The nitrogen bioextraction potential of nearshore *Saccharina latissima* cultivation and harvest in the Western Gulf of Maine

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Received: 11 January 2020 / Revised and accepted: 3 January 2021 / Published online: 20 February 2021
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Abstract

In-water remediation strategies, implemented in conjunction with traditional watershed management, could help minimize the impact of excess nitrogen (N) on marine ecosystems. Seaweed farming and harvesting may have potential as in-water N remediation tools in the Western Gulf of Maine (WGoM), but more understanding of the associated spatial and temporal variability is needed. In this study, *Saccharina latissima* was grown and collected from four WGoM sites in 2016–2019 and analyzed for tissue N content and stable isotopes. The source of N taken by the kelp was not obvious from monthly nor interannual mean $\delta^{15}$N measured in the kelp tissue, and the interannual means were significantly different between sites in the same bay. Mean kelp biomass across all sites and years was 9.84 (± 2.53)–14.84 kg (wet weight) per meter of longline at time of harvest (late May–early June). Nitrogen content of the *S. latissima* tissue was 1.04–3.82% (± 0.22) (dry weight) throughout the growing season and generally decreased through the spring. Using these results, we estimated that harvesting a hypothetical hectare of *S. latissima* after 6–7 months of cultivation in the WGoM would have the potential to remove 19.2 (± 4.8)–176.0 (± 7.7) kg N ha$^{-1}$, depending on the density of longlines. The wide ranges of both biomass at time of harvest, and $\delta^{15}$N and percent N content in the kelp tissue, highlight the need for site-specific pilot studies, even within a specific bay, prior to implementing kelp aquaculture as an in-water tool for N bioextraction.

Keywords Phaeophyceae • Kelp • Mariculture • Nutrients • Ecosystem services • Bioremediation

Introduction

Nutrient pollution is one of the principal causes of poor coastal water quality and habitat degradation (Nixon 1995, 1998; Diaz and Rosenberg 2008; Paerl et al. 2014). Globally, an estimated 245,000 km of coastline are considered “dead zones” triggerd by excessive input of reactive nitrogen (N) and phosphorus (P) (Diaz and Rosenberg 2008). In the United States of America (USA), a nationwide excess of reactive N from anthropogenic sources has caused impairment to an estimated two-thirds of the country’s coastal waters (Bricker et al. 1999; Howarth et al. 2002). Moreover, the degree of coastal nutrient loading to the Northeastern USA coastline is considered one of the highest on Earth (Boesch 2002; Howarth 2008). Nutrient pollution, in combination with other trace elements supporting primary production, results in areas of hypoxia and anoxia, habitat degradation, altered food webs, loss of biodiversity, increased instances of green or harmful algal blooms, and greater susceptibility to localized ocean acidification (Nixon 1987, 1995; Paerl 1997; Paerl and Whitall 1999; Breitburg et al. 2009, 2018; Wallace et al. 2014).

In this study we focus on nutrient concerns in the Western Gulf of Maine (WGoM) bordering Massachusetts (MA), New Hampshire (NH), and Maine (ME), USA. Bays and estuaries adjacent to these states are waterbodies of emerging concern due to both point and nonpoint sources of reactive N (Castro et al. 2003; Liebman et al. 2012). Effluent from wastewater treatment facilities (WWTFs) is the most common point source of N to the WGoM; however, substantial N contributions from nonpoint sources like stormwater runoff, agricultural runoff, and atmospheric deposition also occur in the region (Castro et al. 2003; Liebman et al. 2012; Trowbridge et al. 2014). Atmospheric N deposition is estimated to be 30–40% of the total N load in many locations and stormwater runoff has been estimated to contribute another 30–35% of the nonpoint source...
N loading (Castro et al. 2003; Liebman et al. 2012; Trowbridge et al. 2014). New Hampshire and Massachusetts have implemented N discharge limits and strategies targeting both point and nonpoint source N to address and minimize the deleterious effects of excess nutrients on the WGoM (Reitsma et al. 2017). Maine, the state with the most coastline bordering the WGoM, has yet to establish nutrient criteria.

In addition to improving point source discharges, resource managers in Maine are interested in nutrient bioextraction as part of a system-wide approach integrating watershed load reductions and enhanced nutrient assimilation (Liebman et al. 2012). Nutrient bioextraction strategies, also referred to as bioremediation, aim to remove nutrients that exceed the flushing and assimilation capacity of the system, regardless of their source (Krom 1986; Chopin et al. 2001; Neori et al. 2004). Bioextraction efforts in coastal water bodies typically target dissolved inorganic nitrogen (DIN) because it often limits primary production in temperate marine ecosystems (Ryther and Dunstan 1971; Lobban and Harrison 1994). Excess dissolved inorganic P and dissolved carbon (C), and small amounts of dissolved organic N and P when inorganic nutrient levels are low (Li et al. 2016), are also removed from the environment during bioextraction (Bianchi 2007).

Many primary producers are suitable for use in bioextraction, but recently more attention has been given to the use of macroalgae in this role. Macroalgae naturally extract N from the marine environment because N is one of the key macronutrients required for protein and nucleic acid synthesis; and kelps are highly productive (Gao and McKinley 1994; Valiela et al. 1997; Neori et al. 2004). Previous studies have evaluated a range of macroalgal species and cultivation systems, including temperate and tropical macroalgae, land-based systems, integrated multi-trophic aquaculture (IMTA) systems, and nearshore marine installations. Many of these studies strategically cultivated a desirable alga to remove DIN from the surrounding water (Goldman et al. 1974; Ryther et al. 1975; Neori et al. 1996, 2004; Chopin et al. 1999, 2001, 2012; Buschmann et al. 2001; Troell et al. 2003; Abreu et al. 2011; Sanderson et al. 2012; Wang et al. 2012, 2014; Broch et al. 2013; Handä et al. 2013; Kim et al. 2014, 2015; Yarish et al. 2017; Fossberg et al. 2018). To determine the origin of the removed DIN, the N isotope ratio ($\delta^{15}\text{N}$) in the algal tissue can be compared to the isotopically distinct $\delta^{15}\text{N}$ of nitrogen originating from oceanic, atmospheric, treated wastewater, or fertilizer (Heaton 1986; Owens 1987; Peterson and Fry 1987).

Seaweed aquaculture and harvesting activities are expanding in the WGoM (Grebe et al. 2019; Maine Department of Marine Resources [MEDMR] 2019), which raises the question: can this growing industry potentially contribute to the maintenance or enhancement of the WGoM’s assimilative capacity for nutrients? Aquaculture leaseholders in Maine reported harvesting approximately 127 t wet weight (WW) of cultivated macroalgae in 2019 (MEDR 2019), the majority of which was processed or sold as edible (Piconi et al. 2020). Maine’s seaweed production is projected to grow at 12–15% annually to reach a total annual yield of 1360–2720 t (WW) by 2035 and new market opportunities in livestock feed, fertilizer, pharmaceuticals, and carbon or nutrient offsets are expected (Piconi et al. 2020). Most of the current seaweed aquaculture expansion is focused on kelp (order Laminariales). The most commonly grown species in Maine are the following: Saccharina latissima (Linnaeus) C.E. Lane, C. Mayes, Druel & G.W. Saunders (sugar kelp), Saccharina angustissima (Collins) Augyte, Yarish, & Neefus (skinny kelp), and Alaria esculenta (Linnaeus) Greville (winged kelp or horsetail kelp) (Grebe et al. 2019; Bricknell et al. 2020). Of the three, S. latissima is also the most frequently grown species in the USA (Kim et al. 2015; Yarish et al. 2017). In this study, we focus only on the bioextraction potential of S. latissima.

Previous studies have estimated N bioextraction by S. latissima grown in other regions by multiplying the percent N content in the kelp tissue by biomass harvested and extrapolating to a larger area (Neori et al. 2004; He et al. 2008; Chopin et al. 2012; Kim et al. 2015; Wu et al. 2015; Xiao et al. 2017; Yarish et al. 2017). Findings from these studies suggest that S. latissima aquaculture can be a useful nutrient extraction strategy in specific regions or seasons, but there is a need for more long-term estimates from a wide range of locations. Along the Eastern USA coastline, the need for improved understanding of the temporal and spatial variability of N dynamics and the related bioextraction efficiencies of specific macroalgal species is especially strong (Kim et al. 2007, 2015; Liebman et al. 2012). Kim et al. (2015) and Yarish et al. (2017) provided N bioextraction estimates by S. latissima grown in New York, Connecticut, and southern Massachusetts, but the temperature gradient along the Eastern USA coastline is one of the steepest in the world, and temperature has a strong influence on S. latissima growth (Fortes and Lüning 1980; Bolton and Lüning 1982). A better understanding of the expected macroalgal N bioremediation ranges is essential from a management perspective because bioextraction can be expensive (Neori et al. 2004). Some commonly used nutrient management practices are not adequately assessed and later found to be moderately ineffective (Boesch et al. 2001). Overestimating efficiencies of management measures is costly from a financial perspective, but it also damages social capital that had to be built by resource managers prior to initiating the treatment strategy. Thus, identifying local-regional patterns or commonalities across local studies can help build a better understanding of the range of results expected from bioextraction efforts.

In this study, we aimed to expand on previous work evaluating bioextraction by macroalgae along the Eastern USA coast to gain a more comprehensive understanding of the N extraction potential in the region. First, we estimated the N extraction of kelp harvested from the WGoM in late spring and throughout the growing season to determine the effect of harvest timing, biomass, and percent tissue N content on the total N removed.
from the surrounding water. Then, we sought to characterize the source of DIN taken up by the kelp by measuring the $\delta^{15}N$ in the collected tissue. Lastly, we provided regional context for the potential N removed through harvesting cultivated *S. latissima* from the WGoM by estimating the amount of harvested kelp needed to extract N equivalent to the N loading from atmospheric deposition, wastewater treatment facilities (WWTFs), and stormwater runoff (Garret et al. 1978; Townsend 1991; Pettigrew et al. 1998, 2005; Castro et al. 2003; Trowbridge et al. 2014).

Cultivation and sampling occurred at four sites in the Western Gulf of Maine. Two of the sites were in Casco Bay, Maine. We refer to these sites as Brothers (Bros.) Island and Cow Ledge because they were near these geographical features. The Brothers Island and Cow Ledge sites were < 3 km apart and the longlines were oriented in a similar cardinal direction (north-south) which was parallel to the prevailing current. The other two cultivation and sampling sites, Ram Island and Wood Island, were in Saco Bay, Maine (Fig. 1). The Ram Island and Wood Island sites were < 4 km apart and

### Materials and methods

#### Study site descriptions

The Gulf of Maine (GoM) is a temperate, biologically productive, waterbody extending from Nova Scotia, Canada, to Cape Cod, Massachusetts, USA (Fig. 1). Offshore, much of the GoM’s productivity is from the upwelling of nutrient-rich water from deep on the continental slope (Townsend 1998; Bricknell et al. 2020). In the coastal zone, nutrient delivery and cycling are influenced by vertical mixing by tides, wind-driven transport, small- and large-scale buoyancy forcing, large freshwater sources, atmospheric deposition, wastewater treatment facilities (WWTFs), and stormwater runoff (Garret et al. 1978; Townsend 1991; Pettigrew et al. 1998, 2005; Castro et al. 2003; Trowbridge et al. 2014).

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the longlines were oriented in a similar cardinal direction (east-west) and parallel to the prevailing current.

Casco Bay has a relatively complex, indented shoreline, whereas Saco Bay is a relatively uniform, crescent bay (Tanner et al. 2006). Previous studies have concluded that land-based N sources dominate nearshore N concentrations in Casco Bay (Castro et al. 2003; Gray 2019). Less information is available for Saco Bay, but it is presumably also heavily impacted by land-based N. Both bays receive substantial freshwater and nutrient contributions from rivers draining upland watersheds (Wade et al. 2008; Tilburg et al. 2011, 2015; Gray 2019), WWTFs employing secondary treatment, and combined sewer overflows (Maine Department of Environmental Protection [MEDEP] 2019). Combined sewer overflows (CSOs) contribute land-based nutrients to the bays after heavy rainstorms when stormwater runoff is channeled into the combined sewer collection system at a volume that exceeds the capacity of the treatment facility (MEDEP 2019). In 2019, these CSOs collectively discharged 768,000 m$^3$ of untreated stormwater runoff and wastewater into Casco Bay and 273 m$^3$ into Saco Bay (Riley 2020).

Field measurements and laboratory procedures

Kelp cultivation and collection

Saccharina latissima sporelings were produced using the methodology described in Redmond et al. (2014). Briefly, we collected wild S. latissima reproductive tissue from nearby bays and stressed it in the laboratory to release spores. Thin line was inoculated in water containing the released spores (6000–8000 spores mL$^{-1}$) over night and then transferred to aquaria. The sporelings grew in light and temperature-controlled aquaria for approximately 6–8 weeks. Outplanting occurred between October and December each year. Kelp installations at each site consisted of 1 (Wood Island and Ram Island), 2 (Brothers Island and Cow Ledge in 2018), and 5 (Cow Ledge in 2019) longlines suspended 2 m below the water’s surface. Each longline was 60–120 m long, and the spacing between each line was ≥ 6 m. Each site was less than 1 km from shore. Water depths on site were 7–17 m mean lower low water (MLLW).

Kelp cultivation occurred during four growing seasons: October to June 2016–2019. Sample collection typically began in January or February when the individual sporophytes were 30–50 cm long and 7–8 g (WW). The sporophytes were too small to obtain density estimates at that time. However, in mid-March, mean sporophyte density was typically 200–500 sporophytes m$^{-1}$. At maturity in late May, mean sporophyte density was approximately 200 sporophytes m$^{-1}$.

During sampling events, we maintained the sample integrity by removing the entire organism (holdfast, stipe, and blade) using nitrile gloves. Access to the sampling sites was weather dependent and thus, sampling frequency varied throughout the season and from year to year. During the most rigorous sampling season (2019), we completed approximately 10 sampling events at each site: roughly once per month, December through February, and 2–4 times per month from March to June. The timing of sampling was also variable across tides and time of day. At Cow Ledge in 2019, where there were 5 longlines, we collected kelp from the outermost line. All collected kelp was stored in plastic bags, transported in a covered cooler, and refrigerated at 8 °C until further processing. Transportation between the field and the laboratory was 1–2 h.

Biomass analysis

We removed and weighed all sporophytes from three, 10-cm sections of the longline to generate a mean biomass estimate for each sampling date. The location of the sections along the longline was haphazardly determined. (During a few sampling events and seasons, only one biomass measurement was possible. We do not report standard deviations for these cases). Then, we multiplied the mean biomass (WW) per 10 cm by 10 to obtain an estimate of kelp biomass (WW) per longline meter. We also established a wet to dry ratio for the samples by weighing the collected kelp upon removal from the plastic bag in the laboratory and again immediately after it had been lyophilized. The difference between the two weights was attributed to water loss and used to establish a wet to dry ratio.

Elemental and stable isotope analysis

On each sampling event, we haphazardly collected five individual sporophytes for elemental and stable isotope analysis. Within 12 h of collection, we excised a 4 cm$^2$ cutout from the basal tissue near the meristem, where metabolic activity is concentrated (Nielsen et al. 2014; Boderskov et al. 2016). The tissue was rinsed with deionized water and lightly rubbed between gloved hands for 30 s. No epiphytic algae were visibly present on the sporophytes. A small percentage (<5%) of the sporophytes had snails (Lacuna vincta) or egg rings attached to them, which we manually removed. The tissue samples were stored in a −40 °C freezer. The frozen tissue was lyophilized at −50 °C using a Labconco FreeZone Legacy 2.5 Liter Benchtop Freeze Dryer. After 24 h of drying, the lyophilized samples were homogenized into a fine powder using a mortar and pestle. The powder (2.5–5 mg) was encapsulated in tin capsules and shipped to the University of California Davis Stable Isotope Facility [UC Davis SIF] (https://stableisotopfacility.ucdavis.edu). The SIF analyzed each sample for total N, total C, $^{15}$N, and $^{14}$C using a PDZ Europa ANCA-GSL elemental analyzer interfaced with a PDZ Europa 20-20 isotope ratio mass spectrometer. The
Wave energy has an analytical precision of 0.3‰ for 15N, and the instruments were calibrated before analysis with certified standards (UC Davis SIF 2017). Overall, 364 samples were analyzed for elemental content and stable isotope ratios. We calculated elemental ratios for samples using the measurements obtained from the UC Davis SIF. We obtained percent tissue N or C content by dividing the total weight of N or C measured in each sample by the encapsulated dry sample weight. Then, we calculated the C:N ratio (M:M) for each sample from these percentages.

A common approach for estimating N removed from the marine ecosystem is to multiply the percent tissue N content in the kelp at harvest by the biomass harvested (Neori et al. 2004; He et al. 2008; Kim et al. 2013, 2014, 2015). This methodology stems from the understanding that some N is immediately used to fuel macroalgal growth, and the surplus is stored as pigments, amino acids, and proteins (Martínez et al. 2012). Therefore, we also opted for this approach and estimated total N removed by S. latissima at the time of harvest using the mean percent tissue N calculated for the dry weight (DW) of the kelp during each sampling event (Eq. 1):

\[ \text{N removed} = \frac{gN}{gDW} \times \frac{gDW}{gWW} \times \frac{gWW}{m} \]  

(1)

We adjusted the percent tissue N content by the WW:DW ratio and then multiplied by the estimated mean kelp biomass (WW g m⁻¹) for that site on the same sampling date. No assumptions regarding forms of N were included in these calculations.

The UC Davis SIF calculated the stable isotope ratios for each sample by comparing the difference in the 15N measured in the sample against the 15N in at least four different laboratory reference materials (Eq. 2) (Peterson and Fry 1987):

\[ \delta^{15}N = \left( \frac{R_{\text{sample}}}{R_{\text{reference}}} - 1 \right) \times 1000(\%e) \]  

(2)

where R is 15N/14N. The δ15N of primary producers reflects the inorganic N sources used, plus a variable amount of fractionation (differential use of 15N vs. 14N during N uptake) (Fogel and Cifuentes 1993; Fry 2006). Thus, we compared the calculated ratios to known δ15N ranges for N from specific sources. The δ15N ranges most attributed to each N source are the following: –2 to 0‰ for atmospheric N, –3 to 3‰ for N from commercial fertilizers, 4 to 8‰ marine N from natural sources, and 10‰ for N discharged from wastewater treatment processes (Heaton 1986; Macko and Ostrom 1994; McClelland et al. 1997; McClelland and Valiela 1998; Costanzo et al. 2001; Gartner et al. 2002; Cole et al. 2004; Kendall et al. 2007).

Environmental measurements

Water temperature was continuously measured using Hobo Pendant Temperature/Light 8K Data Loggers (Part #: UA-002-08). If a temperature logger was lost or compromised, we used temperature readings from the nearby University of Maine Land Ocean Biogeochemical Observatory (2016 only) and a buoy maintained by the U.S. National Oceanic and Atmospheric Administration (NOAA) (Station CASM 1). A Tilt Current Meter (TCM-1; Lowell Instruments LLC) hanging inverted from the middle spacer-buoy on each longline continuously measured current velocity and direction. Photosynthetically active radiation (PAR) was measured at 2 m underwater using an LI-193 Spherical Quantum Sensor (LI-COR) during each sampling event May 2018–June 2019. Before these dates, we estimated photosynthetically active radiation (PAR) for each bay by transforming daily Global Horizontal Irradiance obtained from the National Solar Radiation Database (https://maps.nrel.gov/nsrdb-viewer) Physical Solar Model V.3 for the Portland International Jetport station (location ID 1364086). We considered all rainfall up to 60 h before sampling as potential runoff affecting the collected kelp but excluded snowfall. Rainfall data for Casco Bay sites came from the Portland International Airport weather station maintained by NOAA. The University of New England Marine Science Center Weather Station in Biddeford, ME, provided rainfall data for the Saco Bay sites.

We collected triplicate water samples from 2 m underwater during the 2018 and 2019 growing seasons using a horizontal Niskin bottle. All water samples were stored in sealed Whirl-pak bags, transported in a covered cooler to the laboratory, refrigerated at 8 °C, and processed within 4 h of collection. Each water sample was analyzed to estimate salinity, pH, and NO3− concentration at the time of sampling. We measured salinity using a Cole-Parmer RSA-BR90A Refractometer (0–90%) and a HACH benchtop meter (model #: PW172KB0703F01) calibrated to certified standards to measure pH. We determined the concentrations of NO3− in each sample spectrophotometrically using HACH Nitrate TTNplus Low Range Vial Tests and a HACH DR3900 Laboratory VIS Spectrophotometer calibrated before analysis with certified standards. We chose to enumerate NO3− because it is a common form of problematic reactive N in waterways impacted by anthropogenic activities (Galloway et al. 2004) and more easily measured through grab sampling than nitrite or ammonium.

Statistical analysis

We examined all data for assumptions related to normality and homogeneity of variance. We identified and removed outliers using quantile ranges, robust fit, a k-nearest neighbor analysis. Then, we examined data from each study area separately (i.e., individual sites) and collectively (all sites). We used multivariate analyses of variance (MANOVA) to compare the effect of 13 environmental factors on percent tissue N, percent tissue C,
and $\delta^{13}$C, $\delta^{15}$N measured in the kelp tissue. The 13 environmental factors were as follows: site, bay, temperature, current, light, pH, salinity, ambient NO$_3^-$N at the surface, ambient NO$_3^-$N at 2 m deep, total rainfall received 60 h before sampling, grow-out week, distance from shore, and distance from nearest WWTF. When significant effects ($p \leq 0.05$) were detected, each dependent variable was analyzed separately using one-way analyses of variance (ANOVA). We performed post hoc comparisons using Tukey’s honest significant difference HSD tests and measured Pearson $R$ correlations between $\delta^{15}$N, percent tissue N, and the environmental factors. We used JMP Pro 14 for all statistical analyses.

**Results**

**Environmental conditions**

Monthly mean PAR, pH, salinity, NO$_3^-$N, and water temperatures measured during the study varied by bay (Fig. 2). The ambient water conditions at the Saco Bay sites were generally colder, higher in nutrients, and lower in pH and salinity than the Casco Bay sites. The highest salinity ($S$) measured was in February ($S = 35$) and the lowest in early May ($S = 8$). The pH of water collected from the sites was 7.5–8.2. The highest pH values occurred in November, and the lowest values occurred in May. Mean monthly nitrate in the water samples was 0.29–11.8 $\mu$M NO$_3^-$–N. Peak nitrate values occurred in early March to mid-April and then declined mid-April through May. The water temperature at the kelp farms was 1–12 °C from November to June. We observed three distinct temperature intervals. The water temperature steadily declined from 10 °C in November to approximately 2–4 °C in mid-February. Then it oscillated between 2 and 4 °C from mid-February until mid-March. Then, water temperature began to increase before reaching 10–12 °C in late May. The current velocities measured at the Cow Ledge and Wood Island in 2018 and 2019 were 3–54 cm s$^{-1}$. Specific current velocities are not available for Ram Island or Brothers Island, but they are probably like those at Cow Ledge and Wood Island, because tidal cycles drive most of the variability in currents within the nearshore WGoM.

**Biomass and elemental analysis**

Mean kelp biomass across all sites and years was 9.84 ($\pm$ 2.53) to 14.84 kg WW m$^{-1}$ longline at the time of harvest (Table 1). Wet to dry ratios of the kelp were 7.4:1 in March to 8.7:1 in May. The highest sampling frequency, and thus insight into biomass increase, occurred in Spring 2018 and Spring 2019 at the Wood Island site (Fig. 3). In 2018, biomass measurements at this site show that peak growth occurred in late May. Interestingly, measurements from 2019 at this site show peak growth from April until mid-May, followed by a decline in biomass in late May.

Mean N content of the *S. latissima* tissue, calculated for each sampling event, was 1.04–3.82% ($\pm$ 0.22) DW throughout the growing season and generally decreased through the spring. Tissue N content at Wood Island in 2018 and 2019 illustrated the general trend: at this site, percent tissue N decreased 0.08–0.17% week$^{-1}$ from mid-April to late-May (Fig. 3). In late May, percent tissue N contents were 1.04–2.29% ($\pm$ 0.09) DW. Increasing water temperature was negatively associated with percent tissue N (MANOVA; $F_{(9,155)} = 37.49$, $p < 0.0001$). Photosynthetically active radiation (PAR) was positively associated with percent tissue N (MANOVA; $F_{(9,155)} = 4.65$, $p = 0.032$). Site also had significant effects on percent tissue N (MANOVA; $F_{(9,155)} = 4.10$, $p = 0.0078$). There were no significant direct correlations (Pearson’s $r$) between the percent tissue N and environmental data. Across sites and all years, the mean C:N ratio ($M:M$) measured in the *S. latissima* tissue was 9.4 ($\pm$ 0.7) to 23.4 ($\pm$ 10.8). The lowest C:N ratios were in March; the highest ratios were in May and June (Fig. 4).

**Stable isotopes**

The interannual mean $\delta^{15}$N measured in the kelp tissue grown at the two Casco Bay sites was significantly different.
Table 1  Mean *Saccharina latissima* biomass (kg m\(^{-1}\)) \% N content (DW), and tissue C\(\text{N}(M,M)^{15}\text{N}\) measured at harvest from Spring 2012–2019. These observations were used to estimate the potential biomass produced from 1 ha of farming activity at two densities (6 m and 1.5-m spacing between longlines) and the potential N (kg ha\(^{-1}\)) removed when harvesting this biomass. Values are rounded to the nearest tenth. Standard deviations are reported in parentheses when possible, but replicates were not collected for all years.

| Year | Biomass (kg m\(^{-1}\)) | C:N (%) | \% N (DW) | Total N (kg) removed ha\(^{-1}\) |
|------|-------------------------|---------|-----------|-------------------------------|
|      | 6-m spacing             | 1.5-m spacing |
|      | Bros. Island | Cow Ledge | Ram Island | Wood Island | Bros. Island | Cow Ledge | Ram Island | Wood Island |
| 2016 | 1.30                   |           |           | 17.0 (±1.5) |
| 2017 | 1.25 (±0.1)            | 1.1 (±0.1) | 1.2 (±0.1) | 14.8 (±0.4) |
| 2018 | 1.48 (±0.4)            | 2.0 (±0.3) | 1.8 (±0.2) | 2.2 (±0.7)  |
| 2019 | 1.34 (±0.4)            | 0.6 (±0.1) | 0.1 (±0.1) | 0.5 (±0.2)  |

Discussion

**Biomass and bioextraction estimates**

*Saccharina latissima* grew well in both Casco and Saco Bay. Biomass per longline meter at harvest (10–15 kg m\(^{-1}\)) (WW) was on the higher end of the ranges previously reported in the literature (Table 3). The mean wet weight to dry weight ratio (WW:DW) ratio of the sporophytes at harvest was also slightly higher than the 7:1 reported by Sanderson et al. (2012) for cultivated *S. latissima* from an IMTA system in Scotland. In our data, the highest and lowest biomass measurements have almost 1% difference in the tissue N content (Cow 2018 vs. Wood 2019). The observed 1–4% N content is comparable to the range reported by other studies where *S. latissima* was grown in water with high DIN from anthropogenic or fish waste (Handå et al. 2013; Kim et al. 2015; Marinho et al. 2015; Yarish et al. 2017), but the higher 4% N tissue content also exceeded the upper value reported by several studies where *S. latissima* was grown in IMTA or relatively unimpacted water (Sanderson et al. 2012; Bruhn et al. 2016; Freitas et al. 2016; Fossberg et al. 2018). Maximum potential N removal did not coincide with peak percent tissue N and percent tissue N observed in the kelp at the time of May harvest was lower than previously observed for *S. latissima* in the Northwest Atlantic (Kim et al. 2015; Yarish et al. 2017). Almost all percent N and C:N ratios measured in the kelp tissue at harvest indicate that nitrogen was limiting. Like Kim et al. (2015) and Yarish et al. (2017), we observed a high degree of temporal and spatial variability of tissue N content in *S. latissima*.

As previously demonstrated, our results can be extrapolated to generate rough, hectare-scale estimates of potential...
bioextraction by kelp harvesting in the region. With a moderate, 6 m of spacing between longlines, a total of 1767 m of longline fit in one hectare of ocean surface. Multiplying this by the calculated kg kelp m$^{-1}$ would result in a kelp harvest of 17.3 ($\pm$ 4.4)–26.1 t WW ha$^{-1}$. The intensive cultivation scenario, with 1.5-m spacing between longlines, has 6768 longline meters per hectare, which would produce 70.3–100.1 t WW ha$^{-1}$. Converting by the WW:DW ratio (8.7:1), and then multiplying by the mean % tissue N measured in kelp from each site at harvest, results in an estimated 19.2 ($\pm$ 4.8)–46.0 ($\pm$ 2.0) kg N ha$^{-1}$ that could be removed by harvesting a hectare of $S$. latissima with 6-m spacing between longlines (Table 1).

Previously published estimates of N loading to Casco Bay (i.e., atmospheric N deposition, N loading from upland activities in the watersheds, and effluents from large WWTFs) help to put the potential bioextraction from kelp aquaculture in context (Table 2). We can calculate the approximate area of $S$. latissima harvest needed to remove a quantity of N equivalent to the N that is delivered to Casco Bay from these sources. In all examples considered, the quantity of N removed from Casco Bay by harvesting one hectare of $S$. latissima would be greater than the amount of N contributed to the Bay from one hectare of any loading sources. For example, even with 6-m spacing between longlines, the N
extration by 1 ha of *S. latissima* harvest is equivalent to the annual atmospheric deposition of N across 2.7 (± 0.7)–10.7 (±0.5) ha of Casco Bay or 5.1 (± 1.3)–12.1 (± 0.5) ha of activities in a nearby urban subwatershed. Insufficient data on N inputs prevents a direct comparison for Saco Bay, but we expect that the pattern would be similar.

These estimates of total N removed per hectare of kelp harvested from the WGoM (19.2–46 kg ha$^{-1}$ with 6-m longline spacing) are higher than many of the ranges reported by other studies evaluating *S. latissima* for bioextraction at nearshore and IMTA sites (Table 3). Of particular interest, again, is the comparison between this study and those in closest proximity. In the Long Island Sound, CT and the Bronx River Estuary, NY, Kim et al. (2014, 2015) and Yarish et al. (2017) calculated 10–35 kg N ha$^{-1}$ removed with 6-m longline spacing and 29–139 kg N ha$^{-1}$ with 1.5-m longline spacing (Kim et al. 2014, 2015; Yarish et al. 2017). Additionally, Augyte et al. (2017) estimated 88.7 kg ha$^{-1}$ N removal by closely related species, *Saccharina angustissima* (formerly *Saccharina latissima* forma *angustissima*), cultivated near Bristol, ME, and Sorrento, ME, using a 2.5-m spacing between longlines. We recalculated this to be 124 kg ha$^{-1}$ N removal by *S. angustissima* with 1.5-m spacing between longlines and note that this estimate lies in the middle of the range reported by this study for *S. latissima* grown with the same longline spacing in Saco and Casco Bay.

This emphasizes the importance of considering cultivation density and harvest timing when evaluating bioextraction applications. Unsurprisingly, increasing the density of longlines on a hectare of ocean surface produced a much higher estimate of N extraction per hectare. However, we must consider these estimates with caution. The risk of overestimating bioextraction increases when extrapolating from dispersed longlines to higher densities because intensive cultivation reduces the water flow delivering nutrients, and thus the tissue N content, but values from low-density field studies do not reflect this (Kerrison et al. 2015; Marinho et al. 2019). The range of δ$^{15}$N reported by this study for *S. latissima* grown with the same longline spacing in Saco and Casco Bay.

These δ$^{15}$N values spanned from Casco (gray fill) and Saco Bay (white fill) from 2016 to 2019. The range of δ$^{15}$N commonly associated with nitrogen (N) from treated wastewater is δ$^{15}$N = > 10‰, oceanic N δ$^{15}$N = 4–8‰, and fertilizers δ$^{15}$N = −3–3‰. Unshaded ranges represent overlap between N sources. Ranges for δ$^{15}$N from different sources were obtained from: Heaton 1986; Macko and Ostrom 1994; McClelland et al. 1997; McClelland and Valiela 1998; Costaño et al. 2001; Gartner et al. 2002; Cole et al. 2004; Kendall et al. 2007.

Fig. 6 Monthly nitrogen isotope ratios (%$\delta$) measured in *Saccharina latissima* from Casco (gray fill) and Saco Bay (white fill) from 2016 to 2019. The range of δ$^{15}$N commonly associated with nitrogen (N) from treated wastewater is (δ$^{15}$N = > 10‰), oceanic N (δ$^{15}$N = 4–8‰), and fertilizers (δ$^{15}$N = −3–3‰). Unshaded ranges represent overlap between N sources. Ranges for δ$^{15}$N from different sources were obtained from: Heaton 1986; Macko and Ostrom 1994; McClelland et al. 1997; McClelland and Valiela 1998; Costaño et al. 2001; Gartner et al. 2002; Cole et al. 2004; Kendall et al. 2007.

### Table 2 Amount of estimated atmospheric, riverine, and treated wastewater N loading into Casco Bay potentially offset by the harvest of 1 ha of *Saccharina latissima*

| Source of N | Annual N (kg ha$^{-1}$) | 6-m spacing | 1.5-m spacing |
|-------------|--------------------------|-------------|---------------|
| Atmospheric deposition (dry + wet) | 4.3$^a$ | 4.5 (± 1.1)–10.7 (± 0.5) | 17.1 (± 4.3)–40.9 (± 1.8) |
| Low estimate | | | |
| Atmospheric deposition (dry + wet) | 7.2$^a$ | 2.7 (± 0.7)–6.4 (± 0.3) | 10.2 (± 2.5)–24.4 (± 1.1) |
| High estimate | | | |
| Presumpscot river watershed (forested) | 1.5$^b$ | 12.8 (± 3.2)–30.7 (± 1.3) | 49.0 (± 12.3)–117.3 (± 5.1) |
| Royal river watershed (forested) | 5.3$^b$ | 3.6 (± 0.9)–8.7 (± 0.4) | 13.8 (± 3.5)–33.1 (± 1.5) |
| Capisic brook watershed (urban) | 3.8$^b$ | 5.1 (± 1.3)–12.1 (± 0.5) | 19.4 (± 4.9)–46.4 (± 2.0) |
| Effluent from large WWTFs | 3.5$^b$ | 5.5 (± 1.4)–13.2 (± 0.6) | 21.1 (± 0.9)–8.7 (± 0.4) |

$^a$ Sonoma Technology Inc. (2003) estimated that atmospheric N deposition (wet + dry) to Casco Bay is 4.3–7.22 kg ha$^{-1}$ year$^{-1}$ inorganic N

$^b$ Recent work by Gray (2019) suggested that the large and predominantly forested Presumpscot and Royal River watersheds respectively export 1.5 kg to 3.79 kg N ha$^{-1}$ year$^{-1}$ into Casco Bay. She estimated nitrogen loading from the smaller, but urbanized, Capisic Brook watershed to be 5.31 kg N ha$^{-1}$ year$^{-1}$ (Gray 2019)

$^c$ Annually, the six largest WWTFs near Casco Bay discharge an estimated 914 mt of N into the bay which is approximately 3.5 kg N ha$^{-1}$ year$^{-1}$ across the area of Casco Bay. Source: MEDEP [Maine Department of Environmental Protection] (2008) Development of nutrient criteria for Maine’s coastal waters. https://www.maine.gov/dep/water/nutrient-criteria/nutrient_criteria_report_2008.pdf. Accessed 2 Feb 2020
Galicia and Spain

Long Island Sound, USA

Badcall and Calbha, Scotland

Horsen’s Fjord, Denmark

Galicia and Canabria, Spain

Tristeinrasa, Norway

Sogn and Fjordane, Norway

Galicia, Spain

New Brunswick, Canada

Bocabec Bay, Canada

| Location | Cond. notes | Temp (°C) | Salinity | Current (cm s⁻¹) | Harvest time | Biomass (kg WW m⁻²) | % tissue N (DW) | Longline spacing (m) | Total biomass (WW t ha⁻¹) | Total N removed (kg ha⁻¹) | Reference |
|----------|-------------|-----------|----------|------------------|--------------|---------------------|----------------|----------------------|-----------------------------|--------------------------|-----------|
| Gulf of Maine, USA | – | 2–12 | 8–35 | 1–3 | May | 10–15 | 1–4 | 6 | 16–24 | 19–46 | Current study |
| Long Island Sound, USA | – | 0–17 | 21–33 | – | May/June | 1–19 | 1–4 | 6 | 9–10 | 10–35 | Kim et al. (2015); Yarish et al. (2017) |
| Badcall and Calbha, Scotland | IMTA | 7–15 | – | – | June | _* | 1–3 | _* | _* | _* | Sanderson et al. (2012) |
| Horsen’s Fjord, Denmark | – | 0–19 | 12–27 | 0–34 | May | 0–1 | 1–3 | 8–10 | 2–7 | 3–26 | Marinho et al. (2015); Bruhn et al. (2016) |
| Galicia and Canabria, Spain | – | 13–17 | – | 12–92 | May | 4–16 | 1–4 | 8–10 | 7 | 31 | Marinho et al. (2015) |
| Tristeinrasa, Norway | IMTA | 4–14 | 27–34 | 0–20 | June | _* | 2–5 | _* | _* | _* | Handå et al. (2013); Wang et al. (2014) |
| Sogn and Fjordane, Norway | – | 4–15 | – | – | – | 1–2 | – | – | – | Fossberg et al. (2018) |
| Galicia, Spain | IMTA | 11–16 | – | – | April | _* | 2–3 | _* | _* | _* | Freitas et al. (2016) |
| New Brunswick, Canada | – | – | – | – | 8–16 | – | – | – | – | Druehl et al. (1988); Chopin et al. (2004) |
| Bocabec Bay, Canada | IMTA | – | – | – | – | 8–21 | – | – | – | – | Chopin et al. (2004) |

*Information provided was for a cultivation array that does not allow for cross-comparisons to horizontal longlines.

Additionally, higher density cultivation could exceed the environmental or social carrying capacity for kelp aquaculture in the region, which is why we have both evaluated a range of longline densities and underlined the need for integrated management of N pollution. The timing of kelp farm deployment and harvesting also influences bioextraction services of kelp grown in the WGoM. For example, from February through early May 2019, even as percent tissue N decreased throughout the spring, the biomass increased, and thus, so did the potential N removed through the harvest of all cultivated kelp. However, biomass did not increase in the same way during the last couple weeks of May 2019 due to reduced growth rates and sloughing, possibly associated with ambient water temperatures exceeding 10 °C. Therefore, to maximize the N extracted in 2019, the sugar kelp should have been harvested in early May rather than late May. The most dramatic example is from the Wood Island site in 2019, where harvesting one month earlier would have doubled N removal (27.4 kg ha⁻¹ vs. 51 kg ha⁻¹). However, these gains also appear to vary by site. At the other sites that same spring, harvesting 3–4 weeks earlier would have resulted in 3–22 kg ha⁻¹ more N removed. Additionally, in 2018, the highest estimates of N removal were obtained in late May, possibly because the ambient water temperature did not reach 10 °C until that time. This highlights an opportunity for active monitoring of the ambient DIN and dissolved inorganic carbon at kelp aquaculture installations and N and C content in the kelp tissue. Using real-time estimates of N removal and ambient environmental conditions to schedule harvesting could maximize bioextraction effects.
Even with optimizations to harvest timing and density of longline arrays, kelp bioextraction must be part of a comprehensive N management strategy. Human activity has added reactive N to the landscape and changed nearshore habitats in ways that enhance N delivery to coastal ecosystems (Cleveland et al. 1999; Galloway et al. 2004). Comparing the maximum N potentially removed by harvesting a hectare of S. latissima to sources of nitrogen loading in Casco Bay reinforces the magnitude of anthropogenic disturbance in the N cycle. Encouragingly, the hectare-level comparisons generated for Casco Bay suggest that kelp bioextraction may be an efficient in-water tool to intercept nonpoint source pollution like atmospheric N deposition which, again, can be 30–40% of total N load to Casco Bay (Castro et al. 2003; Sonoma Technology Inc. 2003). However, the application of N bioextraction technologies must only be an additional measure for mitigating anthropogenic impacts on the environment. It should not be an alternative to improved management of point source and nonpoint source N by reducing combustion of fossil fuels, decreasing the application of N-based fertilizes, and tertiary treatment of wastewater. Using kelp aquaculture to remediate any substantial quantities of N will require a considerable shift in social acceptance of marine development and would have to be carefully evaluated against other commercial and ecological needs for this bay.

**Environmental conditions**

Careful consideration of environmental variables’ potential effect is important when anticipating how potential yields and nutrient concentrations reported by this study might vary. Many of the measured environmental conditions exhibited patterns like those reported by Kim et al. (2014, 2015) and Yarish et al. (2017); however, ambient salinity at our sites exhibited more dramatic swings than those observed in Long Island Sound. Mean ambient salinity measured at each sampling event declined from 30 to 23 in Saco Bay and 32 to 29 in Casco Bay in March and April, and salinity dropped as low as 16–17 at Cow Ledge and Wood Island in mid-April. This decline in salinity is earlier and steeper than the lowest salinities of 22–26 that Kim et al. observed at their sites in May. This discrepancy is notable regarding the timing of stress on the kelp crop. Saccharina latissima is semi-euryhaline; it can withstand 23–35‰ with no reduction in growth (Druehl 1967; Bartsch et al. 2008), but stress responses often develop at salinities below this range. A sharp decline in growth occurred in S. latissima in salinities consistently below 16 (Bartsch et al. 2008; Nielsen et al. 2014), and Gordillo et al. (2002) found that the closely related, Laminaria digitata, exhibited reduced nitrate uptake rates in low salinity conditions. Thus, the spring flush timing leading to freshening events in the WGoM could have affected the growth and tissue composition of the sampled kelp and may ultimately impact the potential N removed by kelp in this region.

The potential impact of combined stressors should also be considered when interpreting results from this study. No statistically significant relationships were observed between percent tissue N, δ15N, and the measured environmental conditions. One explanation for this may be that an alga’s tolerance range for one environmental factor may be influenced by other environmental factors (Hurd et al. 2014). For example, when Mortensen (2017) grew S. latissima and L. digitata in water enriched with nitrate and phosphate, the algae survived almost 2 weeks in brackish water (salinity = 18). In our study, one or several of the environmental conditions measured were less than optimal for S. latissima growth at some point during the growing seasons. For instance, temperatures at the sampling sites did not reach the 5–15 °C optimal growth range for S. latissima (Fortes and Lüning 1980; Bolton and Lüning 1982; Kim et al. 2015; Yarish et al. 2017) until mid-March. Photosynthetically active radiation measured during some sampling events was lower than the light-saturating level of 150–215 μmol photons m−2 s−1 reported for adult S. latissima sporophytes (Lüning 1979; Bartsch et al. 2008). Similarly, the range of current speeds (3–54 cm s−1) during periods of 2018 and 2019 seasons is broader than the optimal 10–25 cm s−1 flow rate for S. latissima (Kerrison et al. 2015). Lastly, initial sporophyte density (200–500 m−1) may have resulted in clumping and shading preventing adequate light and nutrients from reaching all sporophytes. The statistically significant effect of site on both the δ15N and percent tissue N observed in our results may be the result a combined stressor effect involving any of these ambient conditions and perhaps even other stressors that were not detected. Or inversely, the absence of a clear relationship between percent tissue N, δ15N, and may be because the algae were able to tolerate passing colder temperatures, low light, lower or higher current, or higher cultivation densities because the other environmental conditions were more than adequate.

**Sources of N-stable isotopes**

Mean δ15N measured in the kelp tissue did not show a clear indication that kelp grown and collected from Saco and Casco Bay took up N from anthropogenic sources. This finding contrasts with the general picture of coastal WGoM dynamics presented by Castro et al. (2003), Liebman et al. (2012), and Trowbridge et al. (2014). The absence of a clear N source relationship is also dissimilar to conclusions presented by Kim et al. (2015), who described clear indications that anthropogenic N sources were taken up by S. latissima in the Bronx River Estuary (~2–6‰) and Long Island Sounds (9–19‰). The interannual, site-specific means δ15N for samples from Brothers Island (7.6‰), Cow Ledge (6.6‰), Ram Island (6.3‰), and Wood Island (5.7‰) sites fell within the δ15N ranges commonly attributed to N of marine origin (4–8‰) (Fig. 5). However, the high end of the δ15N range measured
in kelp grown at Brothers Island (4.7–11.51‰) spans into the δ15N values commonly attributed to N from treated wastewater (10–12‰). In Saco Bay, the low end of the δ15N range measured in kelp grown at Wood Island reached into the δ15N values commonly attributed to N from fertilizers (−3–3‰). Looking at δ15N by bay (Fig. 6), the range of tissue δ15N in Saco Bay *S. latissima* reached its lowest values, indicative of N originating from fertilizer, in February. Also, during February, some measurements of tissue δ15N in Casco Bay *S. latissima* had values indicative of N originating from treated wastewater (> 10‰) but the sample mean was much lower (5.78‰). The monthly mean tissue δ15N in Casco Bay *S. latissima* continued to rise through May. Nutrient bioavailability, *S. latissima* ecophysiology, or unmeasured environmental changes may have influenced these results obtained in the present study.

Nutrient bioavailability at the study sites, affected by flushing rates and uptake by wild species, may have also limited exposure of the sampled kelp to anthropogenic N. Slow N supply rates and low amounts of N substrate are key considerations for N isotope distributions in primary producers because they limit reactions important for growth (Peterson and Fry 1987). In N-limited systems, macroalgae do little fractionation of their source material during N uptake (Peterson and Fry 1987; Savage and Elmgren 2004; Thomber et al. 2008); all available N will be consumed regardless of isotope content so long as redox conditions remain relatively stable. Given the stable redox conditions in this well-mixed, highly oxygenated environment, we assume the observed δ15N values in the tissue were representative of the N source (Wada and Hattori 1978; Mariotti et al. 1982; Pennock et al. 1996). However, fractionation by some macroalgae has occurred in water with high DIN concentrations, which resulted in tissue-δ15N values lower than that of the δ15N measured in the source N (Wada and Hattori 1978; Mariotti et al. 1982; Peterson and Fry 1987; Pennock et al. 1996; Wang et al. 2014). Examining the mean percent tissue N and the tissue C:N ratio in the *S. latissima* each month and at harvest indicates that there were periods during many of the growing seasons when the kelp was N limited. In *S. latissima*, > 3% DW tissue N content suggests N sufficiency, 1.9% is the minimum required for maximal growth, and < 1.3% DW tissue N indicates N limitation (Chapman et al. 1978; Wheeler and Weidner 1983; Kim et al. 2015). Therefore, δ15N values measured during or after a period of N limitation may not be comparable to when N was replete in the kelp tissue (Aberle and Malzahn 2007). The natural assimilatory capacity and high flushing rates of the WGoM may also explain the absence of a clear anthropogenic isotopic signature in the cultivated *S. latissima* despite the known contributions of anthropogenic N. Additionally, the WGoM has large, naturally occurring, *Fucus* spp. and *Ascophyllum nodosum* beds in the intertidal and subtidal zones. These wild algae may have also intercepted some anthropogenic N before it reached the study sites.

The nutrient ecophysiology of the sampled kelp (i.e., starving or N saturated) may have affected fractionation rates that are crucial assumptions for the application of stable isotope ratio assessments of primary producers. Fernandes et al. (2012) found that large N reserves in algal tissue can mask the isotopic signal of newly acquired N, and kelp cells have large vacuoles enabling N storage. When ambient N is abundant, kelp cells can store N as nitrate in cellular vacuoles and cytoplasm (Fong et al. 1994). Then, they draw on these reserves when ambient N is low (Chapman and Craigie 1977; Egan and Yarish 1990). It is plausible that this nutrient ecophysiology resulted in a muddled δ15N that is not representative of recent N use. For example, if *S. latissima* took up and stored N from the marine environment in December–February, the stored N would have a δ15N reflecting that source. When this stored N was assimilated into algal tissue later in the spring, because ambient N was insufficient for the sporophytes’ accelerated growth rates, the tissue sampled at that time would still exhibit a δ15N that was influenced by a marine N source despite the possibility that the algae could be using N from another source. Cellular N reserves in the *S. latissima* could also explain why there were no statistically significant relationships between percent tissue N, tissue δ15N, and ambient nitrate at each site.

Lastly, undetected environmental changes in the N sources or at the study sites may have affected the δ15N results. The isotopic composition of N species within aquatic environments is affected by many environmental processes including assimilation, denitrification, nitrification, and mineralization (Wada et al. 1975; Wada and Hattori 1978; McCready et al. 1983). Substantial changes in ambient environmental conditions can result in a shifted δ15N ratio for N sources, making it challenging to use stable isotope techniques to identify nutrient sources in field studies (Fry 2006; Wayland and Hobson 2001). For example, the presence, or pulses, of ammonium at the sites may help to explain why there was no correlation between ambient nitrate concentrations and δ15N. *Saccharina latissima* exhibits a preference for ammonium. Harrison et al. (1986) found that nitrate uptake in *S. latissima* was completely suppressed for 30 min following a pulse of ammonium. We assumed that any ammonium delivered to the sites would be immediately taken up, so we did not attempt to quantify ammonium in this study. However, frequent ammonium supplies or an ammonium pulse shortly prior to a sampling event may have also influenced N uptake rates or provided a contrasting δ15N signal.

Undocumented phytoplankton blooms are another example of an undetected environmental event that may be a source of variability influencing our dataset. Yarish et al. (2017) attributed low tissue N in kelp to a prolonged spring phytoplankton bloom, which may have been supported by mild winter conditions (i.e., harsh winter and spring results in more DIN available for the macroalgae). Anderson et al. (2005) also
found correlations between spring snowmelt and spring phytoplankton blooms in southern New England. Releases of N from $^{15}$N-depleted sediments would have also affected the $\delta^{15}$N measured in the kelp tissue (Altabet 2006; Bianchi 2007; Sigman et al. 2009). Without knowledge or measurement of a release, it would be hard to correct for it when interpreting the data presented here. It is also possible that the natural variation between sites, or between published $\delta^{15}$N values for N sources and those in the WGoM, is so considerable that it exceeds the capacity of stable isotope analysis to differentiate between the N sources (Ostrom et al. 1997; Fry 2006). Due to logistical constraints, characterization of the $\delta^{15}$N in NO$_3^-$ from specific N sources in Casco and Saco Bay was not possible. However, if future work can do this, it will reduce uncertainty regarding unmeasured environmental conditions and support the development of a stable isotope-specific mixing model for these locations.

Importantly, the isotope values reported in this study can help us to understand the current WGoM biogeochemistry and the existing degree of human perturbation in Casco and Saco Bay. If used in future studies, they will also help to better describe the direction and magnitude of nutrient cycling in the WGoM (Peterson and Fry 1987; Ostrom et al. 1997; Dethier et al. 2013). Establishing baseline stable isotope values for S. latissima in this region will help with the detection of potentially incipient eutrophication, which is preferable to restoration (McClelland et al. 1997). Additionally, if future studies can demonstrate a closer relationship between anthropogenic N pollution and bioextraction provided by kelp in the WGoM, it will garner stronger public support for cap and trade programs to include bioextraction as an eligible activity.

**Conclusion**

Identifying and implementing effective nutrient management technologies is critical to mitigating the impact of human activities on coastal ecosystems. This study measured biomass, $\delta^{15}$N, and tissue N content of Saccharina latissima grown from 2016 to 2019 at four sites in Casco and Saco Bay, Maine, to better understand how the N bioextraction achieved by harvesting cultivated kelp varies across space and time. Although the patterns in elemental content of the S. latissima tissue from the WGoM are like those reported from further south, total biomass at time of harvest was higher. Significant variation in biomass and tissue N content was observed between sites between the two bays, potentially due to combined environmental stressors, or the timing of seasonal temperature and salinity changes between the bays. High variation in $\delta^{15}$N also occurred between sites, and the monthly and interannual mean $\delta^{15}$N did not show explicit use of anthropogenic N sources like wastewater or fertilizer. The absence of clear source N relationships may be the result of physiological traits of S. latissima, biogeochemical characteristics of the WGoM, or unmeasured environmental changes. Our results further highlight the need for site-level pilot studies, even within the same bay, to characterize the seasonal and spatial variation of N assimilation before any kelp aquaculture is developed solely for bioextraction purposes in the WGoM. Finally, we extrapolated our results to estimate that harvesting cultivated kelp from the WGoM has the potential to extract 19.2 (+4.8)–176.0 (+7.7) kg N ha$^{-1}$ depending on the cultivation density used, which emphasizes the importance of cultivation density and harvest time on theoretical kelp aquaculture bioextraction efficiencies. We conclude that kelp farming and harvesting could be a component within a broader, integrated approach to N mitigation in the region, but a substantial increase in kelp production and social acceptance of aquaculture will be required.

**Acknowledgments** We thank the following individuals for their innovative ideas and never-ending help in the laboratory and on the water: Arenti T, Beard K, Brawley S, Costa-Pierce B, Hollandbeck M, Jagoutz T, Johndrow K, Jones E, Olson T, Pierce E, St. Gelais A, and Waters A. Much appreciation to Liberti K, and Cleaver C, Koons B, and the anonymous reviewers for their constructive comments.

**Funding** This activity was supported by National Science Foundation award #IA-1355457 to Maine EPSCoR at the University of Maine.

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