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The Amazon River: A Major Source of Organic Plastic Additives to the Tropical North Atlantic?

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ABSTRACT: The release of emerging organic contaminants is identified among the most critical hazards to the marine environment, and plastic additives have received growing attention due to their worldwide distribution and potential deleterious effects. Here, we report dissolved surface water concentrations of two important families of plastic additives (organophosphate esters (OPEs) and bisphenols) and other related organic compounds (perfluorinated chemicals) measured in the North Atlantic from Cape Verde to the West Indies. We found that OPEs were the most abundant contaminants, reaching remarkably high concentrations in open ocean waters (1200 km offshore of the American Coast, at the location of the Amazon river plume during the sampling period), with up to 1.3 μg L⁻¹ (ΣOPEs). A Lagrangian analysis confirmed that these high concentrations of contaminants originated from the Amazon River plume and were transported more than 3000 km by the North Brazil Current and its retroflexion. We thus consider the Amazon River as a major source of organic contaminants of emerging concern to the tropical North Atlantic Ocean and suggest that medium-/long-range contaminant transport occurs, most certainly facilitated by the highly stratified conditions offered by the river plume.

INTRODUCTION

Over the past few decades, the dramatic increase in plastic and oil-resistant properties and therefore are used in outdoor textiles and food packaging as well as in fire-fighting foams. Perfluorinated compounds (PFCs) have water- and oil-resistant properties, which qualifies them as persistent pollutants in the environment. Perfluorooctanesulfonic acid (PFOS) and PFOS precursors have therefore been listed as persistent organic pollutants (POPs) under the Stockholm Convention. PFCs have been detected in remote regions, such as the Central Arctic Ocean, and the most oligotrophic areas of the subtropical Atlantic, Pacific, and Indian oceans as well as in coastal waters. The main source of PFCs in the marine environment is river inputs.

To study their waterborne occurrence and possible sources in the tropical North Atlantic Ocean, surface seawater samples were collected in October 2017 at 14 coastal and open ocean stations (Table S1) during the Transatlantic Sargassum expedition (https://doi.org/10.17600/17016900) on the M/V Yersin from Cape Verde to Martinique Island (Lesser Antilles) along 6000 km. Seawater samples were analyzed for
20 organic contaminants of emerging concern, including OPEs, BPs, and PFCs, by applying a multicontaminant extraction protocol. A Lagrangian analysis was used to investigate the origin of the sampled surface waters.

**METHODS**

Contaminant Analysis. One-liter seawater samples were collected in duplicate at 14 sampling stations (Table S1, Figure 1), amounting to a total of 28 samples. They were collected using an inox collector, poured into corresponding precombusted (450 °C, 6 h) glass bottles, GF/F-filtered (0.7 μm), and stored in a refrigerator for a maximum of 72 h before sample processing. Samples were analyzed for nine organophosphate esters (tripropyl phosphate (TPP), trisobutyl phosphate (TmBP), trimethyl phosphate (TmPP), tris(2-chloroethyl) phosphate (TCEP), tris(2-chloro-1-methylethyl) phosphate (TmBP), triphenyl phosphate (TPhP), 2-ethylhexyldiphenyl phosphate (TEHP), perfluorooctanoic acid (PFOA), perfluorooctanesulfonic acid (PFOS), perfluorooctane sulfonic acid (PFOSF), perfluorooctane sulfonic acid (PFOSF), and perfluorooctane sulfonic acid (PFOSF)), and six bisphenols (bisphenol A (BPA), bisphenol AF (BPAF), bisphenol AP (BPAF), bisphenol F (BPF), bisphenol S (BPS), and bisphenol Z (BPZ)). To do so, organic contaminants were extracted from batches of two replicates from three stations on-board using solid phase extraction cartridges. The seawater samples were spiked with surrogate standards (100 ng sample⁻¹ of D₂₇-TBP, D₁₈-TCPP, D₁₅-TDCP, and D₁₅-TPhP), vigorously shaken and passed through the cartridges containing 250 mg of HLB phase column (50 mm × 2.1 mm, 1.8 μm), five perfluorinated compounds (perfluorohexane acid (PFHA), perfluorohexanesulfonic acid (PFHs), perfluorooctanoic acid (PFOA), perfluorooctanesulfonic acid (PFOS), and perfluorooctane sulfon fluoride (PFOSF)), and six bisphenols (bisphenol A (BPA), bisphenol AF (BPAF), bisphenol AP (BPAF), bisphenol F (BPF), bisphenol S (BPS), and bisphenol Z (BPZ)). To do so, organic contaminants were extracted from batches of two replicates from three stations on-board using solid phase extraction (SPE). Briefly, glass cartridges containing 250 mg of HLB Oasis sorbent (Waters, Milford, MA) were mounted on a manifold (Supelco Visiprep) and rinsed three times sequentially with 10 mL of ethyl acetate and 10 mL of methanol. Next, 10 mL of ultrapure water were passed through the cartridges. The seawater samples were spiked with surrogate standards (100 ng sample⁻¹ of D₂₇-TBP, D₁₈-TCPP, D₁₅-TDCP, and D₁₅-TPhP), vigorously shaken and passed through the cartridges using polytetrafluoroethylene (PTFE) liners, which were previously cleaned thoroughly with methanol.

Afterward, 10 mL of ultrapure water was again passed through the cartridges. The cartridges were then vacuum-dried for approximately 1 h, wrapped in burnt aluminum foil, and stored in the freezer (−25 °C). Upon arrival in the ISO 6 laboratory equipped for trace-clean analysis of organic compounds, a sequential elution was performed as follows: 5 mL of hexane followed by 5 mL of hexane/dichloromethane 50:50 (v/v) (F1) and 5 mL of ethyl acetate followed by 5 mL of methanol (F2). The two fractions were then evaporated with pure N₂ gas until a final volume of approximately 50 μL was left (F1) or until dryness (F2), in which case the extract was then reconstituted using 990 μL of methanol. Internal standards were added as follows: 100 ng sample⁻¹ of D₁₂-TPrP, D₁₅-TCEP and D₇-malathion (F1), and 20 ng sample⁻¹ of BPA-D₁₆, PFHA-¹³C₅, PFBS-¹³C₅, PFHS-¹³C₅, and PFOS-¹³C₈ (F2). OPEs were quantified by GC/MS (Agilent Technology 7820A-5977E) in the splitless selected ion monitoring (SIM) mode. The separation was achieved using a 30 m × 0.25 mm i.d. × 0.25 μm HP-5MS capillary column (Agilent J&W). PFCs and BPs were quantified using LC-QTOF (Agilent 1290 Infinity LC system coupled with Agilent 6530 Accurate-Mass Q-TOF, Agilent Technologies, Les Ulis, France). Separation was achieved using an Agilent Zorbax Eclipse XDB reversed-phase column (50 mm × 2.1 mm, 1.8 μm), with the temperature at 30 °C. GC/MS and LC-QTOF method details are given in the Supporting Information (Text S1 and Tables S2 and S3). CAS numbers of each compound are given in Table S4.

**Quality Assurance/Quality Control (QA/QC).** Strict QA/QC measurements were implemented. Extraction, transport, and laboratory blanks were made and analyzed concurrently with the samples to detect possible contaminations. Blank levels and compound concentrations of samples are presented in Table S5. No contamination was found, except in the case of TCPP, where mean blank levels (n = 5) of ∼16 ng were detected, and for some PFCs and BPs, where trace levels were detected in the blanks (PFHA < LOQ, PFOA n.d. < LOQ, PFOS n.d. < LOQ, BPA n.d. < 1.0 ng L⁻¹, and BPS n.d. <LOQ). The transport blank showed similar blank levels as field blanks, excluding sample contamination during transport and storing. The results presented here are blank-corrected. Limits of quantification (LOQs) ranged from 0.03 to 8 ng L⁻¹ (all LOQs are given in Table S4). If compounds were detected at concentrations < LOQ, they were considered as zero (i.e., not detected) for the calculation of detection frequencies and sums. Mean surrogate recovery rates were as follows: D₂₇-TBP 74%, D₁₈-TCPP 70%, D₁₅-TDCP 76%, and D₁₅-TPhP 71% (see Table S6 for all recovery rates). The results presented here are not recovery corrected. The variability between duplicate samples ranged between 6 and 27%.
Solvents and Reagents. Solvents were purchased from Promochem (hexane, ethyl acetate and dichloromethane, Picograde, LGC standard) and Biosolve (methanol, ULC-MS grade). Isotope-labeled PFCs were purchased from Wellington Laboratories (Canada) and labeled OPEs from C/D/N Isotopes Inc. (Canada) (D_{27}-TBP, D_{15}-TPhP, D_{21}-TPrP, and D_{7}-malathion) and from Cambridge Isotope Laboratories, Inc. (USA) (D_{18}-TCP, D_{15}-TDPP and D_{12}-TCEP). Isotope-labeled bisphenols were obtained from AccuStandard (USA).

Salinity and Velocity Maps. Daily surface salinity and velocity fields at 1/12° (∼8 km) were computed by the Operational Mercator global ocean analysis and forecast system PSY4 V3R1 from MERCATOR-OCEAN. This system used version 3.1 of the NEMO ocean model, which solves the three-dimensional primitive hydrodynamic equations in spherical coordinates discretized on a C-grid and 50 vertical levels, under the hypothesis of boussinesq, hydrostatics, and incompressibility. This model is forced by realistic surface fluxes from ECMWF and jointly assimilates altimeter data, vertical in situ temperature and salinity profiles, and satellite sea surface temperature. Data were downloaded for the surface layer (<1 m depth) of the whole North Atlantic Ocean from the Marine Copernicus Portal (GLOBAL_ANALYSIS_FORECAST_PHY_001_024; http://marine.copernicus.eu/services-portfolio/access-to-products/).

Backward Trajectories. Backward Lagrangian particle trajectories were computed using the Lagrangian transport model Ichthyop, which simulates horizontal and vertical advection and dispersion. In our study, we considered our particles as passive surface tracers and only used horizontal advection in the movement equation. The Ichthyop model was run using the MERCATOR-OCEAN daily surface velocity fields at 1/12° described above. These fields were interpolated...
in space and time using a fourth order Runge–Kutta integration scheme with a constant time step of 1800 s that respected the Courant–Friedrichs–Lewy (CFL) criterion on the entire domain. Each sampled surface water mass was modeled by a circular patch with a radius of 10 km and seeded at the surface with 400 randomly released particles (i.e., approximately 1.3 particles km\(^{-2}\)) at the station times and geographic coordinates. Horizontal particle advection was carried out for 180 days with an output time step of 12 h. At the end of each simulation, the percentage of seeded particles that passed in the Amazon Estuary area was computed, as well as the number of days between the day that they left the Amazon estuary area and the sampling day of the water masses.

The Amazon Estuary area is hereby represented by a box, including longitude between \(-52^\circ\)E and \(-46^\circ\)E and latitude between \(-2^\circ\)N and \(3^\circ\)N. Simulation was not run for station Y16 because of its high proximity to the coast (4 km; lower than the current model resolution).

The choice of a particle patch radius of 10 km for the backward trajectory simulations was motivated by two arguments: the radius needed (i) to be small enough to be representative of the sampled water mass and (ii) to be large enough to include at least two meshes of the velocity field in diameter to take into account possible shifts between modeled and actual velocity fields. Comparison between daily salinity maps and seeded patch areas shows that a 10-km patch size allows for representation of a water mass with a salinity variability lower than \(\pm 0.2\), in comparison with the overall salinity varying between \(\sim 34\) inside the Amazon River plume to 37 in the surrounding seawater (Figure S1). In addition, sensitivity tests for radii from 5 to 25 km and densities from 0.5 to 2.5 particles km\(^{-2}\) were performed to test the robustness of our results (Table S7). The upper limit of 25 km was chosen because higher radii showed too much salinity variability inside patch areas to be consistent with the sampled water mass. These tests showed that the sensitivity of our results to the seeded patch radius and the particle density was very low, with a maximum difference in particle percentage that passed through the Amazon estuary area (Amazon box) lower than 10%, except at stations Y04 (40%) and Y14 (26%).

**Figure 3.** Concentrations (in ng L\(^{-1}\)) of the individual BPs (A) and PFCs (B) detected at sampling stations Y01–Y16. Monthly average salinity distribution in the surface waters of the tropical North Atlantic in October 2017 is indicated as well as the Amazon River mouth (blue star).
RESULTS AND DISCUSSION

OPEs were detected in all samples, while BPs and PFCs were each only detected in 3 of 14 samples. From the nine OPEs analyzed, TTP, TnBP, TEHP, and TDCP were not detected, while TCEP was found only at trace levels <2 ng L$^{-1}$ in 36% of samples. TCPP was detected in all samples, followed by TiBP and TPhP (93 and 57% detection frequency, respectively, Figure S2). Concerning bisphenols, BPA was detected at stations Y01, Y09, and Y10, and BPS was found at station Y10, while the remaining four bisphenols were not detected in any samples. Of the five PFCs studied, only PFOSF was not detected at all. PFHS, PFOA, and PFOS were found at one sampling station (Y11), while PFHA was detected in 21% of samples.

OPEs were the most abundant compound class with a relative abundance of 98–100%, followed by PFCs (0–1.9%) and, finally, bisphenols (0–0.6%). The highest amounts of Σ9OPEs were measured at stations Y11, Y12, and Y10 (1340, 1000, and 955 ng L$^{-1}$, respectively), hence, in offshore waters (Figure 2A). In contrast, the two stations showing the lowest Σ9OPEs concentrations (Y16 and Y14, 74 and 106 ng L$^{-1}$, respectively) are in the most western coastal area. TCPP was hereby the most abundant OPE, with concentrations ranging from 74 (station Y16) to ~1300 ng L$^{-1}$ (station Y11) and a relative abundance ranging from 92 to 100%. TiBP concentrations ranged from <1 (station Y16) to 38 ng L$^{-1}$ (station Y11). In contrast, the maximum TPhP and EHDPP concentrations were measured at station Y03 (7 ng L$^{-1}$ and 5 ng L$^{-1}$, respectively), closer to the NW African coast (Figure 2B). These two compounds exhibit thus a different distribution pattern than Σ9OPEs, possibly due to different sources and/or degradation rates. TPhP and EHDPP are used in hydraulic fluids, among others, and the fact that they exhibited highest concentrations at stations closest to the NW African coast could possibly be due to the high fishing activities in this area and other maritime traffic. This is in agreement with another study reporting that EHDPP concentrations seem to derive mainly from local sources (airports, harbors, and naval bases, urbanized and industrial areas) instead of from waterborne transport.

Station Y11, which was characterized by the highest Σ9OPEs concentrations, is the station closest to the Amazon River estuary, with a distance of approximately 1200 km. Despite the large distance involved, diluted water from the Amazon River has already been found in the Caribbean Sea and has been shown to influence the tropical Atlantic Ocean more than 3500 km away from the river mouth. The freshwater is hereby transported by the North Brazil Current (NBC) and the North Equatorial Counter Current (NECC). Our results indicate, that medium-/long-range transport of OPEs via the water phase is occurring, further challenging the consideration of OPEs as being environmentally friendly alternatives for brominated flame retardants. This observation seems to be especially true for chlorinated OPEs, such as TCPP, since they are more persistent than nonchlorinated OPEs. Furthermore, it has been suggested that chlorinated OPEs have a longer characteristic travel distance in water than in air, supporting the hypothesis of water-borne medium-/long-range transport. This could be a particular reason for concern, since especially chlorinated OPEs are considered to have negative impacts on human health and the environment.

Generally, OPE levels in rivers and their estuaries can be very variable and can depend on the season. TCPO concentrations ranging from 4.6 to 921 ng L$^{-1}$ were measured in rivers in northern China, concentrations of 126 ± 14 ng L$^{-1}$ were found in the Elbe River in Germany, and concentrations of 1.1–4.4 μg L$^{-1}$ were measured in streams in Toronto, Canada. These ranges are comparable with our results (74–1300 ng L$^{-1}$ of TCPP).

Bisphenols and perfluorinated compounds were present at lower concentrations than OPEs (Table S5). BPA concentrations ranged between 1.2 ng L$^{-1}$ (station Y01) and 1.8 ng L$^{-1}$ (Y09 and Y10), while BPS was found at 1.1 ng L$^{-1}$ (Y10) (Figure 3A). While BPS is used as a substitute for BPA in certain products (e.g., in thermal paper or polycarbonate plastics), BPA is still dominating these sectors, probably explaining why BPA was more abundant in our samples than BPS. The finding that bisphenols were only found at concentrations lower than 2 ng L$^{-1}$ or not detected at all could be due to rapid degradation rates. According to the simulation of the backward trajectories (see below), the surface waters needed at least two months after leaving the Amazon River estuary before they reached the sampling stations. It was observed that complete biodegradation of BPA in river water was achieved within a few days, even at high initial concentrations (up to 5500 μg L$^{-1}$). The presence of bisphenols at stations Y09 and Y10, and especially at Y01, is thus probably not necessarily linked to the Amazon River, but could for example originate from ships that recently passed through or near the sampling area. Similar to the OPE results, Y11 was the station that exhibited the highest Σ9PFCs concentration (26 ng L$^{-1}$) and, in particular, the highest PFHA concentration (16 ng L$^{-1}$), followed in this case by the adjacent stations Y12 and Y10 (11 and 9.7 ng L$^{-1}$, respectively). At station Y11, PFHS was found at 4.4 ng L$^{-1}$, PFOA at 2.9 ng L$^{-1}$, and PFOS at 2.3 ng L$^{-1}$ (Figure 3B). This PFOS concentration is quite high, especially for open-ocean waters. As a reference, it clearly exceeded the European Environmental Quality Standard (EQS) for PFOS (0.13 ng L$^{-1}$ for marine waters, directive 2013/39/EU). PFHA concentrations in seawater generally tend to be very low, with 0.02–0.19 ng L$^{-1}$ of PFHA measured in Western Mediterranean waters and <0.003–0.12 ng L$^{-1}$ detected in seawater samples from Northern Europe, the Atlantic Ocean, and the Southern Ocean. A study analyzing PFHA in surface seawater samples close (at a distance of 140–370 km) to our sampling stations Y16, Y14, and Y05 found concentrations between n.d. and 0.0187 ng L$^{-1}$, values that were comparable with ours (n.d. or <LOQ) at the three stations mentioned above. In contrast, PFHA concentrations in rivers have been found to vary between 0.50 and 5.3 ng L$^{-1}$ (Elbe and Weser Rivers in Germany) and between <0.10 and 38 ng L$^{-1}$ (data from 29 Chinese rivers) or reach peak concentrations of up to 16000 ng L$^{-1}$ downstream of a PFC production facility (Samondogawa River in Japan). Our PFHA concentrations at stations Y11, Y12, and Y10 (16, 11, and 9.7 ng L$^{-1}$, respectively) thus seem quite high for seawater samples and indicate once again a riverine source, i.e., the Amazon River.

A recent study proposed fire-fighting foams used at offshore oil and gas platforms to be a potentially important PFC source to the marine environment. Important oil and gas drilling close to our study area mainly occurs in the Campos and Santos basins off the coasts of Rio de Janeiro and Sao Paulo, Brazil. Given the direction of the oceanic currents there (from}
north to south (not toward our sampling stations), these offshore platforms do not seem to be a plausible explanation for the high PFC concentrations detected at stations Y10−Y12. However, multiple studies\textsuperscript{46−48} have investigated the magnitude of oil and gas extractions in the Amazon basin and the far-reaching environmental impacts. These inland oil and gas extraction activities and the associated use of fire-fighting foams could thus be a possible explanation of the PFC concentrations observed. Another potential source of PFCs, OPEs, and BPs in the Amazon basin could be the city of Manaus, which houses an important industry producing chemicals and electronic equipment, among others. Furthermore, more than two million people live in Manaus, but less than 20% of households are connected to the sewage system.\textsuperscript{49}

Low surface salinity (between 33 and 35; Table S1 and Figure S3) was measured at all stations. In the west, this freshening is thought to be primarily due to transport of freshwater from the Amazon, while to the east it mainly emanates from Intertropical Convergence Zone (ITCZ) rainfall.\textsuperscript{50} These trends were also observed for the chlorophyll a spatial concentration distribution, represented by Kd490 (Figure S4). To confirm the origin of the sampled surface waters, a Lagrangian analysis was performed. Simulations using MERCATOR daily surface velocity fields indicated that 98−100% of the surface waters encountered at stations Y06−Y08 and Y10−Y13 passed near or originated from the Amazon River mouth. While the model cannot distinguish whether the surface waters passed by the Amazon River estuary or originated from the river itself, the term "originated from" will be used hereafter in accordance with our hypothesis. According to the simulation, the farthest east where an influence of the Amazon can be observed is at station Y04, where 82% of surface waters originated from the far east corner of the black box enclosing the river estuary (Figure 4; backward trajectories for all stations are provided in Figure S5). These results indicate that the influence of the Amazon River is reaching far into the North Atlantic since even water masses and contaminant concentrations from sampling stations close to the African continent were partially linked to the Amazon River. Hereby, the surface waters of station Y04 (8°N, 21°W) had a calculated travel time of ∼125 days, which is in good agreement with the results obtained by Coles et al., who found that drifters released in the Amazon River estuary reached this zone after approximately 120−150 days.\textsuperscript{51} Stations with high contaminant levels seem generally characterized by (i) a high percentage of surface waters that originated in the Amazon River plume, (ii) a short travel time, and/or a short travel distance and (iii) an origin close to the river mouth where contamination is probably highest. However, we point out that the Lagrangian approach used to simulate the backward trajectories does not take vertical mixing into account, which can complexify (or partly explain some discrepancies in) the relationship between contaminant concentration and percentage, origin, and travel time of

![Figure 4. Water mass backward trajectories for the stations Y01 (a), Y04 (b), Y11 (c), and Y14 (d). Each map has a caption indicating the station ID, the date, the percentage of water masses that were in the black box (indicating the Amazon Estuary), and the number of days that passed between the sampling of the water masses and their journey in the area of the black box (in brackets). The minimum salinity (S) reached by particles during their journey in the black box is indicated as well as the river plume (with a dark to light gray scale; S = 27−37).](image-url)
exports considerable amounts of DOC and dissolved organic that future climate scenarios predict an increase in the importance of the situation is highlighted when we consider geographical zones of scientific interest and to note the current financial and logistical support and for the funding of A.O.'s postdoctoral fellowship. Thanks to the Ministère de l’Écologie et du Développement Durable for financial support. The project leading to this publication has received funding from European FEDER Fund under project 1166-39417. The project has also received funding from the JPI Oceans PLASTOX project and the Agence de l’Eau Rhône Méditerranée Corse Blue-POLUT and CAREMED projects.

**REFERENCES**


discharge of the Amazon River$^{50}$ and that plastic litter in the Amazon watershed is expected to double between 2010 and 2025.$^{51}$

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.est.9b01585.

GC/MS and LC−HRMS method details; salinity maps of seeded particle patches; detection frequencies of organic compounds; daily surface salinity maps; maps of sampling area and Kd490 distribution; water mass backward trajectories; air mass backward trajectories; GPS positions and measured salinity; quantification ions; LOQs; contaminant concentrations in blanks and samples; surrogate recoveries; sensitivity test results for water mass backward trajectories (PDF)

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**Notes**

The authors declare no competing financial interest.

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