13}\text{C}$ did not vary significantly with location in our study (ANOVA, $p > 0.05$), but $\delta^{13}\text{C}$ values in fish from the Saloum station displayed a particular pattern. While low $\delta^{13}\text{C}$ values are generally attributed to pelagic/offshore foraging habitats in marine food webs, organisms from areas exposed to terrestrial inputs like estuaries can be depleted in ^{13}C due to the lower $\delta^{13}\text{C}$ of terrestrial particular organic matter (POM) compared with marine POM (Cresson et al., 2014; Darnaude, 2005). It can therefore lead to the observation of low $\delta^{13}\text{C}$ in fish known to inhabit coastal benthic habitats. In fish sampled off the Saloum estuary, the three species displaying the lowest $\delta^{13}\text{C}$ values were *Chelidonichthys gabonensis*, *Pseudupeneus prayensis* and *Dentex canariensis* ($\delta^{13}\text{C} = -16.96 \pm 0.35 \text{ ‰}$, $-16.93 \pm 0.55 \text{ ‰}$ and $-16.54 \pm 0.37 \text{ ‰}$, respectively), all described as coastal and demersal species (Table 1). Furthermore, the Saloum estuary is subjected to terrestrial carbon sources, such as mangrove leaves, which lead to really low $\delta^{13}\text{C}$ values for both particulate and sediment organic matter (i.e. around -25 ‰) (Faye et al., 2011; Gning et al., 2010). These findings thus suggest that the low $\delta^{13}\text{C}$ values observed in these three particular coastal/demersal species reflected the use of the estuarine habitat of Saloum and Gambia River rather than offshore/pelagic signatures (Figure 6A). In fish from this site, Hg concentrations increased for both the lowest $\delta^{13}\text{C}$ (terrestrial influence) and the highest $\delta^{13}\text{C}$ (inshore/demersal) ($p < 0.001$), showing the higher Hg accumulation of individuals with estuarine foraging. Apart from the local trend off the Saloum estuary, as the $\delta^{13}\text{C}$ pattern was similar between sites, the link between Hg and $\delta^{13}\text{C}$ was investigated on all fish from the

remaining locations together. A positive correlation was also found between individual Hg concentrations and $\delta^{13}\text{C}$ values ($p < 0.001$; Figure 6B), strengthening the hypothesis of the involvement of the coastal environment and sediments in individual Hg exposure. This link between Hg concentration and inshore/demersal affinity inferred by $\delta^{13}\text{C}$ has also been found in fish from other marine environments (Bank et al., 2007; Goutte et al., 2015).

Intraspecific variability determines Hg accumulation

To investigate intraspecific variability in terms of foraging habitat, individual fish were grouped according to their carbon source, *i.e.* group 1: lowest $\delta^{13}\text{C}$, suggesting offshore/pelagic habits; and group 2: highest $\delta^{13}\text{C}$, indicating inshore/demersal affinity (Figure 6C). Individuals from the three species from Saloum discussed above were excluded from the analysis since their terrestrial $\delta^{13}\text{C}$ signature would not assign them to a group reflecting their coastal/benthic described ecology. The k-means clustering in two groups explained 71.6% of the observed variability in $\delta^{13}\text{C}$ values, demonstrating the relevance of this partitioning. Half of the species included in the clustering were split between the two groups (*i.e.* Group: “1 & 2” able 1), highlighting the great discrepancy in habitat occupation of individuals from a same species. Despite this high intraspecific variability, Hg bioaccumulation followed the same pattern as found at the species level, with offshore/pelagic fish being less contaminated than inshore/demersal ones (KW, $p < 0.001$) (Figure 6 and Figure S2).

The trophic niche (*i.e.* combination of trophic level and feeding habitat) can be inferred through the use of both $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ (Sardenne et al., 2016) and has been shown to determine metal exposure for marine fish (Das et al., 2000; Le Croizier et al., 2016). The trophic niche influence on mercury contamination was here investigated on a single species with a sufficient number of individuals ($n=10$): the Atlantic bumper *Chloroscombrus chrysurus*.

Conspecific fish showed two distinct trophic niches based on a k-means clustering, characterized by different $\delta^{15}\text{N}$ (ANOVA, $p < 0.05$) and $\delta^{13}\text{C}$ (ANOVA, $p < 0.01$) (Figure 7). Thus, despite the low selectivity of prey in the feeding behavior of *C. chrysurus* that feeds mainly on pelagic particles (Faye et al., 2011), this result revealed again the large diversity in the trophic sources exploited by coastal consumers (Bird et al., 2018). Individuals belonging to different isotopic niches were also characterized by contrasted Hg concentrations (ANOVA, $p < 0.001$), the fish feeding on a more coastal food chain and at a higher trophic level being the more contaminated. This clear influence of the trophic niche in Hg exposure may be the result of mercury biomagnification even for forage fish such as *C. chrysurus* as well as the higher methylmercury production in estuarine waters compared to shelf waters (Figure 4).

Prediction of Hg concentrations and impact on consumer health

In contrast to other toxic metal elements like cadmium, that are poorly assimilated from food (Le Croizier et al., 2018), bioaccessibility of MeHg to humans from fish consumption can be high depending on the species consumed (Bradley et al., 2017) and can impair human health, mainly through neurological damage (Bosch et al., 2016). Fish consumption is also known to provide health benefits through the intake of essential nutrients such as omega-3 fatty acids that contribute to the prevention of cerebrovascular diseases (Chowdhury et al., 2012; Rheinberger and Hammitt, 2012). However, as fish consumption constitutes the main pathway for Hg exposure (Driscoll et al., 2013) and fish exhibit high inter and intraspecific variability in Hg concentrations, attention should be focused on determining which fish are safe or should be avoided for human consumption. In this study, the low overlap of species between sites makes it difficult to draw conclusion regarding the spatial pattern of Hg contamination along the coast of Senegal. In this context, univariate models were used to

predict Hg concentrations in individual fish taking into account species and site variability. The model with length as a predictor of Hg concentration was the most supported ($\Delta\text{AICc} = 0.00$, $w_i = 1.00$; Table S2, Supplementary Material) and explained 25% of Hg variation in the fish flesh. Similarly, weight explained 15% of the observed Hg variability. These results are in accordance with the well-described bioaccumulation of Hg over time due to its very slow excretion from muscle tissue (Peng et al., 2016). Moreover, taking into account the spatial variability, the habitat of the species accounted for a significant part of Hg variation (i.e. 26% and 16%, for horizontal and vertical habitat respectively). In view of the above conclusions, this highlights the crucial role of the habitat in trophic Hg exposure to marine fish. Finally, models with length as a predictor of Hg were used without species and/or sampling site as random effects. The model with species and sampling site as random effects was again the most supported ($\Delta\text{AICc} = 0.00$, $w_i = 1.00$, $R^2_m = 25\%$; Table S2 Supplementary material). However, the model with only species as a random effect lead to a higher explained variability (i.e. 30%) that demonstrates the importance of considering the species rather than the site (explaining only 4% of Hg variation) in predicting mercury concentrations. To sum up, these results indicate an increasing risk of mercury exposure to humans through the consumption of larger fish (within a same species) with demersal and/or coastal habits.

In the present study, all individuals presented a muscular Hg concentration below the current maximum acceptable limit for seafood consumption (see Supporting Information) set by both the European Union and the Food and Agriculture Organization of the United Nations, that is $0.5 \mu\text{g}\cdot\text{g}^{-1}$ ww for the species investigated here (EU, 2006; FAO, 2003). However, considering the Tolerable Weekly Intake (TWI) set by the European Food Safety Authority (EFSA), that is $1.3 \mu\text{g kg}^{-1}$ (bw) for MeHg (EFSA, 2012), and assuming that nearly all the bioaccumulated Hg is in its organic form, consumption of some species from southern locations such as Saloum

and Casamance should be limited. Indeed, the quantity of muscle tissue consumed per day by a 70-kg person should not exceed 130 g for *Dicologlossa hexophthalma* (Table S3, Supporting material), 118 g for *Pseudupeneus prayensis*, 108 g for *Plectorhinchus mediterraneus* and 93 g for *Caranx rhonchus*, that is among the most exploited species in the Canary Current Large Marine Ecosystem (Sambe et al., 2016). As marine products represent the main source of protein intake for the population of Senegal, especially in southern areas (FAO, 2006), overconsumption of some coastal demersal species could be of concern regarding human exposure to Hg.

Limitations of the study

The present study shows a number of limitations that have to be considered in interpreting the results.

First, a piece of dorsal muscle was used to analyze both stable isotopes and Hg concentration. Thus, the obtained results might not reflect the whole fish fillet characteristics, particularly for fish species with high fat content. Indeed, lipid content can vary between the dorsal and lateral region of muscle tissues, influencing both $\delta^{13}\text{C}$ values and Hg concentrations (Balshaw et al., 2008; Cherel et al., 2011). In this context, a mathematical correction was applied to $\delta^{13}\text{C}$ values to accurately account for the effect of lipids (Post et al., 2007; Skinner et al., 2016). Moreover, no variation in Hg accumulation was detected within the white muscle, that constitutes the largest part of muscle tissue in marine fish (Bosch et al., 2016; O'Bryhim et al., 2017). Thus, the protocol used in this study (i.e. using a dorsal muscle portion) has been extensively applied in recent risk assessment studies (Araújo and Cedeño-Macias, 2016; Bodin et al., 2017; Nicklisch et al., 2017). Second, the risk assessment of mercury intake due to fish consumption is based on 5 individuals for each species (Table 1) and may not be representative of the Hg

concentrations throughout the region. However, to date only few studies have examined Hg concentrations in selected marine fish species from Senegal (Diop et al., 2017; Diop and Amara, 2016), while the present study offers a broader assessment of species that may be at risk for human consumption and allows suggesting that future studies should focus on coastal and demersal fish species.

Finally, interpretation of carbon isotope data to determine feeding habitat for marine fish in coastal environments can be difficult, since carbon sources originating from opposite habitats can show similar isotopic signature (e.g. low $\delta^{13}\text{C}$ values for both oceanic and terrestrial materials), especially in ecosystems under estuarine influence (Cresson et al., 2014; Darnaude, 2005; Senn et al., 2010). In the present study, despite low $\delta^{13}\text{C}$ values observed in three particular coastal species from the Saloum site (red points, Fig. 6A), that is known to receive ^{13}C -depleted terrestrial inputs (Faye et al., 2011; Gning et al., 2010), $\delta^{13}\text{C}$ of "inshore" species was significantly higher than $\delta^{13}\text{C}$ of "offshore" species, all sites combined (see Supplementary material, Figure S3; ANOVA, $p < 0.001$). This result confirms that apart from the particular case of few fish from Saloum, low $\delta^{13}\text{C}$ values are actually characteristic of the influence of offshore marine habitat rather than estuarine habitat throughout the coastal zone. However, this highlights the need to verify the significance of $\delta^{13}\text{C}$ values by comparing with existing ecological data of the studied species (e.g. from stomach contents or fishery data), especially when there are multiple carbon sources such as in complex coastal ecosystems. Here, the same contamination pattern was observed with two different methods (i.e. species-specific ecological data and individual $\delta^{13}\text{C}$ signatures), strengthening the hypothesis that Hg exposure mainly occurs through the sediments of the benthic/coastal habitat, like previously observed in many other coastal ecosystems (Balogh et al., 2015; Gehrke et al., 2011; Sackett et al., 2015)

CONCLUSION

Results from this study confirmed the influence of diet and habitat use in metal transfer to fish in marine environments (Le Croizier et al., 2016; Signa et al., 2017). They further highlight the great diversity in the trophic sources exploited by coastal consumers and the resulting need to consider individual foraging behavior in ecotoxicological studies. Moreover, this study revealed that the mercury previously observed in marine organisms from Senegal (Bodin et al., 2013; Diop and Amara, 2016; Net et al., 2015) is likely to originate from estuarine and sedimentary sources. Recently, Hg isotopic signatures have been used to characterize the complex biogeochemical cycle of this contaminant. Hg isotopes allowed to identify primary food sources of the accumulated-Hg in marine fish (Sackett et al., 2017, 2015) and to determine the habitats leading to Hg exposure, by discriminating between estuarine and oceanic affinities (Li et al., 2016, Senn et al., 2010) or benthic versus pelagic habits (Gehrke et al., 2011). These innovative analytical techniques, as well as the carbon stable isotope analysis of methylmercury (Masbou et al., 2015), are thus promising tools to get better insights in the origin, fate and ecodynamics of mercury in the marine food webs of this region with high fishing potential.

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Figure 1: Sampling sites along the Senegalese coast (West Africa). Saint-Louis, Saloum and Casamance are located off estuaries, while Dakar is located along a rocky coast.

Figure 2: Boxplots of the mercury concentration (Hg in $\mu\text{g}\cdot\text{g}^{-1}$ dry weight) in muscles of fish from different sites (Saint-Louis, $n = 24$; Dakar, $n = 29$; Saloum, $n = 44$; Casamance, $n = 35$). The box length represents the interquartile range, the bar length represents the range, and the horizontal line is the median value. Different letters indicate a significant difference between sites (ANOVA followed by Tukey's HSD test; $p < 0.01$).

Figure 3: Boxplots of the mercury (Hg) concentration ($\mu\text{g}\cdot\text{g}^{-1}$ dry weight) in muscle of fish species according to their proximity to the coast (A: Offshore, $n = 34$; Inshore, $n = 98$) and according to water column distribution (B: Pelagic, $n = 20$; Benthopelagic, $n = 34$; Demersal, $n = 78$). The box length represents the interquartile range, the bar length represents the range, and the horizontal line is the median value. Different letters indicate a significant difference between groups (ANOVA followed by Tukey's HSD test; A: $p < 0.001$, B: $p < 0.01$).

Figure 4: Schematic representation of major processes supposed to be involved in Hg accumulation in marine fish species from the Senegalese coast. River inputs carry organic matter (OM) and particulate inorganic Hg to the coastal environment. After particles sink, Hg is methylated by bacterial activity, becoming highly bioavailable to fish foraging near the sediments (*i.e.* demersal species and, to a lesser extent, benthopelagic species). Methylmercury (MeHg) is rapidly accumulated by the biota, and concentrations decrease away from the coast. MeHg is also supposedly transported nearshore by a coastal current resulting from the upwelling activity. Near the surface, photochemical degradation of MeHg leads to formation of inorganic Hg, which is poorly assimilated by pelagic fish.

Figure 5: Mercury (Hg) concentration ($\mu\text{g}\cdot\text{g}^{-1}$ dry weight) in muscles of individual fish vs. muscle $\delta^{15}\text{N}$ (‰) signature from the different study sites (A: Saint-Louis; B: Dakar; C: Saloum; D: Casamance). Data fit a linear curve in A and an exponential curve in B, no correlation was found in C and D.

Figure 6: Mercury (Hg) concentration ($\mu\text{g}\cdot\text{g}^{-1}$ dry weight) in muscles of individual fish vs. muscle $\delta^{13}\text{C}$ (‰) signature from "Saloum" station (A), from the other sites, *i.e.* Saint-Louis, Dakar and Casamance (B), and from all sites combined except the red points from Saloum (C). Data fit a polynomial curve in A and a linear curve in B. The former is due to the low $\delta^{13}\text{C}$ of

the red-colored individuals despite their well-known coastal ecology, suggesting the assimilation of terrestrial rather than oceanic carbon. In C the two groups determined through k-means clustering on individual $\delta^{13}\text{C}$ values explained 71.6 % of the observed variability and significantly differed in $\delta^{13}\text{C}$ values (C: Group 1, n = 63; Group 2, n = 69; ANOVA, p < 0.001).

Figure 7: Individual stable isotope values ($\delta^{15}\text{N}$ and $\delta^{13}\text{C}$) in the Atlantic bumper *Chloroscombrus chrysurus* sampled in Casamance during this study. Colors indicate two trophic niches determined by a K-means cluster analysis. The two groups are characterized by different $\delta^{15}\text{N}$ (ANOVA, p < 0.05) and $\delta^{13}\text{C}$ (ANOVA, p < 0.01) and muscle mercury concentrations (ANOVA, p < 0.001).

Location	Species	n	T L	Length (cm)	Weight (g)	Shore	Habitat	Fishing depth (m)	$\delta^{15}\text{N}$ (‰)	$\delta^{13}\text{C}$ (‰)	Group	Hg ($\mu\text{g}\cdot\text{g}^{-1}$ dw)
Saint Louis	<i>Brotula barbata</i>	4	3.8	33.6 ± 2.8	287.8 ± 73.0	Off	D	33-37	11.14 ± 0.12	-16.99 ± 0.10	1	0.09 ± 0.01
	<i>Merluccius senegalensis</i>	5	4.5	36.4 ± 1.7	350.8 ± 55.3	Off	D	33-37	11.63 ± 0.59	-16.33 ± 0.36	1 & 2	0.22 ± 0.11
	<i>Pterothrissus belloci</i>	5	5.3	25.6 ± 2.2	115.0 ± 32.3	Off	D	33-37	11.14 ± 0.13	-16.53 ± 0.08	1	0.08 ± 0.02
	<i>Trichiurus lepturus</i>	5	4.5	70.7 ± 4.7	223.3 ± 48.1	Off	B	33-37	11.41 ± 0.31	-16.57 ± 0.21	1	0.06 ± 0.01
	<i>Zeus faber</i>	5	4.5	35.2 ± 5.1	661.1 ± 268.0	Off	B	33-37	12.03 ± 0.51	-15.20 ± 0.55	1 & 2	0.18 ± 0.04
	Dakar	<i>Boops boops</i>	5	3.8	18.7 ± 1.6	63.8 ± 22.2	In	D	66-81	10.12 ± 0.23	-15.77 ± 0.62	1 & 2
<i>Caranx rhonchus</i>		4	3.6	19.9 ± 1.4	91.3 ± 13.1	In	B	66-81	11.53 ± 0.17	-15.42 ± 0.43	2	0.05 ± 0.01
<i>Diplodus bellottii</i>		5	3.5	17.0 ± 0.4	77.9 ± 8.7	In	D	22-23	12.34 ± 0.35	-15.08 ± 0.42	2	0.34 ± 0.09
<i>Pseudupeneus prayensis</i>		5	3.2	21.1 ± 1.8	120.6 ± 32.5	In	D	22-23	11.86 ± 0.36	-15.70 ± 0.44	1 & 2	0.21 ± 0.07
<i>Scomber japonicus</i>		5	3.4	16.2 ± 0.6	32.9 ± 5.2	Off	P	66-81	8.01 ± 0.28	-16.70 ± 0.34	1	0.04 ± 0.01
<i>Trachurus trecae</i>		5	3.5	21.8 ± 0.4	99.6 ± 4.8	In	B	22-23	10.92 ± 0.18	-16.42 ± 0.26	1	0.11 ± 0.02
Saloum	<i>Chaetodon hoefleri</i>	5	3.5	17.4 ± 0.8	147.6 ± 17.2	In	D	49-81	15.16 ± 1.36	-15.35 ± 0.71	1 & 2	0.14 ± 0.08
	<i>Chelidonichthys gabonensis</i>	5	3.6	17.9 ± 6.1	123.8 ± 54.0	In	D	49-81	10.88 ± 0.23	-16.96 ± 0.35	-	0.37 ± 0.14
	<i>Dentex canariensis</i>	5	3.6	25.9 ± 1.4	259.8 ± 48.7	In	D	49-81	12.32 ± 0.39	-16.54 ± 0.37	1	0.24 ± 0.14
	<i>Dicologlossa hexophthalma</i>	5	3.4	19.0 ± 1.0	87.2 ± 11.2	In	D	49-81	11.21 ± 0.14	-14.82 ± 0.14	2	0.42 ± 0.28
	<i>Plectorhinchus mediterraneus</i>	5	3.5	30.0 ± 1.2	376.2 ± 53.2	In	D	49-81	12.65 ± 0.22	-15.30 ± 0.09	2	0.47 ± 0.17
	<i>Pseudupeneus prayensis</i>	5	3.2	21.4 ± 1.2	121.0 ± 15.0	In	D	49-81	11.90 ± 0.30	-16.93 ± 0.55	1	0.43 ± 0.36
	<i>Sphoeroides spengleri</i>	4	3.2	12.6 ± 0.4	34.5 ± 2.1	In	D	49-81	11.34 ± 1.16	-15.57 ± 0.45	1 & 2	0.14 ± 0.06
	<i>Trachurus trecae</i>	5	3.5	24.0 ± 1.7	137.6 ± 27.9	In	B	49-81	10.94 ± 0.22	-15.95 ± 0.60	1 & 2	0.09 ± 0.03
	<i>Umbrina canariensis</i>	5	3.4	23.0 ± 1.0	147.8 ± 16.3	Off	D	49-81	11.48 ± 0.27	-15.99 ± 0.21	1 & 2	0.09 ± 0.05
Casamance	<i>Brachydeuterus auritus</i>	5	3.8	20.8 ± 0.4	127.2 ± 5.0	In	B	14-15	13.69 ± 0.30	-15.31 ± 0.15	2	0.35 ± 0.09
	<i>Caranx rhonchus</i>	5	3.6	33.0 ± 3.0	334.4 ± 68.8	In	B	14-15	13.80 ± 0.83	-15.17 ± 0.27	2	0.58 ± 0.34
	<i>Chloroscombrus chrysurus</i>	10	3.5	20.1 ± 0.9	72.6 ± 8.0	In	P	17-18	13.51 ± 0.45	-16.31 ± 0.61	1 & 2	0.1 ± 0.03

<i>Galeoides</i>	5	3.	17.6 ±	63.0 ±	In	D	14-	13.51 ±	-15.65 ±	1 &	0.08 ± 0
		6	1.3	16.1			15	0.20	0.23	2	
<i>decadactylus</i>	5	4.	24.4 ±	170.0 ±	In	D	14-	13.27 ±	-16.02 ±	1 &	0.23 ±
		1	1.3	22.7			15	0.87	0.48	2	
<i>Selene dorsalis</i>	5	4.	34.4 ±	197.8 ±	In	P	14-	13.78 ±	-15.77 ±	1 &	0.24 ±
		4	3.4	53.6			15	0.84	0.32	2	

Table 1: Summary (mean ± standard deviation) of the different biological parameters measured on fish from different sampling sites. “TL” indicates trophic level based on Froese and Pauly, 2018. Based on the literature (see Materials and methods section), “Habitat” indicates water column distribution (D: Demersal, B: Benthodemersal, P: Pelagic), and “Shore” defines horizontal distribution of the species (“Off” includes bathydemersal and oceanodromous species while “In” includes coastal, brackish and estuarine species). “Group” is relative to k-means clustering of individuals based on $\delta^{13}\text{C}$ values: “1” includes pelagic/offshore fish while “2” includes demersal/inshore individuals.

HIGHLIGHTS

- THg, $\delta^{13}\text{C}$, $\delta^{15}\text{N}$ were determined in 132 marine fish from a tropical coastal region
- Coastal and demersal species were more contaminated than oceanic and pelagic ones
- SIA suggest that Hg exposure derived from bottom feeding and/or in estuarine waters
- Conspecific fish fed on different coastal food webs and Hg increased with fish size
- Hg levels in fish from Senegal were not of concern for human consumption



Figure 1

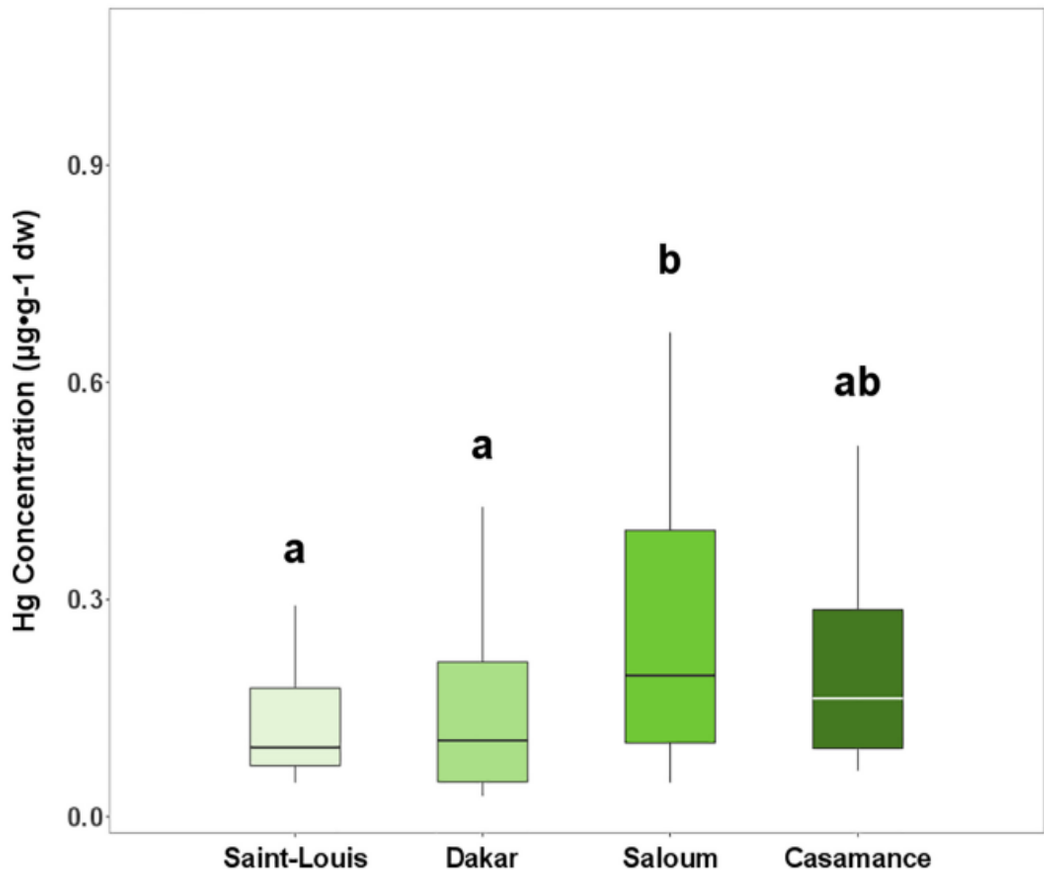


Figure 2

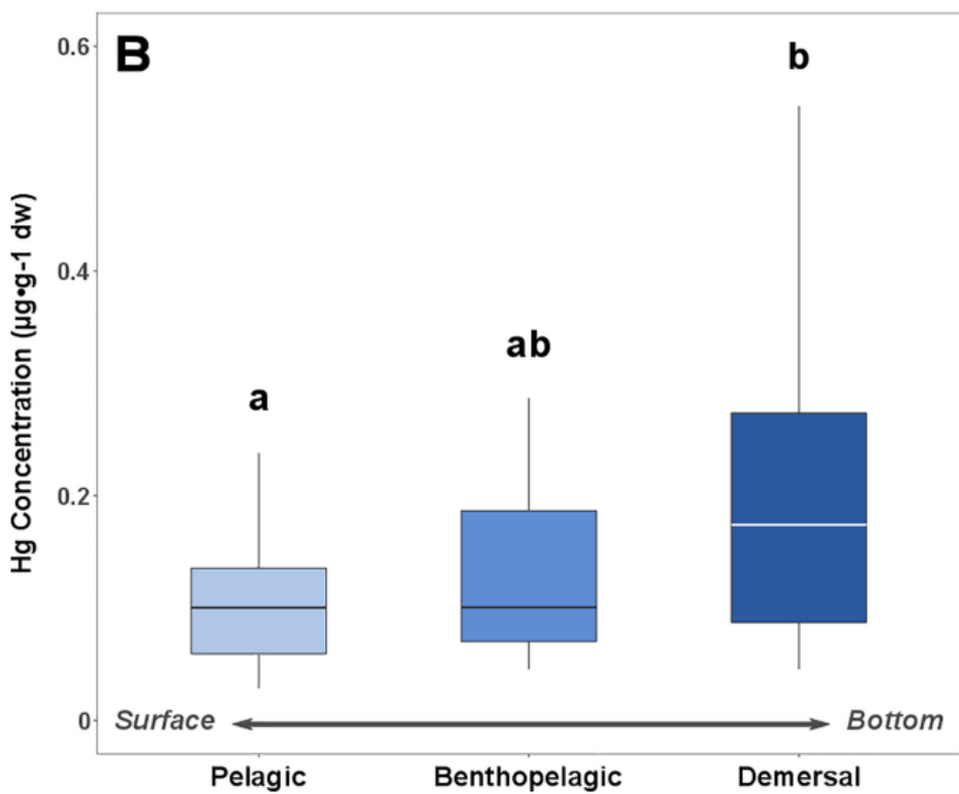
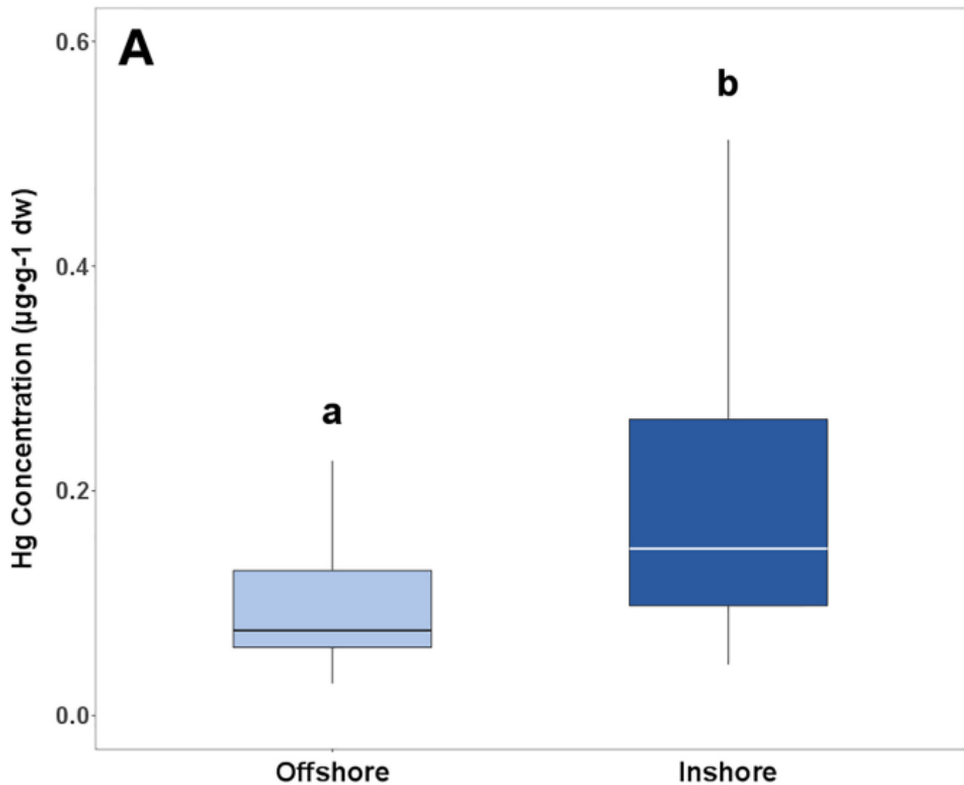


Figure 3

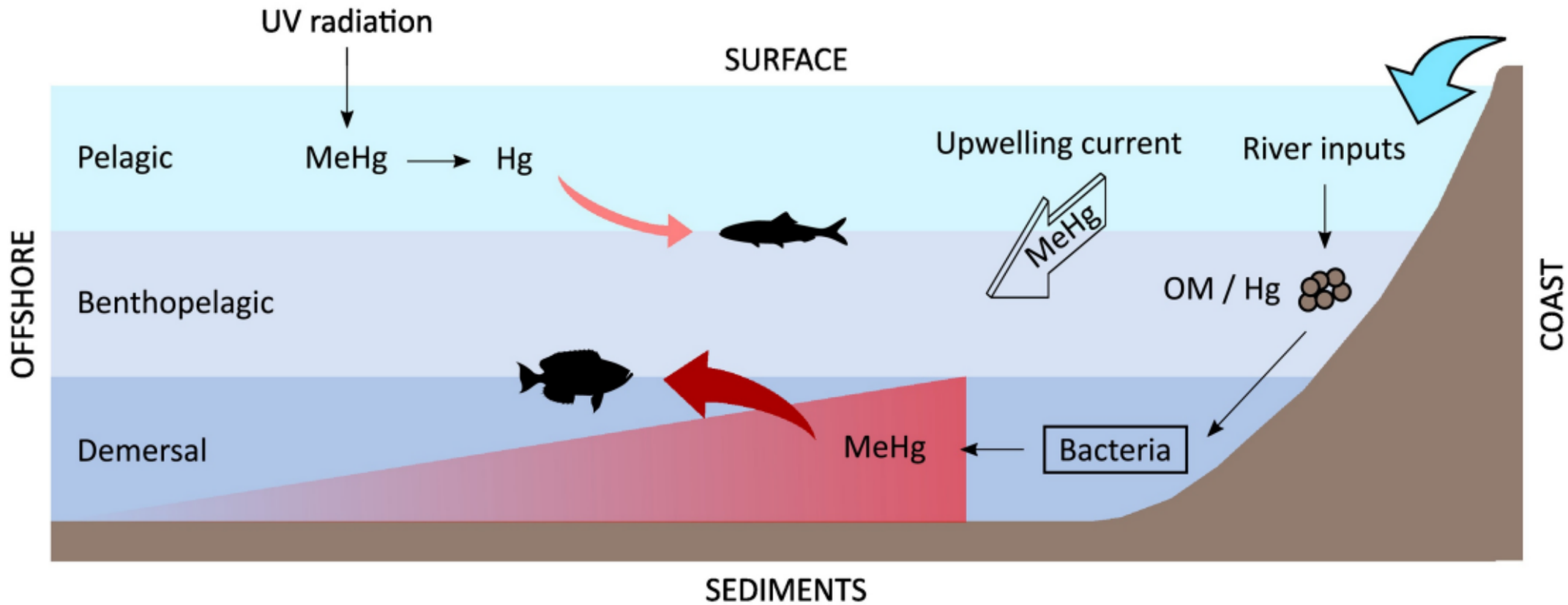


Figure 4

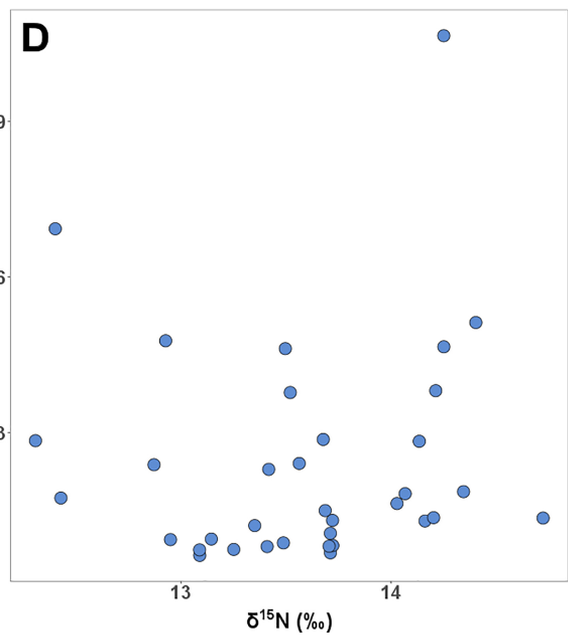
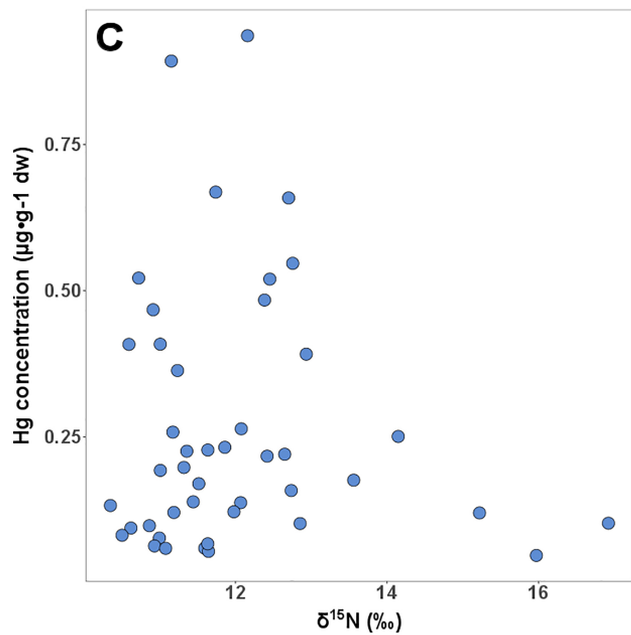
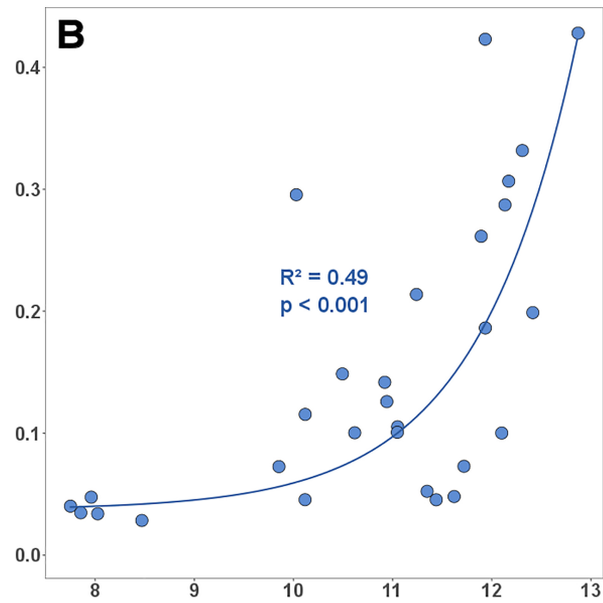
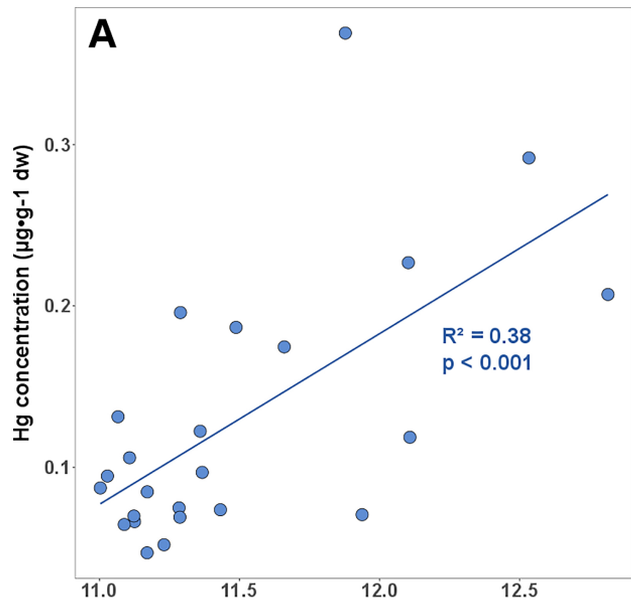


Figure 5

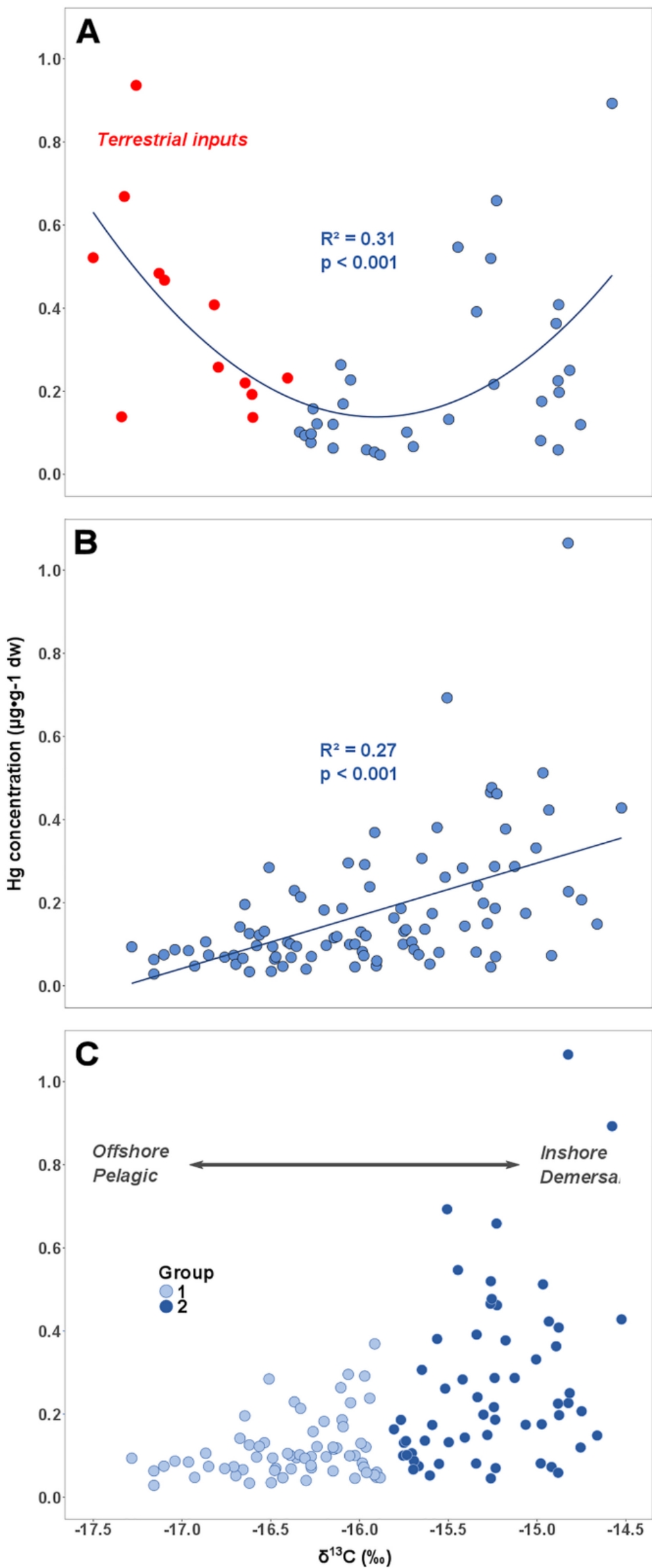


Figure 6

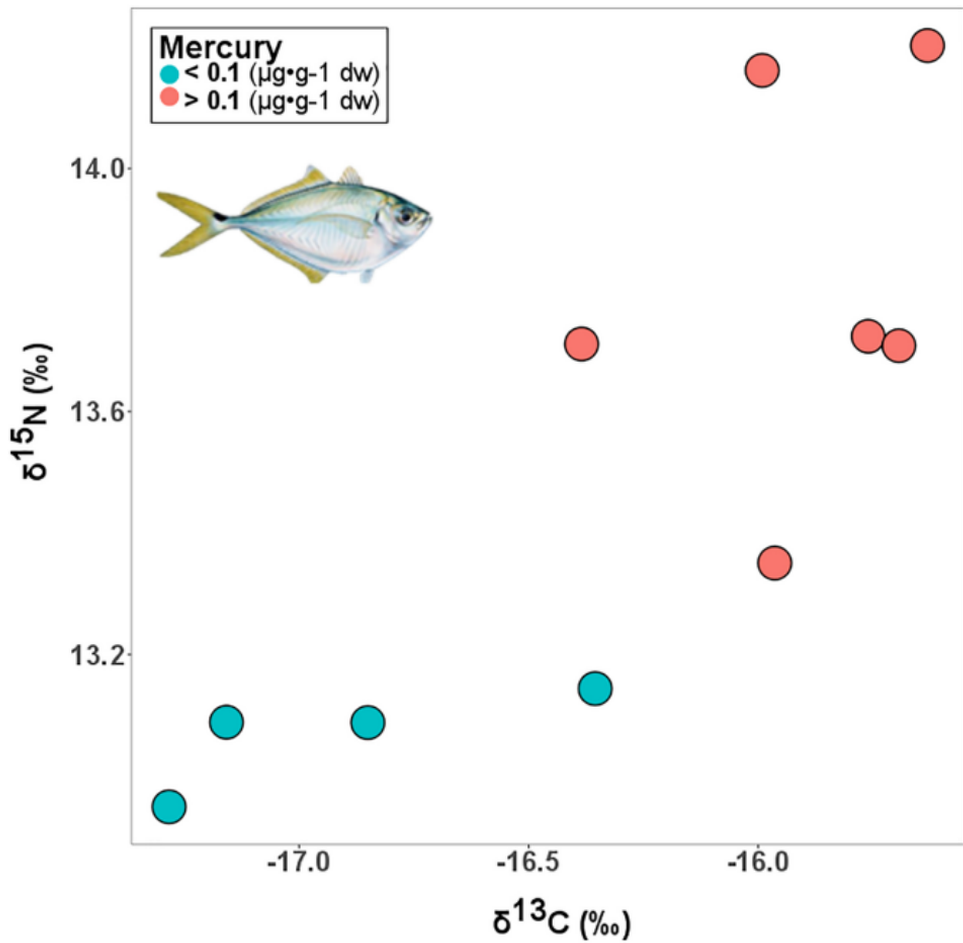


Figure 7