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OPEN



Nutrient content and stoichiometry of pelagic *Sargassum* reflects increasing nitrogen availability in the Atlantic Basin

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The pelagic brown macroalgae *Sargassum* spp. have grown for centuries in oligotrophic waters of the North Atlantic Ocean supported by natural nutrient sources, such as excretions from associated fishes and invertebrates, upwelling, and N₂ fixation. Using a unique historical baseline, we show that since the 1980s the tissue %N of *Sargassum* spp. has increased by 35%, while %P has decreased by 44%, resulting in a 111% increase in the N:P ratio (13:1 to 28:1) and increased P limitation. The highest %N and δ^{15} N values occurred in coastal waters influenced by N-rich terrestrial runoff, while lower C:N and C:P ratios occurred in winter and spring during peak river discharges. These findings suggest that increased N availability is supporting blooms of *Sargassum* and turning a critical nursery habitat into harmful algal blooms with catastrophic impacts on coastal ecosystems, economies, and human health.

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or over five centuries, the floating brown macroalgae of the North Atlantic Ocean (NA) known as pelagic Sargassum has stirred debate and mystery among seafarers and scientists alike. This vegetation was first described by Christopher Columbus and his sailors in 1492, which reminded them of "salgazo," small grapes in Portugal, and thus the name of the central gyre of the NA became the Sargasso Sea¹. The vegetation is comprised of two holopelagic Sargassum species, S. natans and S. *fluitans*, that reproduce solely by vegetative propagation². Early oceanographers and marine botanists thought this vegetation grew primarily in the Sargasso Sea, which they estimated to contain 7 to 10 million tons 3,4 (Fig. 1). However, this presented a paradox to modern oceanographers who considered the Sargasso Sea a biological desert due to the very low nutrient concentrations and biological productivity in its surface waters (Ryther's Paradox)¹.

This paradox has since been explained by the seasonal transport of nutrient enriched and productive Sargassum from the Gulf of Mexico (GOM), Loop Current, and Gulf Stream to the Sargasso Sea. Studies of the productivity and nutrition of pelagic Sargassum showed that neritic plants in the southeastern GOM, Loop Current, and western wall of the Gulf Stream along the southeastern United States had twofold higher productivity and lower carbon:nitrogen (C:N) and carbon:phosphorus (C:P) ratios compared to oceanic populations in the Sargasso Sea^{5,6}. Major advances in remote sensing of Sargassum using Medium Resolution Imaging Spectrometer (MERIS) and Moderate Resolution Imaging Spectroradiometer (MODIS) satellite imagery revealed extensive and frequent windrows of Sargassum (line-shaped aggregations formed by wind forcing) in the western GOM in 2004 and 20057. High biomass strandings of Sargassum along GOM coastlines since the 1980s have led to intensive beach raking⁸ and an emergency shutdown of a nuclear power plant on the west coast of Florida⁹, perhaps as a consequence of increasing N inputs to the GOM from the Mississippi River and its distributary the Atchafalaya River, as well as other land-based sources^{5,10}. The extensive biomass of Sargassum in the western

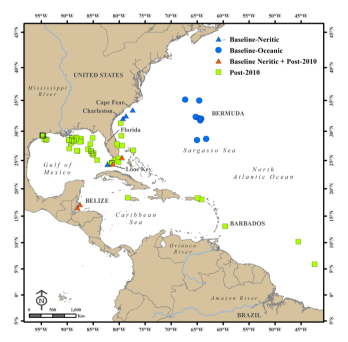


Fig. 1 *Sargassum* **collection locations.** Locations in the North Atlantic Ocean where *Sargassum* samples were collected during the 1980s baseline study⁵ (blue), post-2010 collections (green), and during both time frames (orange).

GOM is proposed to be advected seasonally via the Loop Current and Gulf Stream to the Sargasso Sea¹¹. For the first time, physical connectivity was established linking the abundant *Sargassum* populations in the GOM to nutrient-poor populations in the Sargasso Sea, helping to explain Ryther's Paradox¹.

Beginning in 2011, a new region of concentrated Sargassum biomass developed in the Tropical Atlantic Ocean south of the Sargasso Sea^{12,13}, where it had not been previously observed³. This new region may have been seeded by an extreme negative phase of the North Atlantic Oscillation in 2009 to 2010 that provided windage to transport Sargassum from the Sargasso Sea to the east and ultimately into the North Equatorial Current and central Tropical Atlantic Ocean¹⁴, although this is not evident from satellite imagery. Long-term satellite data, numerical particle-tracking models, and field measurements indicate that a newly formed Great Atlantic Sargassum Belt (GASB) has recurred annually since 2011 and extended up to 8850 km from the west coast of Africa to the GOM, peaking in 2018¹⁵. Over its broad distribution, the GASB can be supported by N and P inputs from a variety of sources including discharges from the Congo, Ama-zon, and Mississippi rivers^{5,15–17}, upwelling off the coast of Africa^{15,16}, vertical mixing⁵, equatorial upwelling¹⁸, atmospheric deposition from Saharan dust, and biomass burning of vegetation in central and south Africa^{16,19}.

Similar to the recent development of macroalgal blooms in the Yellow Sea and the East China Sea^{20,21}, the increasing golden tides of Sargassum in the GOM and GASB could be ecological indicators of large-scale, oceanic eutrophication^{15,22}. Excessive biomass strandings of Sargassum have had catastrophic consequences on ecosystem and human health in coastal areas, negatively impacting seagrasses²³, coral reefs^{24,25}, and a number of suitable sea turtle nesting and hatching areas²⁶. Sargassum removal from Texas beaches during earlier, less severe inundations was estimated at \$2.9 million per year²⁷ and Florida's Miami-Dade County alone estimated recent removal expenses of \$45 million per year. The Caribbean-wide clean-up in 2018 cost \$120 million, which does not include decreased revenues from lost tourism. Sargassum strandings also cause respiratory issues from the decaying process and other human health concerns, such as increased fecal bacteria. During large-scale strandings in 2018, more than 11,400 residents in Martinique and Guadeloupe were diagnosed with acute exposure to toxic H₂S gas produced by decaying Sargassum²⁸.

Increases in harmful algal blooms (HABs) in recent decades are related to global increases in nutrient pollution^{29,30}. Human activities have greatly altered global C, N, and P cycles, and N inputs are considered now high risk and above a safe planetary boundary³¹. Based on scientific research, population growth and land-use changes have increased N pollution and degradation of estuaries and coastal waters since at least the 1950s^{30,32-34}. Despite decreases in N loading in some coastal watersheds, N:P ratios remain elevated in many rivers compared to historic values³⁵. Although the relative importance of N vs. P limitation in the open oceans has been debated^{36,37}, previous analyses of tissue C:N:P data suggest that both N and P potentially limit the growth of pelagic *Sargassum* over its broad geographic range^{5,6}. Here, the objective was to better understand the effects of N and P supply on Sargassum, where a unique baseline tissue C:N:P data set from the 1980s^{5,6} are compared with more recent samples collected since 2010 (Fig. 1).

Results

A total of 488 tissue samples of *Sargassum spp.* were collected during various research projects and cruises in the NA basin between 1983 and 2019. The baseline 1980s samples (41) included

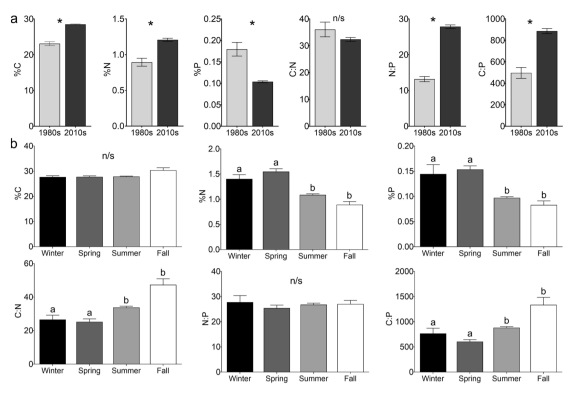


Fig. 2 Sargassum tissue nutrient contents. Tissue elemental composition and C:N:P stoichiometry (mean \pm SE) of Sargassum natans and S. fluitans collected throughout the NA in the 1980s and post-2010. **a** by decade with asterisks representing significant differences and **b** by Northern Hemisphere meteorological season with different lowercase letters representing significant differences identified with Tukey HSD test; "n/s" denotes a non-significant (P > 0.05) ANOVA result.

seasonal sampling of S. fluitans (21 samples) and S. natans (20 samples) at offshore Looe Key reef in the lower Florida Keys in 1983 and 1984 and a broader geographic sampling in 1986 and 1987 from neritic stations offshore the Florida Keys (Looe Key, Dry Tortugas), Gulf Stream (Miami, FL; Charleston, SC; Cape Fear, NC), and Belize, Central America (Glovers Reef, Belize City). Oceanic stations included the northern, central, and southern Sargasso Sea (Fig. 1)⁶. Seasonally these baseline samples consisted of winter (2), spring (15), summer (20), and fall (4) collections. Since 2010, additional samples (447 total) of S. fluitans (302) and S. natans (145) were collected in a variety of locations in the wider NA, including Looe Key, western Florida Bay, the Gulf Stream, coastal waters along the east and west coasts of Florida, various stations in the GOM, Belize, the Caribbean region, and in the Amazon River plume (Fig. 1). The post-2010 samples also spanned winter (28) spring (97), summer (327), and fall (36).

Changes in Sargassum tissue chemistry. Tissue analysis of Sargassum over broad areas of the NA revealed significant changes in N and P contents since the 1980s, indicating widespread N enrichment and increased P limitation. %N and %C increased concurrent with a decrease in %P in Sargassum tissue from the 1980s to 2010s (Fig. 2a). Elemental composition varied significantly between these two decades (MANOVA, Pillai's lambda = 0.201, $F = {}_{3,470}$ 39.4, P < 0.001; Supplementary Table 1). Subsequent univariate analyses revealed significant increases (23%) from the 1980s to the 2010s for %C (ANOVA, F = 153.8, P < 0.001) and %N (35%; ANOVA, F = 15.01, P = 0.026), while %P decreased significantly (-42%; ANOVA, F = 1 31.4, P < 0.001) over the long-term study (Fig. 2a). The C:N:P ratios also varied by decade (MANOVA, Pillai's lambda = 0.236, $F = {}_{3,470}$ 48.4, P < 0.001; Supplementary Table 2). Notably, the biggest change was the N:P ratio, which increased significantly (111%; ANOVA, F = 1 93.4, P < 0.001). C:P ratios also increased similarly

(78%; ANOVA, F = 1 44.9, P < 0.001; Fig. 2a). Although the C:N ratio decreased (-10%) from the 1980s to the 2010s, this change was not significant (ANOVA, F = 1 < 0.001, P = 0.956; Supplementary Table 2, and Fig. 2a). As such, the decadal patterns of increasing %C and %N with decreasing %P observed in *Sargassum* are consistent with observed changes in molar C:N:P ratios.

Seasonal patterns were also observed in elemental composition of Sargassum with higher %N and %P in the winter and spring (Fig. 2b). Elemental composition varied significantly with season (MANOVA, Pillai's lambda = 0.147, $F = _{9,1416}$ 8.13, P < 0.001; Supplementary Table 1). Tissue %N (ANOVA, F = 3 17.1, P <0.001) and %P (ANOVA, F = 3 16.7, P < 0.001) was significantly higher during winter and spring than in summer and fall, but %C was not seasonally variable (ANOVA, F = 3 2.58, P = 0.053). Further, tissue C:N:P ratios also varied with season (MANOVA, Pillai's lambda = 0.115, F = 9.1416 6.25, P < 0.001; Supplementary Table 2). Both the C:N (ANOVA, F = 3 13.4, P < 0.001) and C:P (ANOVA, F = 3 12.5, P < 0.001) ratios were significantly lower in the winter and spring compared to the summer and fall (Supplementary Table 2). The N:P ratio was not seasonally variable (ANOVA, F = 3 0.930, P = 0.427; Supplementary Table 2). These seasonal patterns demonstrate higher %N and %P contents in the winter and spring with no seasonal fluctuations in %C or N:P ratio (Fig. 2b).

Significant interactions between season and decade were observed in *Sargassum* tissue chemistry. For elemental composition, this interaction was significant (MANOVA, Pillai's lambda = 0.055, $F = _{9,1416}$ 2.95, P = 0.002; Supplementary Table 1, Supplementary Fig. 1). For %C the interaction of season and decade was not significant (ANOVA, $F = _3$ 1.90, P = 0.129) but there were significant interactions for %N (ANOVA, $F = _3$ 3.00, P = 0.030) and %P (ANOVA, $F = _3$ 3.91, P = 0.009). For %N, all seasons increased from the 1980s to 2010s, except for winter, which slightly

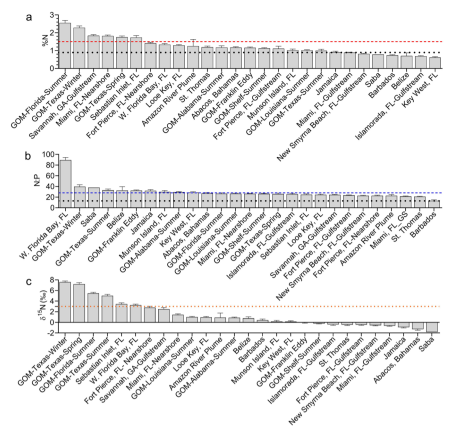


Fig. 3 Post-2010 *Sargassum* **tissue nutrient contents by location.** Post-2010 *Sargassum* tissue nutrient contents by location (mean ± SE), as well as Northern Hemisphere meteorological season for Gulf of Mexico (GOM) samples, indicating where %N and N:P ratios were greater than the 1980s baseline mean for the entire dataset (black dotted lines). **a** For %N, values have significantly increased from the 1980s (decadal mean = 0.89%) to post-2010 (decadal mean = 1.21%); %N values >1.5 (red dashed line) are considered non-limiting to macroalgal growth⁴³. **b** N:P ratios have significantly (111%; ANOVA, F = 1 93.4, P < 0.001) increased from the 1980s (decadal mean = 13.2) to post-2010 (decadal mean = 27.8, blue dashed line). **c** Enriched δ^{15} N values (>+3%e, orange dotted line) are indicative of urbanized wastewater discharges, while more depleted values are indicative of N₂ fixation, atmospheric deposition, and upwelling.

decreased (Supplementary Fig. 1). Conversely, %P decreased between decades for all seasons, except winter, which slightly increased (Supplementary Fig. 1). These interactions suggest that 1980s winter had a different pattern than the other seasons, which may be an artifact of the very small sample size (2) from just one location (Looe Key) for 1980s winter.

Geographic patterns in Sargassum tissue chemistry. Overall, the %N of Sargassum spp. increased 35% over the period of study, while %P decreased by 44%, resulting in more than a doubling of the N:P ratio from 13:1 to 28:1, well above the Redfield Ratio of 16:1. The highest %N, N:P ratios, and stable nitrogen isotope values (δ^{15} N) were in neritic waters heavily influenced by river discharges and land-based runoff (Fig. 3, Supplementary Fig. 2, and Supplementary Table 3). The overall range of %N was 0.15 to 3.05% with the highest mean %N observed in coastal waters of the GOM (2.55% in summer offshore of Florida's west coast, 2.28% in winter offshore of Texas), Florida's east coast (1.82% offshore of Miami, 1.73% near Sebastian Inlet, 1.33% in western Florida Bay), southeast United States (1.84% offshore Savannah, GA) and the offshore Amazon plume (1.25%; Fig. 3a and Supplementary Fig. 2a). The lowest %N was observed offshore of Key West, FL (0.61%; Fig. 3a). Twenty of the post-2010 sampling events had % N values greater than the mean %N from the 1980s (Fig. 3a). The overall range for N:P ratios was 4.66 to 99.2 with the highest in western Florida Bay, FL (89.4), followed by locations in the GOM and Caribbean (Fig. 3b and Supplementary Fig. 2b). The lowest

N:P ratios were observed in the eastern Caribbean at St. Thomas (20.9) and Barbados (13.0; Fig. 3b). Twenty-six of the post-2010 sampling events had N:P ratios greater than the mean N:P from the 1980s (Fig. 3b). δ^{15} N values were variable with an overall range of -5.58 to +8.99%, indicating multiple sources of N were available to *Sargassum* (Supplementary Fig. 2c). High values (>+5‰) occurred along the urbanized Texas coast that is also affected by the Mississippi River plume (Supplementary Fig. 2c). The lowest δ^{15} N values occurred at Saba (-1.83%) in the Leeward Islands of the northeastern Caribbean (Fig. 3c). δ^{15} N values of *Sargassum* collected in the Gulf Stream were also generally low (<-1%), except offshore of Savannah, GA (+2.5%; Fig. 3c).

Discussion

In a series of shipboard experiments during the 1980s, *Sargassum* productivity and growth was enhanced by enrichment with both nitrate⁵ (NO₃⁻) and soluble reactive phosphorus³⁸ (SRP), which resulted in higher tissue levels of N and P. In oligotrophic surface waters of the NA, dissolved inorganic N (DIN) and SRP concentrations are higher within *Sargassum* windrows compared to adjacent waters^{5,39}. This localized enrichment has allowed *Sargassum* to exploit ammonium-rich excretions from associated fishes and invertebrates⁶, recycled nutrients from microbial mineralization of particulate organic matter⁴⁰ (POM) and dissolved organic N forms such as urea and amino acids⁴¹ through its long evolutionary history. While several forms of dissolved

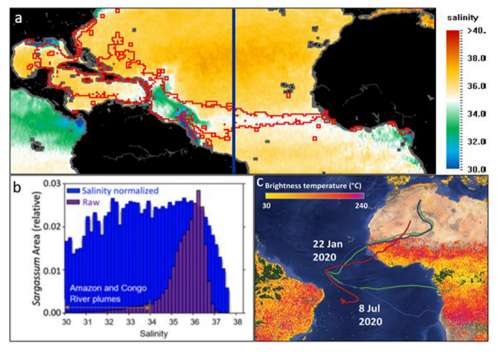


Fig. 4 Spatial distribution of *Sargassum* in relation to salinity and aerosol trajectories. a *Sargassum* distribution (red empty squares) overlaid on salinity derived from the Soil Moisture Ocean Salinity (SMOS) satellite mission, both for July 2018. The blue line marks the longitude of 38°W that the Amazon River plume hardly reaches. **b** Distribution of *Sargassum* as a function of salinity (purple) for 2011 to 2019, where statistics are calculated from 108 monthly mean maps over the cumulative *Sargassum* footprint. Here, the raw data (purple) shows *Sargassum* areal coverage in each salinity increment, relative to the total coverage (i.e., all purple bars sum to 1.0). The *Sargassum* coverage in each salinity increment is divided by the water area in the same salinity increment, resulting in the "salinity normalized" distribution (blue). The *Sargassum* distribution data were obtained on National Centers for Environmental Information (NCEI) Accession 0190272¹⁴. **c** Hybrid Single-Particle Lagrangian Integrated Trajectory model (HYSPLIT) air mass back trajectories (10-day) from 3°N, 34.6°W at 500 m (red), 1000 m (blue), and 1500 m (green), shown with the active fires in Africa and South America observed during Apr 2020 to Mar 2021 (from firms.modaps.eosdis.nasa.gov/download⁸⁵).

nitrogen are available to *Sargassum*, ammonium (NH_4^+) uptake is most efficient⁴².

More recently, the significant increase in tissue N (+35%) and upward shifts in N:P ratios (+111%) since the 1980s suggests that Sargassum is now exploiting the global trend in N enrichment. In the 1980s %N of Sargassum averaged 0.89% compared to higher, non-limiting values for macroalgae (>1.5%43) observed recently in the GOM, peninsular Florida, and the Amazon Plume (Fig. 3). Because of anthropogenic emissions of oxides of N (NO_x), the NO_x deposition rate is about fivefold greater than that of preindustrial times largely due to energy production and biomass burning⁴⁴. Production of synthetic fertilizer N has increased ninefold, while that of P has increased threefold since the 1980s³⁰ contributing to a global increase in N:P ratios. Notably, 85% of all synthetic N fertilizers have been created since 198545, which was shortly after the baseline Sargassum sampling began at Looe Key in 1983. The quantity of global N fixation for fertilizer production and P flowing into the oceans was estimated at 121 and 9.5 million tons/yr, respectively⁴⁶, yielding an anthropogenic N:P molar ratio of 28:1, identical to the mean N:P molar ratio of 28:1 measured in Sargassum since 2010.

A strong connection of *Sargassum* areal cover to land-based runoff is evidenced by the highest tissue %N values occurring in areas influenced by reduced salinity from river discharges and terrestrial runoff (Figs. 3a and 4a). Statistical analysis of the *Sargassum* cover in different salinity ranges from 2011 to 2019 shows that the bulk of *Sargassum* biomass occurs at oceanic salinities of ~36 (Fig. 4b). However, when *Sargassum* cover (or biomass) is normalized by water area in each salinity band (blue bars in Fig. 4b), the distribution is rather flat across the salinity range of 32 to 36 with 32.4 to 33.5 containing slightly higher abundance of Sargassum than 33.5 to 35, indicative of riverine influence. For waters with salinity <31, Sargassum abundance is lower, possibility due to the lower growth rate at low-salinity waters47. The Mississippi River48,49 and South Florida's Everglades and coastal urban belt^{50,51} have experienced trends of increasing N flux and increasing N:P ratios³⁵. For the GOM, the combined annual mean streamflow for the Mississippi and Atchafalaya rivers represents about 80% of the freshwater discharge to the GOM and accounts for 90% of total N load and 87% of the total P load discharged annually to the GOM⁵². Increasing nitrogen (mostly NO₃⁻) along with other nutrients are a cause of hypoxia in a large dead zone along the Louisiana-Texas coast^{53,54}, where the highest Sargassum tissue %N values (Fig. 3a) were observed in waters that tend to have lower salinity (Fig. 4a). In addition, the N:P ratio of the Mississippi River and northern GOM increased from 9 to 15 and 16 to 24, respectively, between 1960s and the 1980s⁵⁴, indicating that this stoichiometric shift began prior to the current study. A 30-year study between 1984 and 2014 at Looe Key reef showed over a twofold increase in seawater DIN and the DIN:SRP ratio, and threefold increase in tissue N:P ratio in a variety of reef macroalgae⁵⁰. Some of the Sargassum collections in the present study were made in blue water offshore of Looe Key reef and paralleled this pattern of tissue N enrichment and an increased N:P ratio from 11.2 in the 1980s to 24.2 since 2010.

The Amazon River is the largest river in the world and accounts for 20% of the world's total river discharges. Data from the 13 Carbon in the Amazon River Experiment (CAMREX) cruise surveys of the Amazon River between 1982 and 1991 show strong, statistically significant correlations between NO_3^- flux and discharge and between SRP flux and discharge (Fig. 5a). The

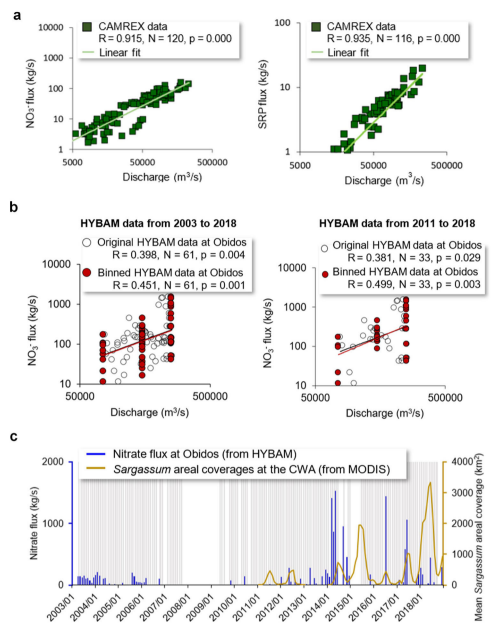


Fig. 5 Nutrient flux from the Amazon River and long-term *Sargassum* **trend. a** Nitrate (NO_3^-) and soluble reactive phosphorus (SRP) flux from all stations of the 13 CAMREX cruises⁸³ (1982 to 1991) are highly correlated with river discharge. These stations are within a 2000 km reach of the Brazilian Amazon River mainstream. Solid lines mark the power law regression lines. **b** NO_3^- flux at Obidos from the HYBAM database¹⁰⁰ is also correlated with discharge, and the correlation is higher if data are binned to different discharge groups. **c** NO_3^- flux at Obidos from the same HYBAM dataset shows apparent increases in recent years. The recent mean monthly *Sargassum* areal coverages obtained from Wang et al. (2019) in the Central West Atlantic (CWA, 0°N to 22°N, 63°W to 38°W) are also shown as reference. All months before 2011 show *Sargassum* coverages. The shaded gray bars indicate when quality controlled HYBAM nutrient flux measurements are available, but data in some months are too low (<0.1 mg/L) to be visible due to the scale.

monthly data at the Obidos station of the Amazon River from the Hydrology and Geochemistry of the Amazon basin (HYBAM) observatory also show a positive, statistically significant (P= 0.049) correlation between NO₃⁻ flux and discharge (Fig. 5b). There have been recent increases in NO₃⁻ fluxes at the Obidos station (Fig. 5b), especially between 2014 and 2016, with lower fluxes between 2016 and 2018 (Fig. 5c). Although a direct annual correspondence between NO₃⁻ flux and *Sargassum* amount is not apparent, the general trend of increased NO₃⁻ and SRP after 2014 suggests that river discharge may have supported *Sargassum* growth around the plume in subsequent years, especially 2015, 2017, and 2018. The HYBAM Obidos results are consistent with the limited field measurements of NO₃⁻ and SRP in the offshore

river plume (salinity 16 to 34) in 2010 and 2018^{15,55}, especially when compared with the lower historical nutrient values in the 1960s⁵⁶ and early 2000s⁵⁷. Such increases in nutrient fluxes could be partially due to Amazonian deforestation (+25% since 2010)¹⁵, which has been shown to alter the hydrochemical balance of streams and soil chemistry. In addition, extreme flooding events in the Amazon basin in this time frame⁵⁸ combined with increased fertilizer use (+67% since 2010)¹⁵ could also contribute. Although the Amazon River plume hardly reaches waters east of 38°W and therefore its contributions to *Sargassum* blooms in the eastern Tropical Atlantic can be neglected, its nutrient flux may have fueled recent blooms since 2014 in the central West Atlantic (Fig. 5c). The importance of the Amazon River nutrient flux to

Sargassum growth was also apparent in *S. natans* collected in plume waters (~10°N, 45.5°W; salinity ~33.8) in late August 2019 that averaged 2.4%N, a very high value that was exceeded only by *S. fluitans* collected in the Mississippi River plume off the Texas and west Florida coasts. Interestingly, while the overall effect of species was not significant (see Supplement), at this location *S. fluitans* was much less enriched in %N (mean = 0.66%) than *S. natans*. Considering that the Amazon Basin dominates P flux to the NA⁵⁹ and that land-based P exports can reach the open ocean⁶⁰ the productivity of *Sargassum* could be enhanced in this region of the western Tropical Atlantic by increased P availability. This is supported by the lowest N:P ratios observed in 2015 in Barbados in the present study (Fig. 3b), which is directly influenced by the Amazon River plume.

Seasonal changes in C:N:P contents of Sargassum closely matched patterns in nutrient flux from the Mississippi and Amazon rivers, further suggesting these river discharges support seasonal growth patterns. These river discharges increase from winter through spring and peak in early summer^{15,35,49} and could support the lower C:N and C:P ratios in Sargassum during this period; in contrast, the higher C:N and C:P ratios in summer and fall indicate nutrient limitation resulting from reduced river discharges^{5,6}. These data suggest seasonal nutrient control of bloom formation by seasonal river discharges, particularly in the GOM, where Sargassum cover expands in the spring and peaks in summer months (Supplementary Fig. 3)^{5,15,61}. Its air bladders allow Sargassum to float and form dense mats which are advected by ocean currents thus allowing episodic access to buoyant, lower salinity, nutrient-enriched river plumes, which can extend for thousands of kilometers from shore¹⁵ (Fig. 4a). Similar to the temperate kelps, Laminaria longicruris⁶² (synonym Saccharina latissimi) and Macrocystis pyrifera⁶³, the floating tropical Sargassum spp. appear to be responders rather than anticipators among macroalgae⁶⁴ by sequestering seasonally available N and P to support annual growth patterns. A seasonal NO₃⁻ related growth strategy occurs in the temperate kelp L. longicruris, which assimilates and stores NO₃⁻ in winter months when its available to support maximum growth rates in the spring and into July, after which tissue N and growth rate decline⁶².

The wide range of δ^{15} N values in *Sargassum* tissue from -2 to +8‰ reinforces previous suggestions that a variety of N sources support growth of Sargassum over its broad geographic range^{5,15,65}. The δ^{15} N values of wet atmospheric deposition across the United States are relatively low, ranging from -11 to +3.5‰ with a median value of -3.1% (*n* = 883)⁶⁶ and are within the low end of the range of $\delta^{15}N$ in Sargassum. Similarly, in Bermuda, rainwater $NH_4^+ \delta^{15}N$ values ranged from -12.5 to +0.7‰67. Synthetic fertilizer N, which has increased ninefold since the 1980s, has δ^{15} N values ranging from -2 to $+2\%^{68}$ and is the mid-range for most Sargassum values in this study. More enriched values of +2.5 to +4.8‰ are indicative of upwelled NO_3^- in the upper 200 m of the NA^{69,70}. Higher values (+3 to +20‰) are indicative of urban wastewater from terrestrial runoff where fractionation associated with volatilization of NH₄⁺ and denitrification of NO₃⁻ occur⁷¹. This enrichment of Sargassum tissue is evident in the highest δ^{15} N values ranging between +3 to +8‰ along urbanized coastal waters in Texas and Florida, illustrating the effect of anthropogenic nitrogen enrichment. The mean δ^{15} N value of POM in the Mississippi River is ~+7‰⁷² and δ^{15} N enrichment of Sargassum by +2‰ has been reported for neritic compared to oceanic regions in the GOM⁶⁵, as well as macroalgal blooms on coral reefs downstream of sewage outfalls in South Florida (+6 to +8‰)⁵¹. In the Amazon River floodplain, δ^{15} N of phytoplankton and macrophytes range from +4.7 to +5.5% respectively⁷³, indicating that the plume could contribute to δ^{15} N enrichment of Sargassum. These findings suggest

that episodic N enrichment in highly populated tourist areas of the Caribbean could help sustain *Sargassum* growth and bloom continuation.

Natural N and P sources, such as upwelling and N₂ fixation, could further support Sargassum growth and would be especially important in offshore and oceanic locations of the GASB¹³. Upwelling occurs at the shelf break in the southeastern United States^{74,75}, and in the eastern equatorial Atlantic⁷⁶ and could supply NO₃⁻ to Sargassum. N₂ fixation by the cyanobacterial epiphyte Dichothrix fucicola occurs in Sargassum windrows and can provide from 2 to 32% of the N needs^{77,78}, which would result in δ^{15} N values close to 0‰. N₂ fixation plays a prominent role in N-nutrient cycling in the Amazon and Congo river plumes^{57,79}. N₂ fixation by diatom diazotroph associations (DDAs) and Trichodesmium support 11% of total primary production in the mesohaline section of the Amazon River plume (salinity ~ 32 to 33)⁵⁷. During a study in which the plume extended into the Caribbean Sea, diazotrophy by DDAs supplied ~25% of water column N demand⁸⁰. In the eastern equatorial Atlantic (Gulf of Guinea), N₂ fixation rates were 2 to 7 times higher when upwelling occurred as compared to non-upwelling conditions, as a result of low NO3:SRP ratios in upwelled waters that leave excess SRP that stimulates N₂ fixation⁷⁶. Considering the high N:P ratio of Sargassum that now occurs in the NA basin, such excess SRP in the upwelled water could stimulate growth of Sargassum in the eastern Tropical Atlantic.

Atmospheric deposition (dry and wet) of lithogenic and anthropogenic-sourced aerosols can supply the central Atlantic with major and trace nutrients that could further support Sargassum growth. Aerosol back trajectories (examples shown in Fig. 4c) show that winds over the central Atlantic change seasonally but are predominantly from northern Africa, where Saharan dust originates, and central and southern Africa, where biomass burning generates anthropogenic-type aerosols^{19,81}. The largest atmospheric supply of nutrients, like Fe and P, comes from seasonal Saharan dust plumes, but the low solubility of these elements in mineral dust limits their bioavailability⁸²⁻⁸⁴. In contrast, biomass burning in central and southern Africa (as shown as active fires in Fig. $4c^{85-87}$) can deliver nutrients like N, P, and Fe to the central Atlantic. Like wind patterns, biomass burning also varies seasonally with the Northern (Southern) Hemisphere burn season occurring in November to March (May to October)88. While nutrient concentrations in aerosols produced during biomass burning might be lower than in Saharan dust, the nutrient solubilities are higher, potentially providing a source of more bioavailable nutrients to the Atlantic surface ocean^{83,89–91}. Fluxes of aerosols from both dust and biomass burning appear to have increased over the past 150 years, according to models of past, present, and future changes to the atmospheric deposition in the North and Central Atlantic⁹². While the fluxes of dissolved P to the NA have likely increased since 1850, which would alleviate P limitation, the fluxes of dissolved Fe have increased faster resulting in higher dissolved Fe/ dissolved P ratios^{91,92}, which could enhance N₂ fixation and balance increased inputs of bioavailable P. Future atmospheric inputs are difficult to predict as aerosol production and processing rely heavily on economic and human factors, including restrictions on biomass burning, fossil fuel consumption, and industrial pollution. Nonetheless, seasonal inputs of natural and anthropogenic-driven aerosol nutrients could at least partially alleviate P and/or Fe limitation, resulting in increased Sargassum growth and abundance across the central Atlantic Ocean.

Almost 50 years ago, scientists recognized that nutrient addition through use of fertilizers can destabilize food webs, leading to loss of biodiversity and ecosystem function through the so called paradox of enrichment^{93,94}. During that time, global river discharges showed a trend of decreasing N:P due to human activities and P was considered the primary limiting nutrient in surface waters⁹⁵. Concurrently, the precept that N, rather than P, was driving marine eutrophication was introduced to the scientific community³³. Since then, N, Fe, and silica have been widely considered to be the most important nutrients that limit phytoplankton growth in the oceans, although mounting evidence is supporting an emerging paradigm in oceanography that P plays a primary role in the Atlantic basin^{96–98}. Recent reviews now show that N:P ratios of rivers are increasing, despite attempts to mitigate application of N fertilizers³⁵. The empirical data presented here for Sargassum supports not only a primary role for P limitation of productivity, but also suggests that the role of P as a limiting nutrient is being strengthened by the relatively large increases in anthropogenic N supply from terrestrial runoff, atmospheric inputs, and possibly other natural sources such as N2 fixation⁹⁶. The increased P limitation in Sargassum could be compensated for by its relatively high capacity for alkaline phosphatase activity, which allows it to sequester SRP from dissolved organic P compounds, a physiological characteristic of adaptive value to growth in oligotrophic waters⁵. Considering the negative effects that the GASB is having on the coastal communities of Africa, the Caribbean, GOM, and South Florida, more research is urgently needed to better inform societal decisionmaking regarding mitigation and adaptation of the various terrestrial, oceanic, and atmospheric drivers of the Sargassum blooms.

Methods

Sample collection. Sargassum samples in the 1980s were collected mostly from University-National Oceanographic Laboratory System research vessels, including the R/V Columbus Iselin (Loop Current, Gulf Stream, Sargasso Sea), R/V Calanus (Belize), RV Cape Hatteras (Sargasso Sea, Gulf Stream, Belize), and R/V Weatherbird (Sargasso Sea); for blue waters offshore Looe Key in the lower Florida Keys, Sargassum was collected from a small boat (20' Mako). Since 2010, Sargassum has been collected from the R/V Point Sur (GOM) and the R/V Thomas G. Thompson (Amazon plume). Other samples were collected by volunteers on private vessels and the M/V Ocearch (Gulfstream). Windrows of Sargassum spp., which result from Langmuir circulation that aligns Sargassum parallel with the wind direction, were frequently encountered at various locations during the research cruises. For all sampling events, Sargassum spp. were collected from small boats either by divers or with a dip net and sorted into the species and morphotypes, S. natans I and S. fluitans III per Parr (1939)⁴. After collection, the plants were placed in clean plastic bags in a cooler. Upon return to the lab or research vessel, the samples were separated into replicate (n = 2 to 3/species for each location and sampling) composite samples (6 to 10 thalli/species), rinsed briefly (3 to 5 s) in deionized water, cleaned of macroscopic epizoa and epiphytes, dried in a laboratory oven at 65 to 70 °C for 48 h, and powdered with a mortar and pestle⁵

For both the 1980s and 2010s tissue analysis, total C and N were determined on a Carlo-Erba CHN Combustion Analyzer, while total P was determined by persulfate digestion followed by analysis for SRP using either a Bausch and Lomb Spectronic 88 or an Alpkem 300 series autoanalyzer. The resulting tissue %C, %N, and %P data were used to calculate molar C:N:P ratios. Additional analysis of 2010 tissue (427) δ^{15} N was conducted on a Thermo Delta V IRMS coupled to a Carlo Erba NA1500 CHN-Combustion Analyzer via a Thermo Conflo III Interface.

Statistical analysis. The relationship between *S. fluitans* and *S. natans* elemental composition (%C, %N, %P) and molar ratios (C:N:P) with species, decade, and season were analyzed using multivariate and subsequent univariate general linear models (MANOVA and ANOVA) in Minitab 19 Statistical Software. All variables were non-normal therefore log transformation was attempted prior to analyses and model fit was assessed through examination of residuals. While log transformation improved the normality, shape, and residual distribution of %N, %P, C:N, N:P, and C:P, %C was not improved and thus the raw values for this parameter were used in analyses. Significant univariate factors and interactions were assessed with Tukey's pairwise comparisons. To better understand nitrogen sources supporting *Sargassum* bloom growth and development, %N, N:P ratios, and δ^{15} N of post-2010 samples were compared by location with ANOVA using similar methods as above in Minitab 19 Statistical Software. Statistical significance was considered at *P* < 0.05 for all analyses.

Satellite-measured distributions of *Sargassum* and salinity and fieldmeasured river nutrient concentrations. Pelagic *Sargassum* distributions covering the GOM and central Atlantic Ocean were derived from MODIS measurements and the data were acquired from NCEI Accession 0190272(¹⁵). Surface salinity distributions were obtained from SMOS Earth Explorer mission and the data were accessed on https://www.catds.fr/Products/Available-products-from-CPDC). Amazon River discharge and water chemistry data, including NO₃⁻⁻, and PO₄³⁻⁻ concentrations, measured at the Obidos station during 2003 to 2018 were downloaded from HYBAM database (http://www.ore-hybam.org/index.php/eng/Data). Similar parameters were also from 1981 to 1991 collected during cruises for the CAMREX⁹⁹ were also analyzed.

Data availability

All data used in this study are available in the main text or the supplemental materials. The raw data that support the findings of this study are available from the corresponding author upon reasonable request. All *Sargassum*-relevant imagery data products are available through the *Sargassum* Watch System (SaWS, https://optics.marine.usf.edu/projects/saws.html).

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References

- 1. Ryther, J. H. The Sargasso Sea. Sci. Am. 194, 98-108 (1956).
- 2. Littler, D. S. & Littler, M. M. Caribbean Reef Plants (Offshore Graphics, 2000).
- 3. Winge, O. The Sargasso Sea, Its Boundaries and Vegetation In *Report of the Danish Oceanographic Expedition*, Vol. III, 1908–1910, (Copenhagen: Andr. Fred. Hòst and Sòn) 34 pp. Miscellaneous Paper Number 2. (1923).
- Parr, A. E. Quantitative observations on the pelagic Sargassum vegetation of the western North Atlantic. Bull. Bingham Oceanogr. Collect. 6, 1–94 (1939).
- Lapointe, B. E. A comparison of nutrient-limited productivity in Sargassum natans from neritic vs. oceanic waters of the western North Atlantic Ocean. Limnol. Oceanogr. 40, 625–633 (1995).
- Lapointe, B. E., West, L. E., Sutton, T. T. & Hu, C. Ryther revisited: nutrient excretions by fishes enhance productivity of pelagic *Sargassum* in the western North Atlantic Ocean. J. Exp. Mar. Biol. Ecol. 458, 46–56 (2014).
- Gower, J., Hu, C., Borstad, G. & King, S. Ocean color satellites show extensive lines of floating *Sargassum* in the Gulf of Mexico. *IEEE Trans. Geosci. Remote Sens.* 44, 3619–3625 (2006).
- Williams, A., Feagin, R. & Stafford, A. W. Environmental impacts of beach raking of *Sargassum spp.* on Galveston Island, TX. *Shore Beach* 76, 63–69 (2008).
- 9. Moritsugu, K. Tampa Bay Times (Times Publishing Company, 1991).
- Turner, R. E. & Rabalais, N. N. Coastal eutrophication near the Mississippi river delta. *Nature* 368, 619–621 (1994).
- Gower, J. F. R. & King, S. A. Distribution of floating *Sargassum* in the Gulf of Mexico and the Atlantic Ocean mapped using MERIS. *Int. J. Remote Sens.* 32, 1917–1929 (2011).
- Johnson, D. R., Ko, D. S., Franks, J. S., Moreno, P. & Sanchez-Rubio, G. The Sargassum invasion of the Eastern Caribbean and dynamics of the Equatorial North Atlantic. In Proceedings of the 65th Annual Gulf and Caribbean Fisheries Institute Conference pp. 102–103 (2013). http://aquaticcommons.org/ 21444/1/GCFI_65-17.pdf.
- Gower, J., Young, E. & King, S. Satellite images suggest a new Sargassum source region in 2011. Remote Sens. Lett. 4, 764–773 (2013).
- Johns, E. M. et al. The establishment of a pelagic Sargassum population in the tropical Atlantic: biological consequences of a basin-scale long distance dispersal event. Prog. Oceanogr. 182, 102269–102269 (2020).
- 15. Wang, M. et al. The great Atlantic Sargassum belt. Science 364, 83-87 (2019).
- Djakouré, S., Araujo, M., Hounsou-Gbo, A., Noriega, C. & Bourlès, B. On the potential causes of the recent Pelagic *Sargassum* blooms events in the tropical North Atlantic Ocean. *Biogeosci. Discuss.* https://doi.org/10.5194/bg-2017-346 (2017).
- Oviatt, C. A., Huizenga, K., Rogers, C. S. & Miller, W. J. What nutrient sources support anomalous growth and the recent *Sargassum* mass stranding on Caribbean beaches? A review. *Mar. Pollut. Bull.* 145, 517–525 (2019).
- McGillicuddy, D. J., Jr, Anderson, L. A., Doney, S. C. & Maltrud, M. E. Eddydriven sources and sinks of nutrients in the upper ocean: results from a 0.1 resolution model of the North Atlantic. *Global Biogeochem. Cycles* 17, 1035 (2003).
- Barkley, A. E. et al. African biomass burning is a substantial source of phosphorus deposition to the Amazon, Tropical Atlantic Ocean, and Southern Ocean. *Proc. Natl Acad. Sci. USA* 116, 16216–16221 (2019).
- Qi, L., Hu, C., Xing, Q. & Shang, S. Long-term trend of *Ulva prolifera* blooms in the western Yellow Sea. *Harmful Algae* 58, 35–44 (2016).
- Qi, L., Hu, C., Wang, M., Shang, S. & Wilson, C. Floating algae blooms in the East China Sea. *Geophys. Res. Lett.* 44, 501–511,509 (2017).

- Smetacek, V. & Zingone, A. Green and golden seaweed tides on the rise. Nature 504, 84–88 (2013).
- Van Tussenbroek, B. I. et al. Severe impacts of brown tides caused by Sargassum spp. on near-shore Caribbean seagrass communities. Mar. Pollut. Bull. 122, 272–281 (2017).
- Alvarez-Filip, L., Estrada-Saldívar, N., Pérez-Cervantes, E., Molina-Hernández, A. & González-Barrios, F. J. A rapid spread of the stony coral tissue loss disease outbreak in the Mexican Caribbean. *PeerJ* 7, e8069–e8069 (2019).
- Cabanillas-Terán, N., Hernández-Arana, H. A., Ruiz-Zárate, M.-Á., Vega-Zepeda, A. & Sanchez-Gonzalez, A. Sargassum blooms in the Caribbean alter the trophic structure of the sea urchin Diadema antillarum. PeerJ 7, e7589–e7589 (2019).
- Maurer, A. S., De Neef, E. & Stapleton, S. Sargassum accumulation may spell trouble for nesting sea turtles. Front. Ecol. Environ. 13, 394–395 (2015).
- 27. Webster, R. K. & Linton, T. Development and implementation of Sargassum early advisory system (SEAS). Shore Beach 81, 1-1 (2013).
- Resiere, D. et al. Sargassum seaweed on Caribbean islands: an international public health concern. Lancet 392, 2691–2691 (2018).
- 29. Glibert, P. et al. The role of in the global proliferation of harmful algal blooms: new perspectives and approaches. *Oceanography* **18**, 196–207 (2005).
- Glibert, P. M. Eutrophication, harmful algae and biodiversity Challenging paradigms in a world of complex nutrient changes. *Mar. Pollut. Bull.* 124, 591–606 (2017).
- Steffen, W. et al. Planetary boundaries: guiding human development on a changing planet. *Science* 347, 6223 https://doi.org/10.1126/science.1259855 (2015).
- Ryther, J. H. The ecology of phytoplankton blooms in Moriches bay and Great South bay, Long Island, New York. *Biol. Bull.* 106, 198–209 (1954).
- 33. Ryther, J. H. & Dunstan, W. M. Nitrogen, Phosphorus, and Eutrophication in the coastal marine environment. *Science* **171**, 1008 LP-1013 (1971).
- Howarth, R. W. & Marino, R. Nitrogen as the limiting nutrient for eutrophication in coastal marine ecosystems: evolving views over three decades. *Limnol. Oceanogr.* 51, 364–376 (2006).
- Oelsner, G. P. & Stets, E. G. Recent trends in nutrient and sediment loading to coastal areas of the conterminous U.S.: insights and global context. *Sci. Total Environ.* 654, 1225–1240 (2019).
- Falkowski, P. G. Evolution of the nitrogen cycle and its influence on the biological sequestration of CO2 in the ocean. *Nature* 387, 272–275 (1997).
- 37. Tyrrell, T. The relative influences of nitrogen and phosphorus on oceanic primary production. *Nature* **400**, 525–531 (1999).
- Lapointe, B. E., Littler, M. M. & Littler, D. S. A comparison of nutrient-limited productivity in macroalgae from a Caribbean barrier reef and from a mangrove ecosystem. *Aquat. Bot.* 28, 243–255 (1987).
- Culliney, J. L. Measurements of reactive phosphorus associated with pelagic Sargassum in the Northwest Sargasso Sea1. Limnol. Oceanogr. 15, 304–305 (1970).
- Schaffelke, B. Particulate organic matter as an alternative nutrient source for tropical Sargassum species (Fucales, Phaeophyceae). J. Phycol. 35, 1150–1157 (1999).
- Vonk, J. A., Middelburg, J. J., Stapel, J. & Bouma, T. J. Dissolved organic nitrogen uptake by seagrasses. *Limnol. Oceanogr.* 53, 542–548 (2008).
- Han, T., Qi, Z., Huang, H., Liao, X. & Zhang, W. Nitrogen uptake and growth responses of seedlings of the brown seaweed *Sargassum hemiphyllum* under controlled culture conditions. *J. Appl. Phycol.* **30**, 507–515 (2018).
- Fujita, R., Wheeler, P. & Edwards, R. Assessment of macroalgal nitrogen limitation in a seasonal upwelling region. *Mar. Ecol. Prog. Ser.* 53, 293–303 (1989).
- Prospero, J. M. et al. in Nitrogen Cycling in the North Atlantic Ocean and its Watersheds (ed. Robert, W. H.) (Springer, 1996).
- 45. Howarth, R. W. Coastal nitrogen pollution: a review of sources and trends globally and regionally. *Harmful Algae* **8**, 14–20 (2008).
- Rockström, J. & Karlberg, L. The quadruple squeeze: defining the safe operating space for freshwater use to achieve a triply green revolution in the Anthropocene. *Ambio* 39, 257–265 (2010).
- 47. Hanisak, M. D. & Samuel, M. A. Twelfth International Seaweed Symposium (Springer, 1986).
- Rabalais, N. N. et al. Hypoxia in the northern Gulf of Mexico: does the science support the plan to reduce, mitigate, and control hypoxia? *Estuar. Coasts* 30, 753–772 (2007).
- Tian, H. et al. Long-term trajectory of nitrogen loading and delivery from Mississippi river basin to the Gulf of Mexico. *Glob. Biogeochem. Cycles* 34, e2019GB006475–e002019GB006475 (2020).
- Lapointe, B. E., Brewton, R. A., Herren, L. W., Porter, J. W. & Hu, C. Nitrogen enrichment, altered stoichiometry, and coral reef decline at Looe Key, Florida Keys, USA: a 3-decade study. *Mar. Biol.* 166, 108–108 (2019).
- Lapointe, B. E., Barile, P. J. & Littler, M. M. & Littler, D. S. Macroalgal blooms on southeast Florida coral reefs: II. Cross-shelf discrimination of nitrogen

sources indicates widespread assimilation of sewage nitrogen. *Harmful Algae* 4, 1106–1122 (2005).

- Dunn, D. E. Trends in Nutrient Inflows to the Gulf of Mexico from Streams Draining the Conterminous United States, 1972-93. Report No. 96-4113 (Austin, TX, 1996).
- Turner, R. E. & Rabalais, N. N. Changes in Mississippi River water quality this century: implications for coastal food webs. *Bioscience* 41, 140–147 (1991).
- 54. Rabalais, N. N. et al. Nutrient changes in the Mississippi River and system responses on the adjacent continental shelf. *Estuaries* **19**, 386-407 (1996).
- Weber, S. C. et al. Amazon River influence on nitrogen fixation and export production in the western tropical North Atlantic. *Limnol. Oceanogr.* 62, 618–631 (2017).
- Ryther, J. H., Menzel, D. W. & Corwin, N. Influence of Amazon River outflow on ecology of Western Tropical Atlantic. I. Hydrography and nutrient chemistry. J. Mar. Res. 25, 69–69 (1967).
- Subramaniam, A. et al. Amazon River enhances diazotrophy and carbon sequestration in the tropical North Atlantic Ocean. *Proc. Natl Acad. Sci.USA* 105, 10460 LP-10410465 (2008).
- Barichivich, J. et al. Recent intensification of Amazon flooding extremes driven by strengthened Walker circulation. *Sci. Adv.* 4, eaat8785–eaat8785 (2018).
- Howarth, R. W. et al. Regional nitrogen budgets and riverine N & P fluxes for the drainages to the North Atlantic Ocean: Natural and human influences. In *Nitrogen Cycling in the North Atlantic Ocean and its Watersheds* (ed. Robert, W. Howarth) (Springer, Dordrecht, 1996). https://doi.org/10.1007/978-94-009-1776-7_3.
- Galloway, J. N. et al. Regional nitrogen budgets and riverine N & P fluxes for the drainages to the North Atlantic Ocean: Natural and human influences. *Biogeochemistry* (ed. Robert, W. Howarth) 35, 181–226 (Springer, 1996).
- Gower, J. & King, S. Satellite images show the movement of floating Sargassum in the Gulf of Mexico and Atlantic Ocean. Nat. Preced. https://doi. org/10.1038/npre.2008.1894.1 (2008).
- Chapman, A. R. O. & Craigie, J. S. Seasonal growth in Laminaria longicruris: relations with dissolved inorganic nutrients and internal reserves of nitrogen. *Mar. Biol.* 40, 197–205 (1977).
- Zimmerman, R. C. & Kremer, J. N. Episodic nutrient supply to a kelp forest ecosystem in Southern California. J. Mar. Res. 42, 591–604 (1984).
- 64. Kain, J. M. The seasons in the subtidal. Br. Phycol. J. 24, 203-215 (1989).
- Dorado, S., Rooker, J. R., Wissel, B. & Quigg, A. Isotope baseline shifts in pelagic food webs of the Gulf of Mexico. *Mar. Ecol. Prog. Ser.* 464, 37–49 (2012).
- Kendall, C., Elliott, E. M. & Wankel, S. D. Wiley Online Books 375-449 (2007).
 Altieri, K. E., Hastings, M. G., Peters, A. J., Oleynik, S. & Sigman, D. M. Isotopic evidence for a marine ammonium source in rainwater at Bermuda.
- Glob, Biogeochem, Cycles 28, 1066–1080 (2014).
 Bateman, A. S. & Kelly, S. D. Fertilizer nitrogen isotope signatures. *Isotopes*
- Bateman, A. S. & Kelly, S. D. Fertilizer nitrogen isotope signatures. *Isotopes Environ. Health Stud.* 43, 237–247 (2007).
- Knapp, A. N., DiFiore, P. J., Deutsch, C., Sigman, D. M. & Lipschultz, F. Nitrate isotopic composition between Bermuda and Puerto Rico: implications for N2 fixation in the Atlantic Ocean. *Global Biogeochem. Cycles* 22, GB3014 https://doi.org/10.1029/2007GB003107 (2008).
- Knapp, A. N., Sigman, D. M. & Lipschultz, F. N isotopic composition of dissolved organic nitrogen and nitrate at the Bermuda Atlantic Time-series Study site. *Global Biogeochem. Cycles* 19, GB1018 https://doi.org/10.1029/ 2004GB002320 (2005).
- Montoya, J. P. Nitrogen stable isotopes in marine environments. Nitrogen Mar. Environ. 2, 1277–1302 (2008).
- Wissel, B. & Fry, B. Sources of particulate organic matter in the Mississippi River, USA. *Large Rivers* 15 105–118 (2003).
- Zaia Alves, G. H., Hoeinghaus, D. J., Manetta, G. I. & Benedito, E. Dry season limnological conditions and basin geology exhibit complex relationships with δ13C and δ15N of carbon sources in four Neotropical floodplains. *PLoS ONE* 12, e0174499 (2017).
- Smith, N. P. Upwelling in Atlantic shelf waters of South Florida. *Florida Scientist* 45, 125–138 (1982).
- Atkinson, L. P., O'Malley, P. G., Yoder, J. A. & Paffenhöfer, G. A. The effect of summertime shelf break upwelling on nutrient flux in southeastern United States continental shelf waters. *J. Mar. Res.* 42, 969–993 (1984).
- Subramaniam, A., Mahaffey, C., Johns, W. & Mahowald, N. Equatorial upwelling enhances nitrogen fixation in the Atlantic Ocean. *Geophys. Res. Lett.* 40, 1766–1771 (2013).
- Carpenter, E. J. Nitrogen fixation by a blue-green epiphyte on Pelagic Sargassum. Science 178, 1207–1209 (1972).
- Phlips, E. J., Willis, M. & Verchick, A. Aspects of nitrogen fixation in Sargassum communities off the coast of Florida. J. Exp. Mar. Biol. Ecol. 102, 99–119 (1986).
- 79. Subramaniam, A., Montoya, J. P., Foster, R. A. & Capone, D. G. Nitrogen fixation in the eastern equatorial Atlantic: who and how much? *European Geosciences Union General Assembly* **11**, 10156–10156 (2009).

ARTICLE

- Carpenter, E. J. et al. Extensive bloom of a N2-fixing diatom/cyanobacterial association in the tropical Atlantic Ocean. *Mar. Ecol. Prog. Ser.* 185, 273–283 (1999).
- Zubkova, M., Boschetti, L., Abatzoglou, J. T. & Giglio, L. Changes in fire activity in Africa from 2002 to 2016 and their potential drivers. *Geophys. Res. Lett.* 46, 7643–7653 (2019).
- Baker, A. R., French, M. & Linge, K. L. Trends in aerosol nutrient solubility along a west-east transect of the Saharan dust plume. *Geophys. Res. Lett.* 33 L07805, https://doi.org/10.1029/2005GL024764 (2006).
- Baker, A. R., Jickells, T. D., Witt, M. & Linge, K. L. Trends in the solubility of iron, aluminium, manganese and phosphorus in aerosol collected over the Atlantic Ocean. *Mar. Chem.* 98, 43–58 (2006).
- Shelley, R. U., Morton, P. L. & Landing, W. M. Elemental ratios and enrichment factors in aerosols from the US-GEOTRACES North Atlantic transects. *Deep Sea Res. Part II Top. Stud. Oceanogr.* 116, 262–272 (2015).
- Giglio, L., Schroeder, W. & Justice, C. O. The collection 6 MODIS active fire detection algorithm and fire products. *Remote Sens. Environ.* 178, 31–41 (2016).
- Giglio, L., Descloitres, J., Justice, C. O. & Kaufman, Y. J. An enhanced contextual fire detection algorithm for MODIS. *Remote Sens. Environ.* 87, 273–282 (2003).
- Giglio, L., van der Werf, G. R., Randerson, J. T., Collatz, G. J. & Kasibhatla, P. Global estimation of burned area using MODIS active fire observations. *Atmos. Chem. Phys.* 6, 957–974 (2006).
- Roberts, G., Wooster, M. J. & Lagoudakis, E. Annual and diurnal african biomass burning temporal dynamics. *Biogeosciences* 6, 849–866 (2009).
- Baker, A. R. & Jickells, T. D. Atmospheric deposition of soluble trace elements along the Atlantic Meridional Transect (AMT). *Prog. Oceanogr.* 158, 41–51 (2017).
- Chance, R., Jickells, T. D. & Baker, A. R. Atmospheric trace metal concentrations, solubility and deposition fluxes in remote marine air over the south-east Atlantic. *Mar. Chem.* 177, 45–56 (2015).
- Myriokefalitakis, S., Nenes, A., Baker, A. R., Mihalopoulos, N. & Kanakidou, M. Bioavailable atmospheric phosphorous supply to the global ocean: a 3-D global modeling study. *Biogeosciences* 13, 6519–6543 (2016).
- Kanakidou, M., Myriokefalitakis, S. & Tsigaridis, K. Aerosols in atmospheric chemistry and biogeochemical cycles of nutrients. *Environ. Res. Lett.* 13, 063004 (2018).
- Rosenzweig, M. L. Paradox of enrichment: destabilization of exploitation ecosystems in ecological time. *Science* 171, 385–387 (1971).
- McCann, K. S. et al. Landscape modification and nutrient-driven instability at a distance. *Ecol. Lett.* 24, 398–414 (2021).
- Meybeck, M. Carbon, nitrogen, and phosphorus transport by world rivers. Am. J. Sci. 282, 401–450 (1982).
- Fanning, K. A. Nutrient provinces in the sea: concentration ratios, reaction rate ratios, and ideal covariation. J. Geophys. Res. Oceans 97, 5693–5712 (1992).
- Ammerman, J. W., Hood, R. R., Case, D. A. & Cotner, J. B. Phosphorus deficiency in the Atlantic: an emerging paradigm in oceanography. *Eos, Trans. Am. Geophys. Union* 84, 165–170 (2003).
- Lomas, M. W., Bonachela, J. A., Levin, S. A. & Martiny, A. C. Impact of ocean phytoplankton diversity on phosphate uptake. *Proc. Natl Acad. Sci. USA* 111, 17540–17545 (2014).
- 99. Richey, J. E. et al. (ORNL Distributed Active Archive Center, 2008).
- 100. Cochonneau, G. et al. The environmental observation and research project, ORE HYBAM, and the rivers of the Amazon basin. In *Climate Variability and Change—Hydrological Impacts* (eds Demuth, S. et al.) vol. 308, 44–50 (2006).

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Author contributions

B.E.L. was responsible for collection and analysis of the *Sargassum* tissue, completed first draft of this manuscript, led data interpretation, and conceived the project. R.A.B. contributed to data analysis and interpretation, geospatial analyses, and manuscript editing. L.W.H. contributed to geospatial analyses. M.W contributed to *Sargassum* collections, data acquisition, and analyses C.H. contributed to project conception, data acquisition, analyses, and interpretation. D.J.M. contributed data acquisition, analyses, and interpretation. F.J.H. contributed data acquisition, analyses, and interpretation. F.L.M. contributed data acquisition, analyses, and interpretation.

Competing interests

The authors declare no competing interests.

Additional information

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