



## Spatiotemporal Patterns in the Biomass of Drift Macroalgae in the Indian River Lagoon, Florida, United States

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Hall LM, Morris LJ, Chamberlain RH, Hanisak MD, Virnstein RW, Paperno R, Riegl B, Ellis LR, Simpson A and Jacoby CA (2022) Spatiotemporal Patterns in the Biomass of Drift Macroalgae in the Indian River Lagoon, Florida, United States. Front. Mar. Sci. 9:767440. doi: 10.3389/fmars.2022.767440 Drift macroalgae plays key roles in the ecology of many coastal systems, including the Indian River Lagoon. In the lagoon, changes in the biomass of drift macroalgae may have interacted with an unprecedented bloom of phytoplankton in 2011. Patterns in the biomass of drift macroalgae were identified using new and original analyses of data from several sampling programs collected between 1997 and 2019. All available data show a relatively low biomass of drift macroalgae in 2010-2012, and surveys of fixed transects and seining as part of a fisheries independent monitoring program also recorded low biomass in 2016. Low light availability and potentially stressful temperatures appeared to be the main influences as indicated by the results of incubations in tanks to determine environmental tolerances and data on ambient conditions. Decreased biomass of drift macroalgae had implications for cycling of nutrients because carbon, nitrogen, and phosphorus not stored in the tissues of drift macroalgae became available for uptake by other primary producers, including phytoplankton. The estimated 14-18% increases in concentrations of these elements in the IRL could have promoted longer and more intense phytoplankton blooms, which would have reduced light availability and increased stress on algae and seagrasses. An improved understanding of such feedback and the ecological roles played by drift macroalgae will support more effective management of nutrient loads and the system by accounting for cycling of nutrients among primary producers.

Keywords: abundance, physiological tolerance, nutrient content, nutrient cycling, habitat suitability

## INTRODUCTION

Interactions among three key primary producers, phytoplankton, macroalgae, and rooted macrophytes, represent influential processes in many estuaries (Duarte, 1995; Kinney and Roman, 1998; Viaroli et al., 2008). In the Indian River Lagoon (IRL), drift macroalgae (DMA) can dominate the total biomass of submersed aquatic vegetation, and even during years when seagrass is abundant, DMA contribute a significant proportion of the total primary productivity (Dawes et al., 1974; Josselyn, 1977; Thompson, 1978; Virnstein and Carbonara, 1985; Jensen and Gibson, 1986).

Beyond being an important primary producer, DMA also provide habitat for small animals and a place to forage for predators (Stoner, 1980; Kulczycki et al., 1981; Virnstein and Howard, 1987; Holmquist, 1997).

As key primary producers in the IRL, DMA play an important role in cycling of carbon, nitrogen, and phosphorus. The ability of macroalgae to take up and store nutrients makes them successful when nutrients are limiting or supplied in pulses, which allows them to compete with phytoplankton for access to elements in the water column (Marshall and Orr, 1949; Hanisak, 1983). However, DMA are less robust and persistent than rooted macrophytes, so their death or lack of growth can add or leave carbon, nitrogen, and phosphorus that become available for uptake by fast-growing phytoplankton (Hanisak, 1983, 1993; Lavery and McComb, 1991; Gao et al., 2013). In fact, shifts from dominance by benthic primary producers to dominance by phytoplankton have been observed in multiple systems with negative impacts on seagrass assemblages and their associated fauna (Jensen and Gibson, 1986; Duarte, 1995; Burkholder et al., 2007; Duarte et al., 2010). Such a shift may have occurred in the IRL because an unprecedented sequence of intense and longlasting blooms of phytoplankton has afflicted the system since 2011 (Phlips et al., 2010, 2011, 2015).

Although important in the IRL and elsewhere, DMA are difficult to sample effectively and efficiently. The fact that DMA are moved by currents complicates the choice of locations to sample, and details regarding their movements in the IRL are poorly known (Kulczycki et al., 1981; Virnstein and Carbonara, 1985; Jensen and Gibson, 1986; Hanisak, 2021). Additionally, DMA occur across a wider range of depths than rooted macrophytes because they have lower requirements for light (Biber et al., 2004; Hily et al., 2004; McGlathery et al., 2007), so the need to survey more area and deeper water makes effective sampling more challenging. Drift macroalgae also are relatively ephemeral, and changes in their abundance within and between years, including 'boom and bust cycles' driven by environmental conditions make the choice of temporal intensity for sampling important (Virnstein and Carbonara, 1985; McGlathery et al., 2007).

In spite of these challenges, the objectives of this paper are to identify the presence, causes, and consequences of differences and changes in the biomass of DMA across space and through time in the IRL. Substantial and consistent patterns in the biomass of DMA are elucidated from data collected at multiple spatial and temporal scales, and those patterns are related to environmental drivers that reflect physiological tolerances identified in incubations. The implications of spatiotemporal variation in biomass of DMA are translated into effects on cycling of elements in the IRL via data on the carbon, nitrogen, and phosphorus content of several species of DMA.

## MATERIALS AND METHODS

#### **Study Area**

The IRL is a shallow, bar-built estuary along the east coast of Florida comprising three, interconnected sublagoons: Mosquito

Lagoon, Banana River Lagoon, and Indian River Lagoon (**Figure 1**). The average water depth in the sublagoons is 1.9 m, and most of the system is microtidal (semidiurnal lunar tidal amplitude range 0.2–18.1 cm), with more substantial changes in water level driven by wind and the seasonal rise and fall of the coastal ocean (Smith, 1987; Pitts, 1989; Steward et al., 2005). Significant exchange with the coastal ocean is limited to five inlets located primarily in the southern portion of the IRL. This configuration contributes to residence times of over a year in the portions of the system that are far from inlets (Steward et al., 2005).

For this paper, the IRL was subdivided into reaches (**Figure 1**). The reaches were delineated by evaluating similarities in time series of water quality parameters (Lasi et al., this volume). The first six reaches were within the St. Johns River Water Management District. The remainder, reaches 7, 8, and 9, were within the boundary of the South Florida Water Management District.

## **Biomass of Drift Macroalgae**

Data generated by visual surveys of fixed transects, seines of two sizes hauled as part of fisheries independent monitoring, and hydroacoustic surveys were analyzed independently because samples were taken at different frequencies, over different areal extents, and across depths that may or may not overlap (**Supplementary Table 1**). Given these variations, there was no attempt to intercalibrate the data, and interpretations focused on changes or differences that were consistent across methods.

A focus on consistent and substantial changes also resulted from a lack of details on the taxonomic composition of DMA. In surveys of transects, which focused on seagrass, and seining, which focused on fish and macroinvertebrates, DMA was considered bycatch; therefore, the biomasses of individual species typically were not measured. Furthermore, hydroacoustic surveys could not differentiate species. Nevertheless, widespread, consistent, and statistically significant spatiotemporal patterns provide valuable insights into the roles of DMA.

#### **Fixed Transects**

Fixed transects were surveyed at least twice a year (summer and winter) approximating times of annual maximum and minimum abundance of seagrasses (**Figure 1**; Virnstein and Morris, 1996). The location of each transect was marked with poles, and the path to be surveyed was delineated by a graduated line extending perpendicularly from the shoreline out to the deep end of the seagrass canopy. In summary, transects extended for 15–1,900 m across depths to 1.8 m.

At pre-designated points along transects, a suite of standardized, non-destructive measurements was taken within a  $1\text{-m}^2$  quadrat divided by strings into 100 cells, each 10 cm by 10 cm. The relevant measurement for DMA was a visual estimate of the biomass of all species expressed as an index, which was converted to biomass using empirically derived coefficients, i.e., 0 = 0 g dry weight (DW) m<sup>-2</sup>, 1 = 0.35 g DW m<sup>-2</sup>, 2 = 1.98 g DW m<sup>-2</sup>, 3 = 21.37 g DW m<sup>-2</sup>, 4 = 47.74 g DW m<sup>-2</sup>, 5 = 141.94 g DW m<sup>-2</sup> (Morris et al., 2001). For this study, we used data collected between 1998 and 2020.



#### **Fisheries Independent Monitoring**

Larger and in some cases deeper areas were sampled as part of long-term fisheries independent monitoring (FIM) conducted by the Florida Fish and Wildlife Conservation Commission, Fish and Wildlife Research Institute. The program sampled sites selected without replacement using a stratified random sampling design (FWC-FWRI, 2016), and we used data from sites sampled between either 1997 or 1998 and 2019.

The FIM program employed multiple types of gear, and we used data from two seines, a 21.3-m  $\times$  1.8-m center-bag haul seine with 3.2-mm nylon mesh and a 183-m  $\times$  2.5-m center-bag haul seine with 38-mm stretched nylon mesh. Seines were deployed along estuarine shorelines to sample areas with emergent vegetation, fringing mangroves, seawalls, and beaches or on flats at least 5 m from the shoreline. The 21.3-m seine was pulled for 9.1 m across a 15.5-m wide strip of bottom, resulting in a sampling area of approximately 140 m<sup>2</sup>. Using a boat, the 183-m seine was set in a rectangular shape adjacent to the shoreline with a maximum depth of 2.5 m at the bag before being retrieved by hand. The area sampled by the net (approximately 40 m  $\times$  103 m = 4,120 m<sup>2</sup>) was standardized by marking 40 m from each end to designate the corners of the rectangular set.

Bycatch of DMA was quantified to 0.1 gallons in the field, with reliable data being quantities of all species due to the challenges associated with identifying species visually. To convert these data to wet weights and dry weights, samples were collected during May and June 2012. Several species of DMA in quantities up to 5 gallons were collected at several sites, transferred to mesh bags, and spun to remove excess water. Mesh bags containing DMA were weighed to yield wet weights per gallon. The algae were dried at 80°C and then weighed to yield dry weights per gallon. A conversion factor was generated by fitting a linear regression, forced through zero. This conversion factor and the area sampled by the seine were combined to convert gallons of DMA to g DW m<sup>-2</sup>.

#### Hydroacoustic Surveys

To quantify the abundance and distribution of DMA found in deeper water, large-scale acoustic surveys were conducted between April and June in 2008, 2010, 2012, 2014, and 2015. While the surveys covered up to 288 km<sup>2</sup>, we focused analysis of spatiotemporal variation on reaches 2, 3, and 4, which were completed in all 5 years.

Hydroacoustic data were acquired with a BioSonics DT-X echosounder and two multiplexed, single-beam digital transducers with full beamwidths of  $10^{\circ}$  (38 kHz) and  $6.4^{\circ}$ (418 kHz), operated at 5-Hz and 0.4 ms pulse duration (Riegl and Bushkirk, 2016; Foster et al., 2018). Surveys focused on water deeper than approximately 1.3 m. The east-west survey lines were spaced a minimum of 200 m and maximum of 400 m apart. Data were post-processed using BioSonics Visual Bottom Typer (VBT; Foster et al., 2018). To estimate percent cover of DMA, samples of algae and video were collected along the line of the hydroacoustic surveys, and wet weights were recorded. Based on these samples, the acoustic signals were grouped into a supervised training catalog for bare substrate, sparse algae, dense algae, and other vegetation. Using the catalog and the estimated conversion factor, counts of the acoustic signals in each class were used to estimate wet biomass, and the conversion factor calculated for samples from seining was used to estimate dry biomass (Riegl and Bushkirk, 2016; Foster et al., 2018).

#### Analysis of Data

To evaluate differences in biomass, dry weights of macroalgae generated by each of the methods were used in separate permutation analyses of variance (PERMANOVAs, Anderson et al., 2008). All the models included reach and years as fixed factors, and models applied to data from transects and seines included seasons as a random factor nested in years. Seasons were defined based on the results of agglomerative hierarchical clustering of water quality data, with a warm wet season running from May to October and a cool dry season running from November in 1 year through April of the following year (Lasi et al., this volume). The PERMANOVAs were based on Bray– Curtis distance measures calculated with a dummy variable of 1 to avoid undefined results.

#### **Physiological Tolerances**

To evaluate the effects of stress caused by extreme salinities, extreme temperatures, low light levels, and combinations of stressors, a series of incubations was conducted under controlled conditions in tanks holding 70 L of water (Hanisak, 2016). Incubations were conducted in batch cultures using approximately 80 g wet weight of Gracilaria tikvahiae and Crassiphycus secundus (formerly Hydropuntia secunda), which are two of the most widespread and abundant rhodophytes in the IRL (Hanisak, 2016, 2021). Data were collected weekly over 35 days after 7 days of acclimation, and quantities of macroalgae were adjusted to maintain  $\sim$ 80 g wet weight. The potential effects of macronutrient and trace metal limitation were obviated by midday additions of 39.0 µM dissolved inorganic nitrogen (i.e., 26 µM ammonium and 13 µM nitrate), 0.94 µM orthophosphate, and trace metals at concentrations found in F/2 medium. These additions achieved a nitrogen:phosphorus ratio of  $\sim$ 41, which paralleled ratios measured in the IRL (Lasi unpublished data), and the  $\sim$ 38 mg of nitrogen and  $\sim$ 2 mg of phosphorus was sufficient to support growth of G. tikvahiae in past cultures (Hanisak, 1990). Salinities were maintained within 1 psu of the target by diluting water from a saltwater well. Temperatures were controlled within 1°C of the targets by chillers and heaters. Light was provided by 100W SOL 1 - LED Grow Lights from 06:00 to 18:00.

Two types of incubations were conducted: single-factor range finding incubations for salinity, temperature, and light availability and incubations manipulating combinations of factors. The factors that were not manipulated were held at 25 psu, 25°C, or 250  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>, as appropriate. Incubations with varying salinity included 12, 18, 25, 35, 45, and 50 psu. Temperatures of 7, 10, 17, 25, and 33°C were tested, and light levels of 0, 10, 30, 50, 100, 250, and 400 µmol m<sup>-2</sup> s<sup>-1</sup> were examined. Algae were harvested each week, and changes in wet weight were recorded. If incremental growth occurred, it was removed so the initial stocking density was returned to the tank. If biomass was lost, the decline was recorded. Wet weights were converted to dry weights using an empirical relationship derived from samples taken at the beginning and end of the incubations. In addition, samples taken each week were rinsed to remove excess salt, dried at 80°C, ground to a fine powder with a mortar and pestle, and shipped to the University of Maryland's Nutrient Analytical Services Laboratory for analysis of carbon, nitrogen, and phosphorus content (University of Maryland, 2022)<sup>1</sup>. The resulting percentages were used to identify any data that were affected by nutrient limitation.

Data on accumulation of biomass (growth) were extracted from seven sets of incubations and analyzed with PERMANOVAs based on Bray–Curtis distance measures with the addition of a dummy variable set at 1. Where necessary, adding the minimum to all values eliminated negative numbers. The design accounted for the repeated measures nature of the incubations. Species and the appropriate stressor or combination of stressors were fixed factors. Tank was a random factor nested in the interaction of species and treatment, and week was a random factor nested in the interaction of species, treatment, and tank. To examine the most extensive set of treatments, some data were used in more than one analysis.

#### **Timing and Extent of Suitable Conditions**

Based on the results from the evaluation of stressors, the timing and extent of suitable conditions for growth of DMA were predicted. The data to characterize stressors were drawn from ongoing sampling of water quality in the lagoon (Lasi et al., this volume) and existing bathymetry.

Salinities, water temperatures, and photosynthetically active radiation (PAR) were measured monthly at 30 fixed stations in reaches 1 through 6 (Figure 1). Measurements were taken with multiparameter instruments that were calibrated and verified with standards and procedures prescribed by state guidelines (Florida Department of Environmental Protection, 2017). Light availability was characterized by coefficients of attenuation for PAR ( $K_d$ ) as determined by applying Beer's Law (Kirk, 1983) to readings recorded simultaneously by three spherical quantum sensors. Sensors were held 0.2 and 0.5 m below the water's surface, and a third sensor was held 0.3 m from the bottom in water < 1.8 m deep or 1.5 m below the surface in deeper water (Morris et al., 2001). The resulting  $K_d$  values were combined with data on solar radiation from the Florida Automated Weather Network (FAWN<sup>2</sup>) (University of Florida, Institute of Food and Agricultural Sciences, 2022) to calculate the depth that received light above the threshold causing stress. Data from FAWN were converted from Watts  $m^{-2}$  to  $\mu$ mol  $m^{-2}$  s<sup>-1</sup> and daily means were calculated from daytime data for 1998 through 2019. The daily means were converted to irradiance  $(I_0)$  immediately below the surface of the water by reducing them by 6.6% to account for surface reflectance (Kirk, 1983), and the resulting values were combined with a threshold from the incubations  $(I_z)$  to determine the depth (z) receiving sufficient light using:

$$z = -\ln\left(I_Z/I_0\right)/K_d$$

The depths with sufficient light were translated to areal extent of suitable habitat for DMA by combining them with bathymetric data collected by Coastal Planning and Engineering (1997). Depth soundings were taken throughout the IRL at 15.2-m intervals along east–west transects that were separated by 150–300 m. This survey resulted in over 230,000 depth measurements that were referenced to North American Vertical Datum of 1988 (NAVD88) and adjusted to depth below mean water level (MWL). From these data, isobaths were generated in 0.1-m increments for reaches 1–6.

Stressful periods in each reach were identified by examining months with salinities, temperatures, or light availability that caused stress during the incubations. In addition, estimates of light penetration combined with bathymetry identified the extent of the lagoon with a suitable light regime.

## Carbon, Nitrogen, and Phosphorus Stored in Drift Algae

The carbon, nitrogen, and phosphorus contents of DMA were determined from samples of common rhodophytes collected bimonthly from six sites in the Indian River and Banana River lagoons (reaches 2, 3, and 4) between July 2014 and August

<sup>&</sup>lt;sup>1</sup>http://umces.edu/nutrient-analytical-services-laboratory

<sup>&</sup>lt;sup>2</sup>https://fawn.ifas.ufl.edu/

2015. Three replicate samples of the two most abundant species were collected from beyond the deep edge of the seagrass beds, typically in 1.5–2.0 m of water. The samples were transported on ice to the laboratory where epiphytes and debris were removed before the algae were rinsed to remove excess salt and dried at 80°C. Dried samples were ground to fine powder and analyzed for carbon, nitrogen, and phosphorus content at University of Maryland's Nutrient Analytical Services Laboratory (see Text Footnote 1).

The resulting data on carbon, nitrogen, and phosphorus content were used to gain insights into changes in the quantities of these elements stored in DMA. Median elemental contents were multiplied by DMA biomass per square meter from transects and surveys with seines to evaluate changes in cycling of key elements.

#### RESULTS

#### Patterns in Biomass of Drift Macroalgae

The biomass of the DMA assemblage varied significantly in space and through time according to analyses of data from surveys of transects, deployment of two different seines, and hydroacoustic surveys (**Table 1**). Biomasses from the transects and seines varied among combinations of reach and season within year, and biomasses detected with hydroacoustics varied significantly among years.

Higher values for the biomass of DMA observed during surveys of transects or as bycatch from seining did not reveal consistent and ecologically meaningful patterns across reaches or among years (**Figures 2A–D**). However, data from surveys of transects indicated that 70% of the biomasses above the 90th percentile were recorded in the warmer, wetter months of summer, and monthly data from seining indicated that the biomass of drift algae tended to be higher from March through July (means ± standard error for 21.3-m seine =  $1.7 \pm 0.2$  g m<sup>-2</sup> and 183-m seine =  $2.6 \pm 0.2$  g m<sup>-2</sup>) than in August–February (means ± standard error for 21.3-m seine =  $1.0 \pm 0.1$  g m<sup>-2</sup> and 183-m seine =  $1.2 \pm 0.1$  g m<sup>-2</sup>).

In contrast, relatively low biomass was recorded by multiple methods during two periods. Low biomass was documented consistently along the transects and in the seines from late 2010 through 2011–2012 in reaches 1–7, but the same pattern was not obvious in reaches 8 and 9 where the abundance of drift algae was lower (**Figures 2A,B**). Additionally, biomass detected by hydroacoustic surveys was lowest in 2012 (**Figure 2E**). Thus, multiple lines of evidence point to 2010–2012 as one period with little drift algae. In addition, drift algal biomass along transects and in the seines was consistently low in reaches 1–6 in 2016 (**Figures 2A,C,D**).

#### **Physiological Tolerances**

Evidence suggested that macroalgae were not limited by carbon, nitrogen, or phosphorus during the 35-day incubations. Cumulative growth of macroalgae over 35 days represented less than 2% of the 80 g incubated, so growth rates were unlikely to exhaust the nutrients, which were replenished

daily. In addition, elemental contents from all incubations were consistent, which indicated that the algae received sufficient carbon, nitrogen, and phosphorus throughout the incubations. The mean content  $\pm$  standard error was 28.169%  $\pm$  0.075% for carbon, 3.504%  $\pm$  0.026% for nitrogen, and 0.389%  $\pm$  0.004% for phosphorus.

Incubations with salinities that spanned 12–50 psu did not identify significant effects on cumulative growth or elemental contents (**Table 2**). On average for each of the salinities, *G. tikvahiae* accumulated  $5.1 \pm 1.4$  g to  $10.2 \pm 1.2$  g of biomass over 35 days, and *C. secundus* accumulated  $4.6 \pm 0.8$  g to  $9.2 \pm 1.4$  g of biomass. Given these results, the effect of salinity was not explored further.

The two species had differing responses to the varying temperatures used in a range finding incubations (**Table 2**). At all temperatures, *G. tikvahiae* accumulated more biomass over 35 days (**Figure 3A**). Both *G. tikvahiae* and *C. secundus* accumulated slightly more biomass at 25°C, although growth at 33°C was similar (**Figure 3A**). On average, *C. secundus* lost biomass at 7°C and only gained  $0.8 \pm 0.1$  g at 10°C.

Light availability characterized by PAR affected the growth of *G. tikvahiae* and *C. secundus* similarly as shown by the lack of a significant interaction between species and amount of PAR (**Table 2**). Unlike *C. secundus*, *G. tikvahiae* did not survive in  $0 \mu \text{mol m}^{-2} \text{ s}^{-1}$ . Both species consistently accumulated biomass over 35 days only when provided with 250 or 400  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> of PAR (**Figure 3B**). On average, *G. tikvahiae* accumulated 2.5 ± 1.2 g of biomass, and *C. secundus* accumulated 3.7 ± 0.7 g of biomass. Given the greater and less variable accumulation of biomass, *C. secundus* appeared to cope with low light better than *G. tikvahiae*.

Two sets of results combined differing temperatures and differing amounts of light, with some data for G. tikvahiae used in two PERMANOVAs (Table 2). In the first set of results, G. tikvahiae and C. secundus were incubated at 7, 10, 17, 25, and 33°C under 10, 50, and 100  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> of PAR, and in the second set, only G. tikvahiae was available to be incubated at 7, 10, 17, 25, and 33°C under 10, 50, 100, and 250 µmol  $m^{-2} s^{-1}$  of PAR. The two species exhibited differing responses to combinations of temperature and PAR (Table 2). For both species, there was loss or accumulation of <2 g of biomass over 35 days at 10  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> of PAR (**Figures 3C–E**). At 50  $\mu$ mol  $m^{-2}$  s<sup>-1</sup> of PAR over 35 days, G. tikvahiae gained 2-3 g of biomass at all temperatures except 25°C, and C. secundus gained about 2 g of biomass at 17 and 33°C (Figures 3C-E). At 100 µmol m<sup>-2</sup> s<sup>-1</sup> of PAR over 35 days, G. tikvahiae gained 2-6 g of biomass, with the maximum gain at 25°C (Figures 3C,E). At 100  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> of PAR, C. secundus lost biomass at 7 and 10°C and gained approximately 3 g over 35 days at 17, 25, and 33°C (Figure 3D). Only G. tikvahiae was subjected to differing temperatures at 250  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>, and it gained a maximum of  $9.1 \pm 0.7$  g of biomass over 35 days at 25°C, with gains decreasing to  $3.4 \pm 0.8$  g of biomass at 7°C and  $7.5 \pm 1.2$  g of biomass at 33°C (Figure 3E). Both species tended to lose biomass when subjected to less than 50  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> of PAR although they survived the 35-day incubations. Loss of biomass was more common at temperatures of 7 or 10°C. Overall, the results indicated that light

TABLE 1 Permutation analyses of variance for biomass of drift macroalgae sampled with four methods
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Survey	Source	df	SS	MS	Pseudo-F ratios	р	Unique permutations
Transects	Rch	8	457,970	57,246	24.58	0.001	997
	Yr	20	266,800	13,340	3.32	0.003	998
	Seas(Yr)	21	84,399	4,019	3.05	0.001	998
	$\operatorname{Rch} \times \operatorname{Yr}$	160	463,520	2,897	1.24	0.063	997
	$Rch \times Seas(Yr)$	168	391,780	2,332	1.77	0.001	995
	Res	2,723	3,582,500	1,316			
	Total	3,100	5,450,000				
21.3-m seine	Rch	4	103,030	25,758	27.02	0.002	500
	Yr	20	72,398	3,620	2.53	0.012	500
	Seas(Yr)	21	30,131	1,435	3.25	0.002	500
	$Rch \times Yr$	80	99,741	1,247	1.28	0.108	499
	$Rch \times Seas(Yr)$	84	82,110	978	2.21	0.002	499
	Res	8,341	3,681,100	441			
	Total	8,550	4,086,600				
183-m seine	Rch	3	132,610	44,204	28.27	0.001	998
	Yr	22	173,380	7,881	2.77	0.007	999
	Seas(Yr)	23	65,587	2,852	3.63	0.001	999
	$Rch \times Yr$	66	123,870	1,877	1.19	0.241	996
	$Rch \times Seas(Yr)$	69	108,950	1,579	2.01	0.001	995
	Res	5,005	3,930,000	785			
	Total	5,188	4,602,700				
Hydroacoustics	Rch	2	1,081	540	1.90	0.127	998
	Yr	4	2,655	664	2.34	0.047	998
	$\operatorname{Rch} \times \operatorname{Yr}$	8	2,985	373	1.32	0.250	999
	Res	40	11,348	284			
	Total	54	24,267				

Rch, reach; Yr, year; Seas, season; Res, residual.

availability was a dominant influence on growth, with an effect of temperature appearing more strongly when the algae received over 100  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> of PAR. In those conditions, optimum temperatures appeared to be 25°C for *G. tikvahiae* and 17–33°C for *C. secundus*.

#### **Timing and Extent of Suitable Conditions**

Salinity did not stress either G. tikvahiae or C. secundus, but incubations in the laboratory indicated that low temperatures (below 17°C), high temperatures (above 25-33°C), and reduced light availability (less than 100  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>) had the potential to reduce growth or cause mortality. Stressful periods for each reach were identified by coding monthly mean  $K_d$  values higher than the 90th percentile of all values (>1.6 m<sup>-1</sup>), monthly mean temperatures above the 90th percentile (>30.3°C), and monthly mean temperatures below the 10th percentile (<17.8°C) as zero and all other values as one to produce a heatmap (Figure 4). Mean temperatures exceeded the 90th percentile in summers of all years, and temperatures dropped below the 10th percentile in multiple winters. The period with the most spatially and temporally extensive distribution of all stressors was 2009-2011, which corresponded with multiple records of low abundance for DMA (Figure 2). In samples taken with the 21.3-m seine from June 2010 through May 2012, the mean biomass of DMA in all reaches was below the 10th percentile of all values 67% of the time and below the 25th percentile of all values 88% of the time. Similarly, in samples taken with the 183-m seine from August 2010 through October 2012, the mean biomass of DMA in all reaches was below the 10th percentile of all values 56% of the time and below the 25th percentile of all values 78% of the time. High  $K_d$  values that translated into low light availability also were noted in 2016 and 2018 along with warm temperatures, which again corresponded with low abundance of DMA, especially in reaches 3-6 (Figures 2A,C,D). In samples taken with the 21.3m seine from August 2015 through November 2016, the mean biomass of DMA in all reaches was below the 10th percentile of all values 12% of the time and below the 25th percentile of all values 50% of the time. For the 183-m seine, the mean biomass of DMA recorded in all reaches from September 2016 through December 2016 was below the 10th percentile of all values 50% of the time and below the 25th percentile of all values 100% of the time.

The incubations to identify tolerances highlighted PAR below 100  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> as a key influence on the growth of DMA; therefore, depths receiving  $\geq$ 100  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> were associated with 0.1-m isobaths to identify the extent of suitable habitat in hectares (**Figure 5** and **Supplementary Figure 1**). The extent of suitable habitat varied through time for each reach, and when compared to the background extent from 1998 to 2009, reaches 2–6 had more substantial losses of suitable habitat in 2010–2011 and in 2016–2018, which would translate to changes in



the amounts of carbon, nitrogen, and phosphorus stored in DMA (**Figure 6** and **Supplementary Table 2**). These results were consistent with other evaluations of the abundance of DMA.

# Carbon, Nitrogen, and Phosphorus Stored in Drift Algae

Five species of DMA were collected from the field, *Acanthophora* spicifera, Agardhiella subulata, G. tikvahiae, C. secundus, and Hypnea spinella. A series of PERMANOVAs using elemental

compositions of the three species collected multiple times, *A. subulata*, *G. tikvahiae*, and *C. secundus*, identified significant variation among the combination of species and sampling event for mean percent carbon  $(p_{10,212} = 0.001)$ , nitrogen  $(p_{10,212} = 0.001)$ , and phosphorus  $(p_{10,212} = 0.002)$ . Mean carbon content ±SE varied from 16.6 ± 0.6% for *A. subulata* in July 2014 to 33.9 ± 0.4% for *C. secundus* in April 2015, mean nitrogen content ± SE varied from 1.5 ± 0.1% for *A. subulata* in August 2015 to 3.6 ± 0.1% for *C. secundus* in April 2015, and mean phosphorus content ± SE varied from 0.04 ± 0.003% for

Stressor	Source	df	SS	MS	Pseudo-F ratios	p	Unique permutations
Sal	Spp	1	705	705	1.64	0.213	999
	Sal	5	2,486	497	1.15	0.392	999
	$Spp \times Sal$	5	4,795	959	2.23	0.082	999
	$Tnk(Spp \times Sal)$	12	5,169	431	1.50	0.115	999
	$Wk[Tnk(Spp \times Sal)]$	72	20,728	288	No test		
	Total	95	33,883				
Temp	Spp	1	6,825	6,825	40.33	0.001	998
	Temp	3	9,137	3,046	18.00	0.001	999
	Spp × Temp	3	3,868	1,289	7.62	0.001	997
	Tnk(Spp × Temp)	18	3,046	169	0.59	0.975	999
	Wk[Tnk(Spp × Temp)]	78	22,211	285	No test		
	Total	103	43,492				
PAR	Spp	1	5,400	5,400	22.79	0.001	999
	PAR	5	15,172	3,035	12.81	0.001	999
	$Spp \times PAR$	5	2,011	402	1.70	0.119	998
	$Tnk(Spp \times PAR)$	12	2,843	237	0.53	0.981	999
	$Wk[Tnk(Spp \times PAR)]$	72	32,462	451	No test		
	Total	95	57,889				
Temp × PAR	Spp	1	1,056	1,056	34.47	0.001	999
	Temp	4	2,276	569	18.57	0.001	998
	PAR	2	3,143	1,572	51.30	0.001	999
	Spp × Temp	4	2,220	555	18.11	0.001	999
	$Spp \times PAR$	2	698	349	11.39	0.001	999
	Temp $\times$ PAR	8	3,396	425	13.86	0.001	999
	Spp $\times$ Temp $\times$ PAR	8	873	109	3.56	0.001	998
	$Tnk(Spp \times Temp \times PAR)$	62	1,900	31	0.38	1.000	995
	$Wk[Tnk(Spp \times Temp \times PAR)]$	276	22,470	81	No test		
	Total	367	38,282				
Temp × PAR	Temp	4	1,886	471	7.32	0.001	999
	PAR	3	8,438	2,813	43.68	0.001	999
	Temp $\times$ PAR	12	4,257	355	5.51	0.001	998
	$Tnk(Temp \times PAR)$	44	2,834	64	0.63	0.994	997
	Wk[Tnk(Temp $\times$ PAR)]	192	19,778	103	No test		
	Total	255	39,914				

Sal, salinity; Temp, temperature; PAR, amount of light; Spp, species; Tnk, tank; Wk, week.

*C. secundus* in April 2015 to  $0.2 \pm 0.006\%$  for *G. tikvahiae* in August 2014 (**Supplementary Table 3**). These results pointed to the value of expanded surveys to determine temporal, spatial, and interspecific variation in elemental compositions of DMA. These data can be used to make multiple estimates of changes in the amounts of carbon, nitrogen, and phosphorus stored in DMA, and as an example to illustrate the importance of such changes, the overall median percent compositions were multiplied by the relevant dry weights.

Carbon (g) = DMA biomass  $(g DW m^{-2}) \times 0.2840$ Nitrogen (g) = DMA biomass  $(g DW m^{-2}) \times 0.0224$ Phosphorus (g) = DMA biomass  $(g DW m^{-2}) \times 0.0008$ 

Based on mean annual biomass of DMA along transects, all reaches had less carbon, nitrogen, and phosphorus stored in DMA tissues in 2020 when compared to the amounts documented from 1998 to 2009 (**Table 3**). The losses of biomass ranged from 21% in reach 9 to 93% in reach 4 (**Table 3**). The decreases suggested that carbon, nitrogen, and phosphorus became available for uptake by other primary producers, such as phytoplankton.

## DISCUSSION

Biomass of DMA varied significantly in space and through time, widespread decreases in biomass were related to thresholds for environmental tolerances as determined by incubations in the laboratory and field data, and less DMA translated into less carbon, nitrogen and phosphorus being stored in DMA. High biomass was recorded in different seasons, years, and reaches by the different methods, but low biomass was recorded by multiple sampling methods during 2010–2012 and in 2016, especially in reaches 1 through 6 where DMA was more common and



percentiles, horizontal line indicates the median, X indicates the mean.

abundant. Both these reductions in biomass of DMA followed periods of low light availability, and the 2010–2012 change also followed periods with potentially stressful low and high temperatures. Incubations indicated that light availability was a dominant influence, with less than ~100  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> leading to stress, and based on this threshold, there were widespread decreases in suitable habitat in 2010–2012, 2016, and 2018. During these periods, less carbon, nitrogen, and phosphorus were stored in DMA, with more of these elements likely to be available for uptake by other primary producers.

Although the biomass of DMA in the lagoon did not vary consistently between seasons in most years, it tended to be

more abundant in March–July, and given the differences in sampling intensity and methodology, these results correspond well with previous surveys in the Indian River Lagoon (Benz et al., 1979; Kulczycki et al., 1981; Virnstein and Carbonara, 1985; Hanisak, 2021). Differences among the reports tended to occur in the cooler months, which was not unexpected given interannual variability in water temperatures. This temporal pattern was disrupted in 2010–2012 and 2016, with reduced light availability and stressful temperatures being apparent influences. The decreased availability of light coincided with phytoplankton blooms that increased light attenuation coefficients above the 90th percentiles for 6–8 months in all reaches during the



28 months from July 2010 to October 2012 and 6–12 months in reaches 1, 2, and 3 during the 18 months from July 2015 to December 2016 (Phlips et al., 2021; Lasi et al., this volume). A similar response to reductions in light availability caused by phytoplankton was noted in the coastal waters off Denmark (Nielsen et al., 2002).

The tolerances identified in incubations in the laboratory agreed with previous reports for various species of Gracilaria and species labeled as Hydropuntia. For example, five species of Gracilaria grew or photosynthesized in salinities ranging from <20 to >40 psu (Lapointe and Hanisak, 1985; Engledow and Bolton, 1992; Yokoya et al., 1999; Orduña-Rojas et al., 2013; de Paiva Alves et al., 2021), but Gracilaria textorii and Gracilaria vermiculophylla from Japan did not survive in 10 or 15 psu (Raikar et al., 2001) so adaptation to local conditions may have been an important influence. Similarly, growth or net photosynthesis typically was recorded across a broad range of temperatures (8-42°C) for fourteen species of Gracilaria and two species of Hydropuntia, with most species doing best at 20-30°C (Lapointe et al., 1984; Lapointe and Hanisak, 1985; Hanisak, 1987; Engledow and Bolton, 1992; Yokoya and Oliveirã, 1992; Dawes et al., 1998; Yokoya et al., 1999; Raikar et al., 2001; Abreau et al., 2011; Nejrup et al., 2013; Orduña-Rojas et al., 2013; Vo et al., 2015; Castro and Yokoya, 2019). Growth and photosynthesis also varied with available light for six species of Gracilaria and two species of Hydropuntia, with algae surviving when held at  $<40 \ \mu mol \ m^{-2} \ s^{-1}$ , exhibiting little increase in

growth rate or photosynthetic rate above  $\sim 100 \ \mu mol \ m^{-2} \ s^{-1}$ , and displaying an ability to cope with more extreme temperatures when sufficient light was available (Lapointe et al., 1984; Lapointe and Hanisak, 1985; Hanisak, 1987; Engledow and Bolton, 1992; Dawes et al., 1998; Yokoya et al., 1999; Abreau et al., 2011; Nejrup et al., 2013; Orduña-Rojas et al., 2013; Vo et al., 2015; Castro and Yokoya, 2019).

Decreases in the biomass of DMA reduced storage of carbon, nitrogen, and phosphorus in this pool, and these elements should have become available to other primary producers. As an example, the amounts stored in DMA were calculated by multiplying the total amount of biomass in reaches 2-6 as estimated from the 2015 hydroacoustic survey (16,000 MT dry weight) by the conversion factors derived from samples of DMA. These calculations yielded estimates of 4,400 MT of carbon, 410 MT of nitrogen, and 13 MT of phosphorus stored in DMA. Applying the mean change in the biomass of DMA along transects in reaches 2-6 calculated for 2019 (-38%; Table 3) to these values indicated that 2,800 MT of carbon, 260 MT of nitrogen, and 8 metric tons of phosphorus would have become available. Given the volumes of reaches 2-6 and an assumption of no uptake, concentrations of carbon, nitrogen, and phosphorus in the water column would have increased by 1.419, 0.132, and 0.004 mg  $L^{-1}$ , respectively. These changes represented 14, 14, and 18% of the mean concentrations of dissolved organic carbon, dissolved nitrogen, and dissolved phosphorus in reaches 2-6 from 1997



FIGURE 5 | Extent of suitable light regime for drift macroalgae in a portion of reaches 1 and 2, with the minimum extent from 1998 to 2020, background extent from 1998 to 2009, and maximum extent from 1998 to 2020 displayed. Topographic map source: ESRI, USGS, contour lines represent 0.1-m isobaths.

to 2009, respectively (St. Johns River Water Management District, unpub. data). Thus, DMA played an important role in the cycling of elements in the lagoon that managers should consider.

Given that DMA play an important role in cycling of elements in the lagoon, managers could consider harvesting DMA to remove nitrogen and phosphorus and prevent shading of seagrass (Virnstein and Carbonara, 1985; Sfriso et al., 2020). Such an action demands careful consideration because removing DMA may increase the quantities of carbon, nitrogen, and phosphorus available to phytoplankton, but this issue could be addressed by timing the harvest to the period following maximum growth (Braun, 2020). In addition, less DMA translates into less structural habitat in the lagoon (Gore et al., 1981; Kulczycki et al., 1981; Virnstein and Howard, 1987; Holmquist, 1997; Monagail et al., 2017). For example, evidence has suggested that loss of DMA reduces refuge from predation for many macroinvertebrates (Heck, 1979; Heck and Thoman, 1981; Stoner, 1985; Stoner and Lewis, 1985; Edgar, 1987) and alters the composition of assemblages of benthic invertebrates due to species specific responses (Norkko et al., 2000).



TABLE 3 | Estimated carbon (C), nitrogen (N), and phosphorus (P) found in drift macroalgae along transects.

Period	Parameter	Reach								
		1	2	3	4	5	6	7	8	9
1998–2009	Mean C (g m <sup>-2</sup> )	1.562	3.364	2.480	1.963	1.891	2.875	4.747	1.927	0.848
	Mean N (g m <sup>-2</sup> )	0.123	0.265	0.196	0.155	0.149	0.227	0.374	0.152	0.067
	Mean P (g m <sup>-2</sup> )	0.004	0.009	0.007	0.006	0.005	0.008	0.013	0.005	0.002
2020	Mean C (g m <sup>-2</sup> )	1.127	1.618	0.263	0.133	0.698	2.223	1.570	0.497	0.669
	Mean N (g m <sup>-2</sup> )	0.089	0.128	0.021	0.010	0.055	0.175	0.124	0.039	0.053
	Mean P (g m <sup>-2</sup> )	0.003	0.005	0.001	0.000	0.002	0.006	0.004	0.001	0.002
Change	(%)	-28	-52	-89	-93	-63	-23	-67	-74	-21

Overall, management of the lagoon would be improved by a better understanding of the ecological roles played by DMA and attached macroalgae. Beyond elucidating its roles in cycling of elements and as a habitat, regular surveys that document the distribution, abundance, and diversity of DMA, experiments to elucidate the drivers of changes in the distribution and abundance of DMA more fully, and experiments to untangle the relationship between DMA and seagrass would supply valuable information to decision makers. In all cases, effort needs to be expanded to additional species of DMA and due consideration should be given to attached macroalgae. For example, DMA in reaches 2 and 4 during 2014-2015 were dominated by Chaetomorpha sp., large blooms of this green alga have blanketed seagrass in other locations (Pulich et al., 1997; Kennish et al., 2010; Gao et al., 2013), and approximately 35% of drift algae in parts of the lagoon are not Gracilaria (Hanisak, 2021). An increased likelihood of such blooms could be driven by climate change because green algae tolerated relatively high temperatures (Menéndez and Comín, 2000). Furthermore, blooms of DMA have been shown to shade seagrasses or self-shade, which increased concentrations of nutrients in the water column due to decomposition (Holmquist, 1994, 1997; Hauxwell et al., 2001; Cummins et al., 2004; Gao et al., 2013; Foster et al., 2018). Additionally, Fox et al. (2008) stated that Cladophora vagabunda and G. tikvahiae in locations with higher nitrogen loads from their watersheds stored up to 250% of the annual load. Similar roles may be played by attached algae. For example, Caulerpa prolifera has been documented as the dominant vegetation in the northern Indian River Lagoon and in deeper areas in Banana River Lagoon (White and Snodgrass, 1990; Provancha and Scheidt, 2000), and this species has demonstrated an ability to take up significant amounts of ammonium (Alexandre and Santos, 2020) so its role in cycling of nutrients should be considered.

In conclusion, reduced light availability and extreme temperatures in the IRL likely contributed to decreased growth or mortality of DMA. In turn, less DMA likely promoted phytoplankton blooms because more carbon, nitrogen, and phosphorus became available. The presence of DMA in the IRL has been recognized for decades (Thompson, 1978), but an increased understanding of its roles will support improved management of the system.

#### DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation, to any qualified researcher.

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#### **AUTHOR CONTRIBUTIONS**

All authors contributed to designing components of the study. MH conducted the incubations to determine tolerances and generated data on elemental composition. provided valuable RP and BR biomass data. CI performed statistical analysis. LH and LM drafted the manuscript.

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## SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars. 2022.767440/full#supplementary-material

Supplementary Figure 1 | Extent of suitable habitat for drift macroalgae in hectares based on mean depths receiving 100  $\mu mol \ m^{-2} \ s^{-1}.$  Note different scales.

Supplementary Table 1 | Metadata for each method of sampling biomass of drift macroalgae.

**Supplementary Table 2 |** Estimates of carbon, nitrogen, and phosphorus found in drift macroalgae along transects extrapolated to extent of suitable habitat and reach volume.

**Supplementary Table 3 |** Elemental contents of common rhodophytes collected between July 2014 and August 2015.

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