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Optimization and assessment of phytoplankton size class algorithms for ocean color data on the Northeast U.S. continental shelf

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Abstract

The size structure of phytoplankton communities influences important ecological and biogeochemical processes, including the transfer of energy through marine food webs. A variety of algorithms have been developed to estimate phytoplankton size classes (PSCs) from satellite ocean color data. However, many of these algorithms were developed for application to the global ocean, and their performance in more productive, optically complex coastal and continental shelf regions warrants evaluation. In this study, several existing PSC models were applied in the Northeast U.S. continental shelf (NES) region and compared with in situ PSC estimates derived from a local HPLC pigment data set. The effect of regional re-parameterization and incorporation of sea surface temperature (SST) into existing abundance-based model frameworks was investigated and model performance was assessed using an independent data set. Abundance-based model re-parameterization alone did not result in significant improvement in model performance compared with other models. However, the inclusion of SST led to a consistent reduction in model error for all size classes. Of two absorption-based algorithms tested, the best performing approach displayed similar performance metrics to the regional SST-dependent abundance-based model. The SST-dependent model and the absorption-based method were applied to monthly composites of the NES region for April and September 2019 and qualitatively compared. The results highlight the benefit of considering SST in abundance-based models and the applicability of absorption-based PSC methods in optically complex regions.
### Table 1. Symbols and definitions.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>$a_d$</td>
<td>absorption coefficient of non-algal particles</td>
<td>m$^{-1}$</td>
</tr>
<tr>
<td>$a_{dg}$</td>
<td>absorption coefficient of colored dissolved organic matter + non-algal particles</td>
<td>m$^{-1}$</td>
</tr>
<tr>
<td>$a_g$</td>
<td>absorption coefficient of colored dissolved organic matter</td>
<td>m$^{-1}$</td>
</tr>
<tr>
<td>$[\text{Allo}]$</td>
<td>alloxanthin concentration</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>$a_{ph}$</td>
<td>absorption coefficient of phytoplankton</td>
<td>m$^{-1}$</td>
</tr>
<tr>
<td>$a_{ph}^*$</td>
<td>chlorophyll-specific absorption coefficient of phytoplankton</td>
<td>m$^2$ mg$^{-1}$</td>
</tr>
<tr>
<td>$[\text{But-fuco}]$</td>
<td>19'-butanoyloxyfucoxanthin concentration</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>$[\text{Chl-a}]$</td>
<td>chlorophyll-a concentration</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>$C_{DP}$</td>
<td>chlorophyll-a concentration reconstructed from the weighted sum of diagnostic pigments</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>$C_{HPLC}$</td>
<td>chlorophyll-a concentration measured by high-performance liquid chromatography</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>$C_{size}$</td>
<td>chlorophyll-a concentration specific to size class “size”</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>$C_{m_{size}}$</td>
<td>asymptotic maximum chlorophyll-a concentration of size class “size”</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>$D_{size}$</td>
<td>fraction of size class “size” as total chlorophyll-a tends to zero</td>
<td>unitless</td>
</tr>
<tr>
<td>$F_{size}$</td>
<td>fraction of size class “size”</td>
<td>unitless</td>
</tr>
<tr>
<td>$[\text{Fuco}]$</td>
<td>fucoxanthin concentration</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>GB</td>
<td>Georges Bank</td>
<td>–</td>
</tr>
<tr>
<td>GoM</td>
<td>Gulf of Maine</td>
<td>–</td>
</tr>
<tr>
<td>$[\text{Hex-fuco}]$</td>
<td>19'-hexanoyloxyfucoxanthin concentration</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>MAB</td>
<td>Middle Atlantic Bight</td>
<td>–</td>
</tr>
<tr>
<td>MAD</td>
<td>mean absolute difference</td>
<td>unitless</td>
</tr>
<tr>
<td>NES</td>
<td>northeast U.S. continental shelf</td>
<td>–</td>
</tr>
<tr>
<td>$[\text{Perid}]$</td>
<td>peridinin concentration</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>PSC</td>
<td>phytoplankton size class</td>
<td>–</td>
</tr>
<tr>
<td>$r$</td>
<td>Pearson correlation coefficient</td>
<td>unitless</td>
</tr>
<tr>
<td>$R_{rs}$</td>
<td>remote sensing reflectance</td>
<td>sr$^{-1}$</td>
</tr>
<tr>
<td>$S$</td>
<td>slope of a Type-II linear regression</td>
<td>unitless</td>
</tr>
<tr>
<td>$SST$</td>
<td>sea surface temperature</td>
<td>ºC</td>
</tr>
<tr>
<td>SFF</td>
<td>size-fractionated filtration</td>
<td>–</td>
</tr>
<tr>
<td>$[\text{TAcc}]$</td>
<td>total concentration of accessory pigments</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>$[\text{TCChl-b}]$</td>
<td>total chlorophyll-b concentration</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>$[\text{Zea}]$</td>
<td>zeaxanthin concentration</td>
<td>mg m$^{-3}$</td>
</tr>
<tr>
<td>$\delta$</td>
<td>bias</td>
<td>unitless</td>
</tr>
</tbody>
</table>
1. Introduction

Phytoplankton form the base of pelagic food webs and are a key component in biogeochemical cycles that ultimately impact global climate (Behrenfeld et al., 2006; Field et al., 1998). Phytoplankton in the ocean are taxonomically diverse and exhibit an array of unique morphological and physiological characteristics (Caron et al., 2012; Finkel et al., 2010). Phytoplankton community composition and biomass are highly variable in time and space, changing in response to both bottom-up (e.g., environmental conditions, nutrient availability) and top-down (e.g., grazing) controls. Understanding the dynamics of phytoplankton abundance and community structure is critical for studies of marine ecology and biogeochemistry.

Functional traits or groupings have been applied as a means of characterizing phytoplankton communities (IOCCG, 2014; Le Quéré et al., 2005; Nair et al., 2008). Phytoplankton cell size is considered a fundamental trait as it affects many important biological and ecological processes, including photosynthesis (Uitz et al., 2008), nutrient uptake (Raven, 1998), growth rate (Marañón, 2015), light absorption (Bricaud et al., 2004; Ciotti et al., 2002), carbon export (Guidi et al., 2009; Mouw et al., 2016), and the transfer of energy to higher trophic levels (Boyce et al., 2015). Thus, the size structure of phytoplankton assemblages can serve as a valuable indicator of the state of marine ecosystems and their response to environmental changes, including ocean warming (Marinov et al., 2010; Morán et al., 2010; Platt and Sathyendranath, 2008). Following the classification of Sieburth et al. (1978), phytoplankton are conventionally partitioned into three phytoplankton size classes (PSCs; see Table 1 for a list of symbols and
abbreviations used in this text): picoplankton (0.2-2 µm), nanoplankton (2-20 µm) and microplankton (20-200 µm).

A number of methods exist for quantifying PSCs *in situ*, including microscopy, size-fractionated filtration (SFF), conventional and imaging flow cytometry (Olson and Sosik, 2007), and high-performance liquid chromatography (HPLC) pigment analysis, each with their own advantages and limitations (IOCCG, 2014). While these methods are extremely useful, they can be labor-intensive, time consuming, and expensive, and as a result, the availability of *in situ* PSC data remains limited, thus limiting their use in studying and modeling large-scale, temporally dynamic ocean and ecosystem processes. Satellite remote sensing enables frequent, synoptic coverage of upper ocean optical properties and provides a means to characterize PSCs at spatial and temporal resolutions unattainable with *in situ* sampling. Given this fact, deriving information on phytoplankton size structure from satellite ocean color data is an active area of research, and a variety of algorithms have been developed for both global ocean (Brewin et al., 2015; Hirata et al., 2011) and regional applications (Brito et al., 2015; Di Cicco et al., 2017; Gittings et al., 2019; Sun et al., 2018, 2019a). Most current approaches for detecting PSCs from remote sensing can be categorized as either abundance-based or absorption-based, differing in terms of their theoretical frameworks and the remotely sensed parameters used as inputs (IOCCG, 2014; Mouw et al., 2017b).

Abundance-based algorithms exploit the generally observed co-variance of phytoplankton cell size and total biomass, indexed by its proxy, chlorophyll-a concentration ([Chl-a]; mg m⁻³), to estimate the dominance or relative biomass fractions of PSCs (Brewin et al., 2010; Hirata et al., 2011; Uitz et al., 2006). For example, it is well
established that picoplankton, such as the cyanobacteria *Prochlorococcus* and *Synechococcus*, dominate in low-nutrient oligotrophic environments (e.g., central ocean gyres) and microplankton, including diatoms and dinoflagellates, comprise a greater fraction of total biomass in eutrophic, nutrient-rich regions (e.g., upwelling zones) (Chisholm et al., 1988; Margalef, 1978; Yentsch and Phinney, 1989). Abundance-based methods rely on empirical or semi-empirical relationships based on coincident in situ observations of size-fractionated [Chl-a], typically from HPLC pigments or SFF, and total [Chl-a] to model PSCs as a function of total [Chl-a]. Given that [Chl-a] is perhaps the most widely used and well-validated satellite ocean color product, abundance-based methods offer a straightforward, “user-friendly” approach for estimating PSCs from remote sensing. Yet, these methods are an indirect approximation of size structure, and the empirical relationships underlying them are subject to change both temporally and geographically, requiring ongoing assessment and re-calibration (Mouw et al., 2017b). Recent studies have demonstrated that the incorporation of additional environmental information attainable from remote sensing, such as sea surface temperature (SST), can improve the retrieval accuracy of abundance-based models (Brewin et al., 2017a; Moore and Brown, 2020; Ward, 2015).

Absorption-based algorithms distinguish PSCs directly from spectral variations in phytoplankton absorption ($a_{ph}(\lambda)$; $\lambda$ denotes wavelength dependence), the amount of light absorbed by phytoplankton across the visible spectrum, which influences and can be derived from the reflectance signal measured by a satellite ocean color sensor (remote sensing reflectance ($R_{rs}(\lambda)$)) (Ciotti and Bricaud, 2006; Devred et al., 2011; Mouw and Yoder, 2010). Larger cells are less efficient at absorbing light than smaller cells as a result
of pigment packaging, which results in a flattening of the chlorophyll-normalized phytoplankton absorption spectrum \( a_{ph}(\lambda) \) with increasing cell size, with the most pronounced change around 440 nm (Bricaud et al., 1988; Morel, 1987; Morel and Bricaud, 1981). Ciotti et al. (2002) demonstrated that despite physiological and taxonomic variability, cell size could explain >80% of the variance in \( a_{ph}(\lambda) \) spectral shape over the wavelength range 400-700 nm. An advantage of absorption-based methods over abundance-based approaches is that they can detect changes in PSCs that do not co-vary with total \([Chl-a]\) (e.g., blooms of smaller cells). Moreover, as absorption-based methods are based on a direct optical response rather than indirect empirical relationships, they are less likely to require re-calibration over time or for different ocean regions. However, the limited spectral resolution of current multi-spectral ocean color instruments and inherent ambiguity in the optical signals of in-water constituents can make retrieving accurate \( a_{ph}(\lambda) \) spectral shape and magnitude challenging. This is particularly true in optically complex coastal and continental shelf waters with high concentrations of colored dissolved organic matter (CDOM) and non-algal particles (NAP), which overlap with phytoplankton in their contribution to the total light absorption in the blue region of the spectra, where satellite retrievals are subject to large uncertainties related to atmospheric correction and issues with sky and sun glint (Estrella et al., 2020).

Given the unique strengths and limitations of these different satellite PSC algorithm approaches, evaluating their performance in different ocean regions and whether they may be optimized for regional application is essential. Several studies have successfully retrieved PSCs at regional scales, including the Red Sea (Gittings et al., 2019), the
Mediterranean Sea (Di Cicco et al., 2017), the Western Iberian coast (Brito et al., 2015), and the continental shelf seas of China (Sun et al., 2018, 2019a) through re-parameterization of global abundance-based models with local in situ data sets. These studies demonstrate the potential for PSC model optimization in other ocean regions, which may provide benefits for regional-scale applications, including food web modeling and ecosystem-based fisheries management.

The northeast U.S. continental shelf (NES) is a highly productive, temperate marine ecosystem that supports many commercially and recreationally important fisheries (Hare et al., 2016; National Marine Fisheries Service, 2018). The NES is physically dynamic and optically complex (Mannino et al., 2014; Pan et al., 2008), thus necessitating evaluation and potential optimization of existing PSC algorithms to ensure their accuracy. Phytoplankton species composition and biomass in the NES vary seasonally, with diatoms dominating in a typical winter-spring bloom and other taxa, such as cryptophytes and cyanobacteria, becoming more prevalent during the summer (O'Reilly and Zetlin, 1998; Pan et al., 2011; Richaud et al., 2016).

In this study, we aim to evaluate and optimize several existing abundance-based and absorption-based PSC algorithms for application to ocean color imagery in the NES region, with the goal of improving PSC imagery products for long-term time series investigations and integration into regional ecosystem and fisheries modeling efforts. Primarily, we address the following questions:

- To what extent does regional re-parameterization using a local in situ data set improve the performance of abundance-based PSC models in the NES?
• Does incorporating SST into an abundance-based model improve PSC retrieval accuracy in the NES?
• How do abundance-based and absorption-based algorithms compare in their estimations of PSCs in the NES?

2. Data and methods

2.1 Study area

The NES region (35°N-45.5°N, 64°W-77°W) extends along the east coast of the U.S. from Cape Hatteras, NC, USA to Nova Scotia, Canada (Fig. 1). The region includes three primary subregions: the Mid-Atlantic Bight (MAB; 36°N-41°N, 70°W-76°W), Georges Bank (GB; 40°N-42°N, 66°W-70°W), and the Gulf of Maine (GoM; 42°N-45°N, 66°W-71°W). The shelf is commonly defined as inshore of the 200 m isobath, with deeper basins in the GoM exceeding 300 m depth. The NES is influenced by two major current systems: the warm, saline, northward flowing Gulf Stream, and the colder, fresher, southward flowing Labrador Current. Mesoscale features (e.g., eddies, fronts) and interannual variations in the path and flow of these two currents affect nutrient fluxes, productivity, and phytoplankton community composition across the region (Saba et al., 2015; Schollaert et al., 2004). The NES has experienced rapid warming in recent decades (Pershing et al., 2015), which has been connected to changes in phytoplankton bloom dynamics (Hunter-Cevera et al., 2016) and the distributions of fish and other marine species (Friedland et al., 2020; Kleisner et al., 2017). The NES is a well-studied region relative to other parts of the global ocean, with routine oceanographic surveys conducted throughout the region since the late 1970s (O’Reilly and Zetlin, 1998). At present, NOAA’s Northeast Fisheries Science
center Ecosystem Monitoring (EcoMon) program provides a range of hydrographic and biological data for the region (National Marine Fisheries Service, 2020).

2.2 In situ data

Discrete surface (upper 10 m) samples of HPLC pigments and the spectral absorption of phytoplankton ($a_{ph}(\lambda)$), CDOM ($a_g(\lambda)$), and NAP ($a_d(\lambda)$) collected throughout the NES region from 2003 to 2018 were acquired from NASA’s SeaWiFS Bio-optical Archive and Storage System (SeaBASS; https://seabass.gsfc.nasa.gov/) (Fig. 1). The compiled data were from several different cruises and experiments, with the largest portion from the Impacts of Climate Variability on Primary Production and Carbon Distributions in the Middle Atlantic Bight and Gulf of Maine (CliVEC) field campaign (Mannino et al., 2009) and the University of New Hampshire Western Gulf of Maine time series (Moore, 2006) (Table 2). Only HPLC pigment samples with a full set of seven diagnostic pigments required for estimating PSCs (fucoxanthin, peridinin, 19’-hexanoyloxyfucoxanthin, 19’-butanoyloxyfucoxanthin, alloxanthin, total chlorophyll-b, and zeaxanthin) were included in this analysis (see Section 2.6). For the absorption data, only measurements with 1-nm spectral resolution over the range 400-700 nm with coincident HPLC pigment measurements were retained. In order to calculate the combined absorption of CDOM and NAP ($a_{dg}(\lambda)$), a parameter routinely retrieved by satellite inversion algorithms, measurements of $a_g(\lambda)$ without matching $a_d(\lambda)$ (and vice-versa) were excluded. For stations with multiple samples in the upper 10 m, the data were averaged. To limit the effects of very shallow water and near-coastal processes, stations with a depth <25 m were removed prior to analysis.
In addition to the data obtained from SeaBASS, 40 HPLC, 24 $a_{ph}(\lambda)$, and 14 $a_{g}(\lambda)/a_{d}(\lambda)$ samples collected by our group on the Summer 2018 and Fall 2018 NOAA EcoMon surveys were also included. HPLC samples from these cruises were collected according to the NASA Ocean Optics Protocols (Mueller et al., 2003) and analyzed at Horn Point Laboratory (University of Maryland Center for Environmental Science). Absorption measurements were collected and analyzed following the procedures detailed in Mouw et al. (2017a).

**Fig.1.** Locations of the *in situ* data and satellite match-ups used in this study. The 200 and 2000 m isobaths from the 2019 General Bathymetric Chart of the Oceans (https://www.gebco.net/) are shown for reference. See Table 2 for information on data sources.
Table 2. Summary of in situ data sources. $N$ denotes the number of samples (after QA) and the number in parentheses refers to the number of satellite match-ups. Citations for the individual data sets from SeaBASS are also provided.

<table>
<thead>
<tr>
<th>Cruise/Experiment – P.I.(s)</th>
<th>Year(s)</th>
<th>Month(s)</th>
<th>$N$, HPLC</th>
<th>$N$, $a_{ph}(\lambda)$</th>
<th>$N$, $a_{adj}(\lambda)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Impacts of Climate Variability on Primary Production and Carbon Distributions in the Middle Atlantic Bight and Gulf of Maine (CiVEC) – Mannino et al., 2009</td>
<td>2009-2012</td>
<td>Feb, May, Jun, Aug, Nov</td>
<td>424 (212)</td>
<td>182 (101)</td>
<td>153 (86)</td>
</tr>
<tr>
<td>NOAA Ecosystem Monitoring (EcoMon) – Mannino et al., 2013</td>
<td>2013, 2018</td>
<td>Feb, Aug, Nov</td>
<td>71 (41)</td>
<td>24 (18)</td>
<td>14 (9)</td>
</tr>
<tr>
<td>Optical and Nutrient Dependence of Quantum Efficiency (OnDeque3) – Marra et al., 2008</td>
<td>2008</td>
<td>Jul</td>
<td>26 (15)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Tara Oceans Expedition – Boss et al., 2009</td>
<td>2012</td>
<td>Jan, Feb</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>East Coast Ocean Acidification (ECOA-1) – Mannino et al., 2015</td>
<td>2015</td>
<td>Jun, Jul</td>
<td>37 (16)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>LOBO timeseries – Roesler, 2009</td>
<td>2009</td>
<td>Mar</td>
<td>6</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Western Gulf of Maine – Moore, 2006</td>
<td>2006-2009</td>
<td>All months</td>
<td>188 (68)</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Ocean Color Cal Val (OCV) – Hooker et al., 2005</td>
<td>2007, 2009</td>
<td>May, Nov</td>
<td>16 (7)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>COASTAL (C7) – Hooker, 2000</td>
<td>2008</td>
<td>Oct</td>
<td>6</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>Delaware and Chesapeake Bay Fluorescence – Chekalyuk, 2008</td>
<td>2008</td>
<td>May</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2009oct_Chesapeake – Gould, 2009</td>
<td>2009</td>
<td>Oct</td>
<td>7 (3)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>BIOCOMPLEXITY – Harding, 2001</td>
<td>2003</td>
<td>Aug</td>
<td>2 (1)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Totals:</td>
<td></td>
<td></td>
<td>786 (368)</td>
<td>214 (123)</td>
<td>173 (99)</td>
</tr>
</tbody>
</table>
Fig. 2. Relative frequencies of *in situ* observations from the NES data set used in this study (blue stairs): (a) HPLC-measured \([Chl-a]\) \((N = 786)\), (b) \(a_{ph}(443)\) \((N = 214)\) and (c) \(a_{dg}(443)\) \((N = 173)\), with their respective monthly distributions (d-f, black bars). Global distributions from an OC-CCI v4.2 annual satellite composite for 2018 are overlaid for comparison (red line). Frequencies were normalized by the maximum value.

Quality assurance (QA) for the HPLC pigment data was carried out following Uitz et al. (2006). First, to account for differences in the detection limits and sensitivities of different HPLC processing methods, pigment concentrations <0.001 mg m\(^{-3}\) were set
to zero. Then, utilizing the relationship of Trees et al. (2000), who demonstrated that 
[Chl-a] and the total concentration of major accessory pigments ([TAcc]) co-vary in 
log-linear fashion within the euphotic zone of diverse oceanic regions, a robust linear 
regression (MATLAB function robustfit.m) of [TAcc] on [Chl-a] was performed to 
identify outliers. Any points exceeding three standard deviations with respect to the 
mean co-variation were excluded. QA for the \(a_{ph}(\lambda)\), \(a_g(\lambda)\), and \(a_d(\lambda)\) data consisted of 
the following steps: (1) overly noisy spectra were manually identified and removed; (2) 
spectra with negative values not exceeding -0.1 were offset by the most negative 
value (Grunert et al., 2019); (3) spectra were smoothed using a Savitsky-Golay 
filtering technique (MATLAB function sgolayfilt.m) with a 9-nm smoothing window 
(Torrecilla et al., 2011). The \(a_g(\lambda)\) and \(a_d(\lambda)\) values were then summed to obtain \(a_{dg}(\lambda)\). 
After QA, a total of 786 HPLC, 214 \(a_{ph}(\lambda)\), and 173 \(a_{dg}(\lambda)\) measurements remained. 
The pigment data were representative of every month of the year, with a slight bias 
towards summer months, whereas \(a_{ph}(\lambda)\) and \(a_{dg}(\lambda)\) data were almost exclusively 
limited to summer and fall months (Fig. 2).

2.3 Satellite data

Daily, Level-3 mapped (4-km resolution, sinusoidal projected) estimates of \(R_{rs}(\lambda)\), 
[Chl-a], \(a_{ph}(\lambda)\), and \(a_{dg}(\lambda)\) from the most recent version of the European Space 
Agency’s (ESA) Ocean Colour – Climate Change Initiative product (OC-CCI v4.2; 
Sathyendranath et al., 2019) at the time of the initiation of this study were downloaded 
from https://esa-oceancolour-cci.org/. OC-CCI v4.2 consists of merged, bias- 
corrected data from the Sea-viewing Wide-Field-of-View Sensor (SeaWiFS), Medium 
Resolution Imaging Spectrometer (MERIS), Moderate Resolution Imaging
Spectroradiometer (MODIS-Aqua), and Visible Infrared Imaging Radiometer Suite (VIIRS-SNPP) satellite sensors over the period 1997-2019. The multi-sensor data are band-shifted to SeaWiFS reference wavelengths (412, 443, 490, 510, 555, and 670 nm) and include per-pixel uncertainty estimates. OC-CCI v4.2 also incorporates the latest NASA re-processing (R2018), which corrected for significant drift in the MODIS-Aqua sensor. The reader is referred to the OC-CCI v4.2 Product User Guide (https://esa-oceancolour-cci.org/documents-list) for a more detailed overview.

The standard OC-CCI [Chl-a] algorithm uses a blended combination of NASA’s OC3, OCI, and OC5 algorithms (Gohin et al., 2002; Hu et al., 2012; O’Reilly et al., 1998) based on optical water classes (Jackson et al., 2017; Moore et al., 2009), which improves performance in optically complex waters. In addition to the standard algorithm, we also calculated [Chl-a] using the regional algorithm of Pan et al. (2010). This regional empirical algorithm was developed using coincident in situ measurements of HPLC pigments and $R_{rs}(\lambda)$ collected at various locations across the MAB and GoM and was applied using the published coefficients for SeaWiFS wavebands. The standard OC-CCI $a_{ph}(\lambda)$ and $a_{dg}(\lambda)$ products, derived using the Quasi-Analytical Algorithm (QAA_v5) of Lee et al. (2009), were used in this study.

For validation of the satellite input products ([Chl-a], $a_{ph}(\lambda)$, and $a_{dg}(\lambda)$) and PSC algorithm output, in situ samples were matched in time and space with the satellite data. Following convention, match-ups were determined as the geometric median of a 3x3 pixel box centered on the sampling location (nearest latitude and longitude), and only match-ups with at least 5 valid pixels and a median coefficient of variation of <0.15 for $R_{rs}(\lambda)$ bands between 412 and 555 nm were used to ensure spatial
homogeneity (Bailey and Werdell, 2006). As OC-CCI is a daily, multi-sensor product, a same-day coincidence window was used rather than the more stringent ±3-hour window recommended for a single mission by Bailey and Werdell (2006). This resulted in 368 [Chl-a], 123 \(a_{ph}(\lambda)\), and 99 \(a_{dg}(\lambda)\) match-ups (Table 2). Furthermore, all in situ pigment samples were matched with daily estimates of SST from the Multi-scale Ultra-high Resolution SST analysis (MUR v4.1) (Chin et al., 2017). MUR is a gap-filled, 1-km resolution gridded global SST product that combines night-time infrared SST retrievals from the MODIS sensor with multiple other in-orbit infrared/microwave instruments and data from NOAA’s in situ SST Quality Monitor (iQuam) database (Xu and Ignatov, 2014). Data for the period June 1, 2002 - December 31, 2019 were downloaded freely from NASA’s Physical Oceanography Distributed Active Archive Center (PO.DAAC; https://podaac.jpl.nasa.gov).

2.4 Data partitioning for model re-parameterization and validation

To allow for both the re-parameterization of abundance-based PSC models (see Sections 2.7.1.6 and 2.7.1.7) and independent model validation, the pigment data were split into two separate data sets. Of the 786 total samples, the 368 samples with a valid satellite [Chl-a] match-up (47% of the data) were removed and reserved for independent validation and are referred to as the validation data set. The remainder of the data \((N = 418)\) were used for model re-parameterization and are referred to as the parameterization data set.

2.5 Statistical metrics

We used several statistical metrics to compare algorithm estimates with the in situ data and evaluate algorithm performance. To measure accuracy, we used the mean
absolute difference \((MAD)\). While many studies commonly consider the root mean square difference \((RMSD)\), \(MAD\) has been recommended as a more unambiguous and appropriate metric for model assessment (Seegers et al., 2018; Willmott and Matsuura, 2005). To measure systematic bias, we used the mean bias \((\delta)\). The \(MAD\) and \(\delta\) are calculated according to:

\[
MAD = \frac{1}{N} \sum_{i=1}^{N} |M_i - O_i| \tag{1}
\]

and

\[
\delta = \frac{1}{N} \sum_{i=1}^{N} (M_i - O_i), \tag{2}
\]

where \(M\), \(O\), and \(N\) represent the modeled value (i.e., satellite estimate), the observed value (i.e., \textit{in situ}), and the number of samples, respectively. A positive (negative) \(\delta\) indicates a model’s tendency to systematically overestimate (underestimate) the variable of interest. We also computed the Pearson correlation coefficient \((r)\) and slope of a Type-II linear regression \((S)\) for additional comparison between modeled and \textit{in situ} values (Brewin et al., 2015b; Werdell et al., 2013). Type-II regression (MATLAB function \textsc{lsqfitgm.m}) was applied rather than Type-I regression as it accounts for the inherent measurement uncertainties of \textit{in situ} field data (Laws and Archie, 1981). While values of \(r\) and \(S\) that are close to one generally indicate better agreement between model estimates and \textit{in situ} observations, \(r\) and \(S\) alone provide no information on the accuracy or bias of a given model, and thus are not viewed in isolation from the \(MAD\) and \(\delta\) when assessing model performance. Statistical calculations involving total or size-specific \([Chl-a]\), \(a_{\text{ph}}(\lambda)\), or \(a_{\text{dg}}(\lambda)\) were performed in \(\log_{10}\) space, while calculations involving size fractions were performed in linear space.
2.6 Estimation of PSCs from HPLC pigments

For algorithm re-parameterization and validation, PSCs were estimated from the pigment data using the Diagnostic Pigment Analysis (DPA) method (Brewin et al., 2015a; Claustre, 2005; Devred et al., 2011; Uitz et al., 2006; Vidussi et al., 2001). This method has been used extensively for PSC algorithm development and validation given the relative abundance of HPLC pigment measurements compared with other in situ PSC methods. The DPA approach involves first re-constructing the measured \([\text{Chl-a}]\) (denoted here as \(C_{\text{HPLC}}\)) from the weighted sum of seven biomarker pigments (denoted \(C_{\text{DP}}\)) according to:

\[
C_{\text{DP}} = \sum_{i=1}^{7} W_i P_i, \tag{3}
\]

where \([W]\) represents pigment-specific weighting coefficients and \([P]\) is the set of seven biomarker pigments: \{fucoxanthin ([Fuco]), peridinin ([Perid]), 19'-hexanoyloxyfucoxanthin ([Hex-fuco]), 19'-butanoyloxyfucoxanthin ([But-fuco]), alloxanthin ([Allo]), total chlorophyll-b ([TChl-b]), zeaxanthin ([Zea])]\}. To derive an optimal set of weighting coefficients from the NES pigment data set, a multi-linear regression of \([P]\) on \(C_{\text{HPLC}}\) was performed. The newly computed weighting coefficients compared reasonably with those obtained from previous studies (Table 3), with the exception of large differences observed for the weights attributed to \([\text{But-fuco}]\) and \([\text{Allo}]\), which may be due to differences in community composition in the NES compared with the global ocean. The new weights yielded close agreement between
Fig. 3. Comparison of the HPLC-measured \([Chl\text{-}a] (C_{\text{HPLC}})\) and the reconstructed \([Chl\text{-}a] (C_{\text{DP}})\) from the sum of seven diagnostic pigments \((C_{\text{DP}})\) using the weighting coefficients derived from this study (green circles; \(MAD = 0.12, r = 0.98\)), the weights derived by Uitz et al. (2006) from a global data set (blue triangles; \(MAD = 0.47, r = 0.96\)), and no weighting coefficients (magenta squares; \(MAD = 0.62, r = 0.96\)).

\(C_{\text{DP}}\) and \(C_{\text{HPLC}}\) \((MAD = 0.12, r = 0.98)\), and demonstrated better results relative to using the unweighted sum of the diagnostic pigments (Vidussi et al., 2001), or the commonly applied weights of Uitz et al. (2006), which were derived from a large global pigment database (Fig. 3, Table 3).

The fractional contributions of micro-, nano-, and picoplankton were estimated from the ratios of the diagnostic pigments attributed to each size class to \(C_{\text{DP}}\). Two
Table 3. Diagnostic pigments $[P]$ and their associated taxonomic groups and attributed size classes, along with weighting coefficients $[W]$ obtained from this study and previous studies. The number of data points and geographical regions of each study are also provided.

<table>
<thead>
<tr>
<th></th>
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<th></th>
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<th></th>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>$P_1$ Fucoxanthin</td>
<td>Diatoms</td>
<td>Micro/ nano</td>
<td>2.20</td>
<td>1.41</td>
<td>1.51</td>
<td>1.65</td>
<td></td>
</tr>
<tr>
<td>$P_2$ Peridinin</td>
<td>Dinoflagellates</td>
<td>Micro</td>
<td>1.08</td>
<td>1.41</td>
<td>1.35</td>
<td>1.04</td>
<td></td>
</tr>
<tr>
<td>$P_3$ 19'-Hex</td>
<td>Prymnesiophytes</td>
<td>Nano</td>
<td>0.86</td>
<td>1.27</td>
<td>0.95</td>
<td>0.78</td>
<td></td>
</tr>
<tr>
<td>$P_4$ 19'-But</td>
<td>Pelagophytes</td>
<td>Nano</td>
<td>3.63</td>
<td>0.35</td>
<td>0.85</td>
<td>1.19</td>
<td></td>
</tr>
<tr>
<td>$P_5$ Alloxanthin</td>
<td>Cryptophytes</td>
<td>Nano</td>
<td>-0.10</td>
<td>0.6</td>
<td>2.71</td>
<td>3.14</td>
<td></td>
</tr>
<tr>
<td>$P_6$ Total chlorophyll-b</td>
<td>Prochlorophytes, Chlorophytes</td>
<td>Pico</td>
<td>1.21</td>
<td>1.01</td>
<td>1.27</td>
<td>1.38</td>
<td></td>
</tr>
<tr>
<td>$P_7$ Zeaxanthin</td>
<td>Prochlorophytes, Cyanobacteria</td>
<td>Pico</td>
<td>0.99</td>
<td>0.86</td>
<td>0.93</td>
<td>1.02</td>
<td></td>
</tr>
</tbody>
</table>

Number of data points | 786 | 2419 | 5841 | 2239 |

Geographic region | NES | Global | Global | North Atlantic |

diagnostic pigments were attributed to microplankton: [Fuco] and [Perid], associated with diatoms and dinoflagellates, respectively. Acknowledging that [Fuco] is also present in prymnesiophytes and chrysophytes, and that diatoms can also occupy the nano size range, Devred et al. (2011) introduced a modification that attributes a portion of [Fuco] ($P_1$) to nanoplankton, such that $P_1 = P_{1,nano} + P_{1,micro}$. In their approach, $P_{1,nano}$ is estimated from the equation:

$$P_{1,nano} = 10^{[q_1 \log_{10}(P_3) + q_2 \log_{10}(P_4)]},$$

(4)

where $P_3$ and $P_4$ are the pigments [Hex-fuco] and [But-fuco], respectively, and $q_1$ and $q_2$ are the coefficients of a 1% multi-linear quantile regression of $P_1$ on $P_3$ and $P_4$. 

20
Following the same approach, the coefficients $q_1$ and $q_2$ were re-computed for the NES pigment dataset, obtaining values of $q_1 = 0.999$ and $q_2 = 0.271$, and $P_{1,nano}$ was estimated using Eq. (4). In any instance where the estimated $P_{1,nano}$ was found to be greater than $P_1$, it was set equal to $P_1$. The fraction of microplankton ($F_{micro}$) was then calculated according to:

$$F_{micro} = \frac{\sum_{i=1}^{3} W_i P_i - W_i P_{1,nano}}{C_{DP}}.$$  (5)

Three diagnostic pigments were used to estimate nanoplanckton: [Hex-fuco], [But-fuco], and [Allo], attributed to prymnesiophytes, pelagophytes, and cryptophytes, respectively (Brewin et al., 2015a; Roy, 2011; Uitz et al., 2006). Brewin et al. (2010) proposed a linear adjustment that attributes a portion of [Hex-fuco] to picoeukaryotes (picoplankton) in ultra-oligotrophic environments ([Chl-a] < 0.08 mg m$^{-3}$). However, considering that only one sample in the data set used in this study met this criterion ([Chl-a] = 0.07 mg m$^{-3}$) and the adjustment was found to make only minor difference (not shown), it was excluded for simplicity. Incorporating the [Fuco] modification of Devred et al. (2011), the fraction of nanoplanckton was calculated as:

$$F_{nano} = \frac{\sum_{i=2}^{5} W_i P_i + W_i P_{1,nano}}{C_{DP}}.$$  (6)

The final two diagnostic pigments: [$TChl-b$] and [Zea], were attributed to the picoplankton class, the former associated with prochlorophytes and chlorophytes and the latter with prochlorophytes and cyanobacteria (Chisholm et al., 1988; Roy, 2011; Uitz et al., 2006). The fraction of picoplankton ($F_{pico}$) was computed as:

$$F_{pico} = \frac{\sum_{i=6}^{7} W_i P_i}{C_{DP}}.$$  (7)
Once \( F_{\text{micro}} \), \( F_{\text{nano}} \), and \( F_{\text{pico}} \) (collectively referred to as \( F_{\text{size}} \)) were determined, the [Chl-a] specific to each size class (collectively referred to as \( C_{\text{size}} \)) was calculated by multiplying \( F_{\text{size}} \) by \( C_{\text{HPLC}} \), such that:

\[
C_{\text{micro}} = F_{\text{micro}} C_{\text{HPLC}}, \tag{8a}
\]

\[
C_{\text{nano}} = F_{\text{nano}} C_{\text{HPLC}}, \tag{8b}
\]

\[
C_{\text{pico}} = F_{\text{pico}} C_{\text{HPLC}}. \tag{8c}
\]

2.7 PSC algorithms

A variety of PSC algorithms, including purely abundance-based methods, abundance-based methods that include SST, and absorption-based approaches, were selected for optimization and/or evaluation in this study. The abundance-based models chosen are among the most commonly applied in the literature and have been successfully re-parameterized for studies in diverse ocean regions, including continental shelf systems (Brito et al., 2015; Sun et al., 2018). The absorption-based models were chosen based on their global performance metrics (Mouw et al., 2017b) and their demonstrated consistency in capturing multiple phytoplankton phenology metrics in the North Atlantic (Kostadinov et al., 2017). The following sections provide brief overviews of each model, including their principal frameworks, methods used for model development/parameterization, and key differences. For more comprehensive information, the reader is referred to the original publications and the reviews of Mouw et al. (2017b) and IOCCG (2014).

2.7.1 Abundance-based

2.7.1.1 Brewin et al. (2010, 2015a)
The three-component model of Brewin et al. (2010) relates the fractional contribution of combined pico- and nanoplankton ($F_{pico,nano}$) and picoplankton ($F_{pico}$) to [Chl-a] using two exponential functions (Sathyendranath et al., 2001) according to:

\[
F_{pico,nano} = \frac{c_{m}^{nano,nano} \left(1 - \exp\left(-\frac{D_{pico,nano}[Chl-a]}{c_{m}^{nano,nano}}\right)\right)}{[Chl-a]}, \tag{9}
\]

\[
F_{pico} = \frac{c_{m}^{pico} \left(1 - \exp\left(-\frac{D_{pico}[Chl-a]}{c_{m}^{pico}}\right)\right)}{[Chl-a]}, \tag{10}
\]

where the model parameters $c_{m}^{nano,nano}$ and $c_{m}^{pico}$ represent asymptotic maximum [Chl-a] for the associated size classes, and $D_{pico,nano}$ and $D_{pico}$ represent the fraction of each size class as [Chl-a] tends toward zero. The model parameters are derived by fitting Eqs. 9 and 10 to an in situ data set of $F_{pico,nano}$, $F_{pico}$, and [Chl-a] via nonlinear least squares regression. $F_{micro}$ and $F_{nano}$ are then determined as $F_{micro} = 1 - F_{pico,nano}$ and $F_{nano} = F_{pico,nano} - F_{pico}$, respectively. Brewin et al. (2010) used a data set of HPLC pigments from the Atlantic Ocean ($N = 1935$) (Atlantic Meridional Transect cruises 5-15) and estimated PSCs using DPA to derive model parameters. Brewin et al. (2015a) utilized a much larger global data set of HPLC measurements ($N = 5841$) to compute the model parameters. These two models are denoted B10 and B15 throughout the remainder of the text. For simplicity, the notation B10 is also used to refer to the general framework that underlies both of these models (i.e., Eqs. 9 and 10), in addition to the parameterization specific to that study. Further, while Brewin et al. (2015a) also investigated the influence of average irradiance in the mixed layer on model parameters, in this study B15 refers to the model without this modification. Parameter values obtained from various studies are provided in Table 4.

2.7.1.2 Brewin et al. (2017)
Brewin et al. (2017) used a merged *in situ* HPLC/SFF data set from the North Atlantic \((N = 2239)\) to compute the parameters of the B10 model (Eqs. 9 and 10). They then modified the model parameters to vary as a function of SST by matching the *in situ* PSC data with satellite-derived SST and conducting a running fit of the model to the data binned by increasing SST. They represented the resulting relationships between SST and model parameters using logistic functions, such that \(C_{m \text{pico}, \text{nano}}, C_{m \text{pico}}, D_{\text{pico}, \text{nano}}\) and \(D_{\text{pico}}\) are expressed as:

\[
C_{\text{pico}, \text{nano}}^m = 1 - \left\{ \frac{G_1}{1+\exp \left( -G_2(SST-G_3) \right) } + G_4 \right\}, \tag{11}
\]

\[
C_{\text{pico}}^m = 1 - \left\{ \frac{H_1}{1+\exp \left( -H_2(SST-H_3) \right) } + H_4 \right\}, \tag{12}
\]

\[
D_{\text{pico}, \text{nano}} = \frac{J_1}{1+\exp \left( -J_2(SST-J_3) \right) } + J_4, \tag{13}
\]

\[
D_{\text{pico}} = \frac{K_1}{1+\exp \left( -K_2(SST-K_3) \right) } + K_4, \tag{14}
\]

where \(G_i, H_i, J_i,\) and \(K_i\) (where \(i = 1-4\)) are empirical parameters controlling the shape of the respective logistic curve and are provided in Table 4 of Brewin et al. (2017). In the remainder of this text, the notation B17 is used to refer to the SST-independent parameterization of the model, which uses a single set of model parameters derived from their full data set (Table 4). The SST-dependent parameterization, which uses Equations 11-14 with the published coefficients, is denoted as B17-SST.

2.7.1.3 Devred et al. (2011)

The model of Devred et al. (2011) (denoted D11) is based on the same exponential functions as the B10 model (Eqs. 9 and 10). The primary difference is that the model parameters \(C_{m \text{pico}, \text{nano}}^m\) and \(C_{m \text{pico}}^m\) were not derived from HPLC pigment-based size classes, but rather by successive application of the two-population absorption model
of Devred et al. (2006) to \( a_{ph}(\lambda) \) and \([Chl-a]\) data from the Northwest Atlantic and NASA’s NOMAD data set. While the model was originally applied as a spectral-based approach, in this study it is implemented as an abundance-based method, using the parameters provided in Table 2 of Brewin et al. (2015a). In a comparison of nine existing PSC algorithms in the Northwest Atlantic region near Newfoundland, Liu et al. (2018) found the application of this model as an abundance-based method to be the most successful (Model E in their study). The reader is referred to Chapter 4 of IOCCG (2014) for more information on this approach.

2.7.1.4 Hirata et al. (2011)

The empirical model of Hirata et al. (2011) (denoted H11) estimates the fractional contribution of pico- and microplankton to \([Chl-a]\) according to:

\[
F_{\text{pico}} = -[a_1 + \exp(a_2 X + a_3)]^{-1} + a_4 X + a_5, \tag{15}
\]

\[
F_{\text{micro}} = [b_1 + \exp(b_2 X + b_3)]^{-1}, \tag{16}
\]

where \(a_i\) and \(b_i\) (where \(i = 1-5\) and \(i = 1-3\), respectively) are empirical coefficients specific to each size class and \(X\) is log10-transformed \([Chl-a]\). \(F_{\text{nano}}\) is then calculated by difference \((1 - F_{\text{micro}} - F_{\text{pico}})\). The H11 model was developed using PSCs derived from a global HPLC data set \((N = 2776)\) following a unique version of DPA that attributes \([TChl-b]\) to nanoplankton rather than picoplankton, as in Brewin et al. (2010, 2015a, 2017) and the present study (see Section 2.6).

2.7.1.5 Moore and Brown (2020)

The model of Moore and Brown (2020) utilizes the H11 microplankton model (Eq. 16) to estimate both \(F_{\text{pico}}\) and \(F_{\text{micro}}\). Using a data set of surface HPLC pigments from the Atlantic Ocean \((N = 1083)\), they developed two separate models: one
parameterized using the DPA method (following the procedure of Brewin et al. (2015a)) and one parameterized using CHEMTAX (Mackey et al., 1996). They then incorporated different remotely sensed environmental variables into the models, following a similar approach to that of Brewin et al. (2015a, 2017), and created a look-up table (LUT) for each model parameter indexed by the environmental data. Of the environmental variables tested, they found that the inclusion of SST resulted in the largest reduction of model error. In this study, the DPA version of the model was applied, both with and without the incorporation of SST. The SST-independent model (denoted MB20) was applied using Eq. (16) with the coefficients provided in Table 4 of Moore and Brown (2020). The SST-dependent model (denoted MB20-SST) was applied using the set of parameters from their LUT indexed by SST (obtained from Timothy Moore via personal communication).

2.7.1.6 Re-parameterized B10 and H11 models

We derived new model parameters for the B10 and H11 models using the pigment-based estimates of $F_{size}$ from the NES parameterization data set (Table 3). To re-parameterize the B10 model, Eqs. (9) and (10) were fit to $F_{pico,nano}$, $F_{pico}$, and $C_{HPLC}$ using a standard nonlinear least squares curve fitting method (MATLAB function lsqcurvefit.m, Levenberg-Marquardt algorithm) (Brewin et al., 2017; Brewin et al., 2015a). We followed the same procedure for the H11 model, however with a slight modification. Considering the findings of Moore and Brown (2020), who showed better results when using the H11 microplankton formula (Eq. 16) for estimating both $F_{micro}$ and $F_{pico}$, we tested the fit of both Eq. (15) and Eq. (16) to our data set of $F_{pico}$ and $C_{HPLC}$. We likewise found improved fit statistics when using Eq. (16) to estimate $F_{pico}$
Therefore, we fit Eq. (16) to both $F_{\text{micro}}$ and $F_{\text{pico}}$ and $C_{\text{HPLC}}$ to derive new model parameters. We refer to these two regionally re-parameterized abundance-based models as B-NES and H-NES, respectively.

**Table 4.** Parameter values for the abundance-based models of Brewin et al. (2010) (Eqs. 9 and 10) and Hirata et al. (2011) (Eq. 16), obtained from this study and from previous studies.

<table>
<thead>
<tr>
<th>Study</th>
<th>Geographic region</th>
<th>Years</th>
<th>Parameters for Equations (9) and (10)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>$C^m_{\text{pico,nano}}$</td>
</tr>
<tr>
<td>Brewin et al. (2010)</td>
<td>Atlantic</td>
<td>1997-2004</td>
<td>-</td>
</tr>
<tr>
<td>Brewin et al. (2015)</td>
<td>Global</td>
<td>1992-2012</td>
<td>-</td>
</tr>
<tr>
<td>Brewin et al. (2017)$^a$</td>
<td>N Atlantic</td>
<td>1995-2015</td>
<td>-</td>
</tr>
<tr>
<td>Devred et al. (2011)</td>
<td>NW Atlantic</td>
<td>1996-2003</td>
<td>-</td>
</tr>
<tr>
<td>This Study</td>
<td>NES</td>
<td>2003-2018</td>
<td>-</td>
</tr>
</tbody>
</table>

<table>
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<tr>
<th>Parameters for Equation (16)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$b_{1,\text{micro}}$</td>
</tr>
<tr>
<td>Hirata et al. (2011)$^b$</td>
</tr>
<tr>
<td>Moore and Brown (2020)$^c$</td>
</tr>
<tr>
<td>This Study</td>
</tr>
</tbody>
</table>

$^a$ Refers to the SST-independent model from their study  
$^b$ Hirata et al. (2011) used a different empirical formula to estimate picoplankton (Eq. 15; see their Table 2)  
$^c$ Refers to coefficients derived using the DPA method [see Table 4 of Moore and Brown (2020)]

### 2.7.1.7 Regional SST-modified B10 model

Following a similar methodology to recent studies (Brewin et al., 2019, 2017; Moore and Brown, 2020; Sun et al., 2019), we investigated the influence of SST on
Fig. 4. Brewin et al. (2010) model parameters (a) $C_{\text{size}}^m$ as a function of SST. Circle markers show the median parameter values obtained by performing a running bootstrap fit of the model (Eqs. 9 and 10) to the in situ parameterization data set ($N = 418$) sorted by increasing SST. The SST-parameter relationships were smoothed using a 5-point running average. Dashed lines indicate the SST-independent model (B-NES) parameters obtained when fitting the model to the full parameterization data set.
the parameters of the B10 model. This was done by sorting the pigment-based estimates of $F_{\text{pico}}$ and $F_{\text{pico,nano}}$ from the parameterization data set by increasing SST and conducting a running fit of the model from low to high SST, using a bin size of 125 samples. Starting at the lowest temperature, the bin was moved at one-sample intervals, and at each interval Eqs. (9) and (10) were fit to the data within the bin using the same the same approach as described in the previous section. From each fit, the median values of the model parameters $C_{\text{m size}}$ and $D_{\text{size}}$, along with the average SST of the binned data, were incorporated into a LUT. Finally, the LUT relationships were smoothed using a 5-point running mean (Fig. 4). We chose a LUT approach as opposed to fitting logistic functions to represent the SST-parameter relationships in order to capture variability in the relationships that may be ecological meaningful and would otherwise be smoothed out by a logistic curve. The LUT included 293 unique sets of model parameters covering a range of SST of 6.8 – 21.4 °C at intervals of approximately 0.06 °C. Application of the LUT enables a dynamic set of model parameters to be derived based on remotely sensed SST. We refer to this SST-dependent re-parameterization as B-NES-SST henceforth.

2.7.2 Absorption-based

2.7.2.1 Ciotti et al. (2002)

The model of Ciotti et al. (2002) (denoted C02) estimates the fractional contribution of picoplankton ($F_{\text{pico}}$), by weighting $a_{\text{ph}}(\lambda)$ between two basis spectra representing "pure" micro- and picoplankton. The basis spectra were determined by lab measurements of $a_{\text{ph}}(\lambda)$ of 16 natural phytoplankton communities of varying dominant
cell sizes, and are provided in Ciotti et al. (2002) with an updated picoplankton spectra provided by Ciotti and Bricaud (2006). The model can be expressed as:

$$a_{ph}^*(\lambda) = [F_{pico} * \tilde{a}_{pico}^*(\lambda)] + [(1 - F_{pico}) * \tilde{a}_{micro}^*(\lambda)],$$  

(13)

where $\tilde{a}_{pico}^*(\lambda)$ and $\tilde{a}_{micro}^*(\lambda)$ represent the basis spectra of pico- and microplankton, respectively. $F_{pico}$ was estimated from Eq. (13) by performing a linear least squares optimization (MATLAB function lsqlin.m), using satellite-derived $a_{ph}^*(\lambda)$ (calculated by dividing the satellite $a_{ph}(\lambda)$ by the satellite [Chl-a]) and the published basis spectra at four wavelengths (443, 490, 510, and 555 nm). These four wavelengths were used due to the poor retrieval of $a_{ph}(670)$ (see Section 3.1) and better statistical performance when excluding 412 nm. The inverse of $F_{pico}$ ($1 - F_{pico}$) was considered equivalent to the combined fraction of micro- and nanoplankton ($F_{micro,nano}$).

2.7.2.2 Mouw and Yoder (2010)

The algorithm of Mouw and Yoder (2010) (denoted MY10) employs an optical LUT containing ranges of $F_{micro}$ (binned to increments of 0.1), [Chl-a], and $a_{dg}(443)$, from which $R_{rs}(\lambda)$ was calculated using the radiative transfer software HydroLight (Mobley and Sundman, 2013). The model uses satellite [Chl-a] and $a_{dg}(443)$ as inputs to first narrow the search space of the LUT, then the closest matching LUT $R_{rs}(443)$ to the satellite-derived $R_{rs}(443)$ is found to retrieve the corresponding $F_{micro}$. $F_{pico,nano}$ is then calculated as $1 - F_{micro}$. Based on determined thresholds for the detectability of cell size from SeaWiFS $R_{rs}(\lambda)$ (Mouw and Yoder, 2010), the algorithm masks pixels with [Chl-a] < 0.05 mg m$^{-3}$, [Chl-a] > 1.75 mg m$^{-3}$, or $a_{dg}(443)$ > 0.17. Additionally, when applied to a satellite image, a 2D-averaging filter (MATLAB function filter2.m) with a 3x3 pixel size is applied in the algorithm routine.
PSC algorithm performance was assessed statistically (see Section 2.5) as follows. First, using the *in situ* \([\text{Chl-a}]\) data as input, estimates of \(F_{\text{size}}\) and \(C_{\text{size}}\) from the SST-independent abundance-based models (B10, B15, B17, D11, H11, MB20, B-NES, and H-NES) were compared with the *in situ* pigment-based estimates of \(F_{\text{size}}\) and \(C_{\text{size}}\) for both the parameterization and validation data sets. Then, using the *in situ* \([\text{Chl-a}]\) and matching satellite SST as input, estimates of \(F_{\text{size}}\) and \(C_{\text{size}}\) from the SST-dependent models (B17-SST, MB20-SST, B-NES-SST) were also compared with the *in situ* \(F_{\text{size}}\) and \(C_{\text{size}}\) from both the parameterization and validation data sets, and the influence of SST on model performance relative to the SST-independent models was quantified. Lastly, using the satellite products as input, independent satellite validation of \(C_{\text{size}}\) was conducted for the regionally re-parameterized abundance-based models (B-NES-SST, B-NES, and H-NES) and the absorption-based algorithms (C02 and MY10).

To illustrate the application of the models to ocean color imagery and explore spatial-seasonal variations of PSCs in the NES region, the best performing abundance-based and absorption-based models were applied to monthly imagery composites for April 2019 and September 2019, and qualitatively compared.

### 3. Results

#### 3.1 Satellite validation of \([\text{Chl-a}], a_{\text{ph}}(\lambda)\) and \(a_{\text{dg}}(\lambda)\)

As the performance of PSC algorithms is largely dependent on the quality of the satellite products used as inputs, the satellite retrievals of \([\text{Chl-a}], a_{\text{ph}}(\lambda)\) and \(a_{\text{dg}}(\lambda)\) in the NES were first validated with the available *in situ* observations. Of the two \([\text{Chl-a}]\)
algorithms assessed, the standard OC-CCI algorithm (Fig. 5a) displayed lower
difference and bias \((\text{MAD} = 0.21, \delta = -0.03)\) than the regional empirical algorithm of
Pan et al. (2010) \((\text{MAD} = 0.27, \delta = -0.20; \text{Fig. 5b})\) when compared with the \textit{in situ}
HPLC \([\text{Chl-a}]\). The OC-CCI algorithm was associated with overestimation at low \([\text{Chl-a}]\) and underestimation at high \([\text{Chl-a}]\) (>0.6 mg m\(^{-3}\)). This is a relatively common
feature of \([\text{Chl-a}]\) estimated from empirical band-ratio algorithms (e.g., OCx), that can
be attributed primarily to the impact of phytoplankton cell size and underlying
variability in the concentrations of CDOM and inorganic particulates (Dierssen, 2010;
Mouw et al., 2012; Sauer et al., 2012). In contrast, the algorithm of Pan et al. (2010)
exhibited a systematic underestimation across the entire \([\text{Chl-a}]\) range \((\delta = -0.20)\).
This difference is reflected in a lower Type-II regression slope for the OC-CCI
algorithm \((S = 0.66)\) compared with a slope closer to one for the Pan et al. (2010)
algorithm \((S = 0.89)\). The two algorithms displayed similar correlation coefficients \((r =
0.75\) and 0.74 respectively). The performance of the OC-CCI algorithm was
comparable to the global \([\text{Chl-a}]\) validation of OC-CCI v4.2 as shown in the Product
User Guide \((\text{RMSD} = 0.32, \delta = 0.07, r = 0.75, S = 0.72; \text{see their Fig. 5})\). Considering
the lower difference and bias of the OC-CCI product, \([\text{Chl-a}]\) from this algorithm was
used as the satellite input to the PSC models in this study.
Fig. 5. Bivariate histograms showing the satellite-to-in situ comparisons of [Chl-a], $a_{ph}(\lambda)$, and $a_{dg}(\lambda)$, shaded by number of observations: (a) [Chl-a] from the standard OC-CCI algorithm, (b) [Chl-a] from the regional algorithm of Pan et al. (2010), (c) $a_{ph}(443)$, (d) $a_{ph}(490)$, (e) $a_{ph}(510)$, (f) $a_{dg}(443)$, (g) $a_{dg}(490)$, and (h) $a_{dg}(510)$ from the standard OC-CCI algorithm (QAA_v5). The solid black line is the 1:1 line, dashed
black lines indicate the 1:1 line ± 30%, and the red line is the Type-II regression line. $N$ denotes the number of match-ups, $MAD$ denotes the mean absolute difference, $\delta$ denotes the bias, $r$ denotes the correlation coefficient, and $S$ denotes the regression slope. The $a_{ph}(\lambda)$ and $a_{dg}(\lambda)$ data are shown using the same x- and y-axis range for comparison.

The standard OC-CCI estimates of $a_{ph}(\lambda)$ and $a_{dg}(\lambda)$, derived using QAA_v5, compared reasonably with in situ measurements, with most points falling within the ±30% uncertainty range at the wavelengths 443, 490, and 510 nm (Fig. 5 c-h). The lowest differences for $a_{ph}(\lambda)$ were observed at 490 and 510 nm ($MAD = 0.22$ and $0.24$, respectively), with retrievals of $a_{ph}(443)$ exhibiting slightly higher uncertainty ($MAD = 0.28$). For $a_{dg}(\lambda)$, the best performance was observed at 443 nm ($MAD = 0.17$, $\delta = -0.14$). There was a consistent negative bias in the retrieved $a_{dg}(\lambda)$ which became more negative at longer wavelengths, corresponding with an increasingly positive bias for $a_{ph}(\lambda)$. Retrievals of $a_{ph}(555)$ and $a_{ph}(670)$ were associated with larger errors ($MAE = 0.32$ and $0.62$, respectively), the same being true for $a_{dg}(555)$ and $a_{dg}(670)$ ($MAE = 0.38$ and $0.41$, respectively). The decrease in performance at longer wavelengths for QAA_v5 is consistent with results from the global inter-comparison of bio-optical algorithms conducted by Brewin et al. (2015b) (Model E in their study). Considering the reasonable performance of the standard OC-CCI $a_{ph}(\lambda)$ and $a_{dg}(443)$ products ($443$ nm is the only wavelength required for $a_{dg}(\lambda)$ as input into the MY10 algorithm), no regional optimization of the absorption retrievals was attempted in this study.

3.2 Comparison of SST-independent abundance-based models
Figure 6 shows the in situ pigment-based estimates of $F_{\text{size}}$ and $C_{\text{size}}$ from the parameterization data set ($N = 418$) with the SST-independent abundance-based models overlain. $F_{\text{size}}$ and $C_{\text{size}}$ exhibited trends with $C_{\text{HPLC}}$ that are consistent with established relationships of phytoplankton size structure and total biomass (Brewin et al., 2010; Hirata et al., 2011; Uitz et al., 2006). $F_{\text{micro}}$ generally increased, while $F_{\text{pico,nano}}$ and $F_{\text{pico}}$ generally decreased with increasing $C_{\text{HPLC}}$. $F_{\text{nano}}$ displayed a unimodal relationship $C_{\text{HPLC}}$, peaking at intermediate $C_{\text{HPLC}}$. $C_{\text{micro}}$ increased in near log-linear fashion with $C_{\text{HPLC}}$, becoming more tightly correlated at higher $C_{\text{HPLC}}$, when microplankton are the dominant size class. $C_{\text{pico,nano}}$, $C_{\text{nano}}$, and $C_{\text{pico}}$, also displayed an overall positive relationship with $C_{\text{HPLC}}$, with greater variability at higher $C_{\text{HPLC}}$. The abundance-based models all followed to first order these general trends, with some variations that can be attributed to differences in the model frameworks, data sets, and approaches used for model development/parameterization. For example, the B10 model parameters $C_{\text{m,pico,nano}}$ and $C_{\text{m,pico}}$ impose asymptotic maximum values for $C_{\text{pico,nano}}$ and $C_{\text{pico}}$ respectively, while the purely empirical H11 model does not define any strict maximums. This can be seen when comparing $C_{\text{pico}}$ predicted by the H-NES model, which increases continuously with $C_{\text{HPLC}}$, with that of the B-NES model, which levels off at the imposed maximum concentration ($C_{\text{m,pico}} = 0.2 \text{ mg m}^{-3}$) (Fig. 6h). The H11-modeled $F_{\text{pico}}$ is based on a different empirical function (Eq. 15) than the one used in this study (see Section 2.7.1.6), and goes to zero at $C_{\text{HPLC}} > 4 \text{ mg m}^{-3}$ (Fig. 6d), accounting for the breakdown in this model at higher $C_{\text{HPLC}}$ for $C_{\text{pico}}$, $C_{\text{nano}}$, and $C_{\text{pico,nano}}$ (Fig. 6f-h). Compared with the other models, the B-NES model predicted a slightly higher $F_{\text{micro}}$ and lower $F_{\text{pico,nano}}$ and $F_{\text{pico}}$ at low $C_{\text{HPLC}}$ (Fig. 6a,b,d).
Fig. 6. Pigment-based estimates of $F_{\text{size}}$ (a-d) and $C_{\text{size}}$ (e-h) as a function of $C_{\text{HPLC}}$ from the parameterization data set ($N = 418$) with abundance-based models overlaid: re-parameterized B10 model (B-NES, solid black), re-parameterized H11 model (H-NES, dashed black), Brewin et al., 2010 (B10, blue), Brewin et al., 2015a (B15, green), Brewin et al., 2017 (B17, red), Devred et al., 2011 (D11, violet), Hirata et al., 2011 (H11, brown), and Moore and Brown, 2020 (MB20, yellow).

Furthermore, the B-NES-modeled $F_{\text{nano}}$ leveled off at low $C_{\text{HPLC}}$ rather than decreasing as with the other models (Fig. 6c). Despite the variability between the different models, the range of variability in the pigment-based estimates of $F_{\text{size}}$ and $C_{\text{size}}$ was generally greater across the entire trophic domain. Statistical comparison between the in situ and modeled $F_{\text{size}}$ and $C_{\text{size}}$ for both the parameterization and validation data sets yielded very similar metrics between models (Appendix A). Overall, minimal...
improvement in performance was observed for the regionally re-parameterized models (B-NES and H-NES) compared with the other models examined, although there was a reduction in error and bias for the nanoplankton size class compared with the original global models (B10 and H11).

3.3 Regional SST-dependent model (B-NES-SST)

Figure 7 shows the modeled $F_{\text{size}}$ and $C_{\text{size}}$ from the B-NES-SST model as a function of [Chl-a] at varying SST along with the SST-independent (B-NES) model for reference. SST had a clear influence on the predicted $F_{\text{size}}$ and $C_{\text{size}}$ for all of the size classes. Lower SST was associated with a higher fraction of microplankton, and a lower fraction of nanoplankton and picoplankton (Fig. 7a-d), consistent with the results of previous studies (Brewin et al., 2017b; Moore and Brown, 2020; Sun et al., 2019b). This relationship was generally observed across the entire [Chl-a] domain but was more pronounced at lower [Chl-a]. $F_{\text{nano}}$ increased with SST for [Chl-a] > 1 mg m$^{-3}$, whereas for [Chl-a] < 1 mg m$^{-3}$, $F_{\text{nano}}$ increased with SST only up to ~18 ºC, beyond which there was a decrease in $F_{\text{nano}}$ corresponding with an increase in $F_{\text{pico}}$ at SST > 18 ºC. The largest variability in modeled $C_{\text{micro}}$ with SST was at [Chl-a] < 1 mg m$^{-3}$, whereas the largest variability in modeled $C_{\text{pico,nano}}$, $C_{\text{nano}}$, and $C_{\text{pico}}$ with SST was at [Chl-a] > 1 mg m$^{-3}$ (Fig. 7e-h).

Results from the statistical comparison of modeled $F_{\text{size}}$ and $C_{\text{size}}$ from the B-NES-SST, B17-SST, and MB20-SST models with the in situ pigment-based estimates of $F_{\text{size}}$ and $C_{\text{size}}$ are presented in Table 5. For both the parameterization and validation data sets, the B-NES-SST model performed with lower error and significantly higher correlation coefficients than the other two SST-dependent models across all size
There was also a consistent improvement in performance (i.e., reduction in error, increase in correlation coefficient) for the B-NES-SST model relative to the SST-independent B-NES model for both data sets. Considering the statistical results from the validation set, the inclusion of SST led to a reduction in MAD of 10-12% for $F_{\text{size}}$ and 4-10% for $C_{\text{size}}$, with the largest reductions for $F_{\text{nano}}$ and $C_{\text{pico,nano}}$, respectively. Likewise, the inclusion of SST increased the correlation coefficient ($r$) for $F_{\text{size}}$ and $C_{\text{size}}$, with the largest increases for $F_{\text{nano}}$ and $C_{\text{pico}}$, respectively. Interestingly, the B17-SST model exhibited slightly worse performance relative to the SST-independent B17 model.

**Fig. 7.** Modeled $F_{\text{size}}$ (a-d) and $C_{\text{size}}$ (e-h) from the B-NES-SST model plotted as a function of [Chl-a], with the color gradient illustrating the changes in the model when model parameters vary as a function of SST (see Section 2.7.1.7, Fig. 4). The black line indicates the SST-independent model, with a single set of model parameters (B-NES).
Table 5. Mean absolute difference (MAD) and correlation coefficients ($r$) for $F_{size}$ and $C_{size}$ (values for $C_{size}$ shown in parentheses) for the SST-dependent abundance-based models applied to the *in situ* parameterization and validation data sets. The percent change in the metrics when incorporating SST relative to the SST-independent models is included for reference. Percentages are rounded to the nearest 1%.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Model</th>
<th>Parameterization set (N = 418)</th>
<th>Validation set (N = 368)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>MAD % change with SST $r$ MAD % change with SST $r$</td>
<td>MAD % change with SST $r$ MAD % change with SST $r$</td>
</tr>
<tr>
<td>$F_{micro}$ ($C_{micro}$)</td>
<td>B-NES-SST</td>
<td>0.16 (0.21) -11 (-13) 0.68 (0.89) +21 (+3)</td>
<td>0.17 (0.24) -11 (-4) 0.58 (0.87) +32 (+2)</td>
</tr>
<tr>
<td></td>
<td>B17-SST</td>
<td>0.19 (0.23) +6 (-4) 0.54 (0.87) -4 (+1)</td>
<td>0.20 (0.25) +5 (0) 0.44 (0.85) 0 (0)</td>
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<tr>
<td></td>
<td>MB20-SST</td>
<td>0.18 (0.23) -5 (-4) 0.59 (0.83) +7 (-3)</td>
<td>0.19 (0.24) -5 (-4) 0.49 (0.86) +9 (+1)</td>
</tr>
<tr>
<td>$F_{pico,nano}$ ($C_{pico,nano}$)</td>
<td>B-NES-SST</td>
<td>0.16 (0.17) -11 (-11) 0.68 (0.75) +21 (+12)</td>
<td>0.17 (0.18) -11 (-10) 0.58 (0.77) +32 (+6)</td>
</tr>
<tr>
<td></td>
<td>B17-SST</td>
<td>0.19 (0.20) +5 (+5) 0.54 (0.59) -4 (-11)</td>
<td>0.20 (0.20) +5 (0) 0.44 (0.72) 0 (+1)</td>
</tr>
<tr>
<td></td>
<td>MB20-SST</td>
<td>0.18 (0.20) -5 (0) 0.59 (0.56) +7 (-16)</td>
<td>0.19 (0.19) -5 (-5) 0.49 (0.73) +9 (0)</td>
</tr>
<tr>
<td>$F_{nano}$ ($C_{nano}$)</td>
<td>B-NES-SST</td>
<td>0.13 (0.21) -13 (-13) 0.55 (0.74) +49 (+12)</td>
<td>0.15 (0.26) -12 (-7) 0.39 (0.79) +225 (+6)</td>
</tr>
<tr>
<td></td>
<td>B17-SST</td>
<td>0.14 (0.24) -7 (-4) 0.44 (0.66) +19 (0)</td>
<td>0.16 (0.27) -6 (-4) 0.29 (0.75) +45 (+1)</td>
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<td>MB20-SST</td>
<td>0.16 (0.26) -6 (-4) 0.28 (0.58) +8 (-13)</td>
<td>0.16 (0.27) -11 (-7) 0.18 (0.72) 0 (-3)</td>
</tr>
<tr>
<td>$F_{pico}$ ($C_{pico}$)</td>
<td>B-NES-SST</td>
<td>0.07 (0.19) -13 (-10) 0.70 (0.64) +15 (+16)</td>
<td>0.09 (0.20) -10 (-9) 0.73 (0.62) +16 (+15)</td>
</tr>
<tr>
<td></td>
<td>B17-SST</td>
<td>0.08 (0.23) 0 (-4) 0.63 (0.46) 0 (+7)</td>
<td>0.09 (0.23) -10 (-4) 0.73 (0.53) +4 (+23)</td>
</tr>
<tr>
<td></td>
<td>MB20-SST</td>
<td>0.08 (0.21) 0 (-13) 0.67 (0.53) +6 (+56)</td>
<td>0.10 (0.22) 0 (-8) 0.72 (0.52) +3 (+37)</td>
</tr>
</tbody>
</table>
model for estimating $F_{micro}$, $F_{pico,nano}$, and $C_{pico,nano}$, with essentially no change for $C_{micro}$. The MB20-SST model displayed general improvement over the MB20 model.

3.4 Satellite validation of $C_{size}$

Using the satellite-derived data as input, estimates of $C_{size}$ from the regional abundance-based (B-NES-SST, B-NES, and H-NES) and absorption-based algorithms (C02 and MY10) were compared with the in situ pigment-based $C_{size}$ from the independent validation data set. The B-NES-SST, B-NES, and H-NES models displayed similar statistical performance (Fig. 8), although the SST-dependent model performed considerably better across all statistical metrics for $C_{micro}$ (Fig. 8a). The B-NES-SST model generally performed better than the SST-independent B-NES model, and was less constrained by static maximums for $C_{pico,nano}$, $C_{nano}$, and $C_{pico}$ (Fig. 8 f-h, dashed green lines), particularly for $C_{pico}$, for which a substantial increase in the correlation coefficient was observed, consistent with previous studies (Brewin et al., 2017b; Sun et al., 2019b). Like the OC-CCI [Chl-a] input product, the satellite-estimated $C_{size}$ from these models tended to be underestimated at higher concentrations and overestimated at low concentrations, especially for $C_{nano}$ and $C_{pico}$ <0.1 mg m$^{-3}$ and 0.05 mg m$^{-3}$, respectively.

The C02 and MY10 algorithms performed comparably to the re-parameterized abundance-based models (Fig. 9). The MY10 algorithm estimated $C_{micro}$ and $C_{pico,nano}$ with similar differences as the B-NES-SST model, but with higher correlation coefficients ($r = 0.74$ and 0.63, respectively) and improved regression slopes ($S = 0.97$ and 0.89, respectively), although it is noted that the number of validation points was
reduced from $N = 368$ to $N = 352$ and $N = 332$ for C02 and MY10, respectively. For the former, this was due to 16 match-ups with negative satellite $a_{ph}(555)$ retrievals.

**Fig. 8.** Bivariate histograms showing the satellite-to-*in situ* comparisons of $C_{size}$ estimated from the regionally parameterized B-NES-SST (a-d), B-NES (a-d), and H-NES (i-l) abundance-based models, shaded by number of observations. The solid black line is the 1:1 line, dashed black lines indicate the 1:1 line ± 30%, and the red line is the Type-II regression line. Dashed green lines indicate the maximum chlorophyll concentrations imposed by the B-NES model. $N$ denotes the number of match-ups for each parameter, MAD denotes the mean absolute difference, $\delta$ denotes the bias, $r$ denotes the correlation coefficient, and $S$ denotes the regression slope.
Fig. 9. Bivariate histograms showing the satellite-to-in situ comparisons of $C_{size}$ estimated from the absorption-based algorithms applied in this study: (a) $C_{\text{micro,nano}}$ and (b) $C_{\text{pico}}$ from the algorithm of Ciotti et al. (2002) and (c) $C_{\text{micro}}$ and (d) $C_{\text{pico,nano}}$ from the algorithm of Mouw and Yoder (2010). The solid black line is the 1:1 line, dashed black lines indicate the 1:1 line $\pm$ 30%, and the red line is the Type-II regression line. $N$ denotes the number of match-ups, $MAD$ denotes the mean absolute difference, $\delta$ denotes the bias, $r$ denotes the correlation coefficient, and $S$ denotes the regression slope.
while for the latter, 36 match-ups exceeded the \([Chl-a]\) and \(a_{dg}(443)\) detection thresholds of the MY10 algorithm (1.75 mg m\(^{-3}\) and 0.17 m\(^{-1}\), respectively; see Sections 2.7.2.1 and 2.7.2.2). Although the overall bias was generally higher for the absorption-based approaches, they did not exhibit the same overestimation (underestimation) at low (high) concentrations as seen with the abundance-based methods, with the exception of \(C_{micro, nano}\) estimated by C02 model (Fig. 9a).

3.5 Examples of satellite imagery

Considering the overall improved performance of the B-NES-SST algorithm compared with the other abundance-based models, and the statistically similar comparison metrics of the MY10 absorption-based approach, monthly composite imagery from these algorithms was generated for April 2019 and September 2019 for visualization and spatial-temporal comparison. These months were chosen as they were relatively cloud free and displayed contrasting SST and \([Chl-a]\), thus providing some insight into the seasonal variability of phytoplankton size structure in the NES.

Figure 10 shows the monthly imagery of OC-CCI \([Chl-a]\) (Fig. 10a) and MUR SST (Fig. 10b), along with the size class fractions \((F_{micro}, F_{nano}, F_{pico};\) Fig. 10c-e) and size-specific \([Chl-a]\) \((C_{micro}, C_{nano}, C_{pico};\) Fig. 10f-h) from the B-NES-SST algorithm for April 2019. In April, around the time of the typical North Atlantic spring bloom (Friedland et al., 2016), \([Chl-a]\) exceeding 1 mg m\(^{-3}\) was observed both on the shelf and off-shore, with the highest \([Chl-a]\) observed around GB, south of Nova Scotia, and in the near-shore coastal waters along the MAB and GoM. SST ranged from \(<5^\circ C\) in the northern GoM to \(~25^\circ C\) within the Gulf Stream. Microplankton were dominant in low SST, high
Fig. 10. Monthly composite imagery for April 2019: (a) OC-CCI \([\text{Chl-a}]\), (b) MUR SST, (c) \(F_{\text{micro}}\), (d) \(F_{\text{nano}}\), (e) \(F_{\text{pico}}\), (f) \(C_{\text{micro}}\), (g) \(C_{\text{nano}}\), and (h) \(C_{\text{pico}}\) from the B-NES-SST model. Color scales for \(F_{\text{size}}\) are adjusted to reflect the range of the model for each size class (see Fig. 8). The black line indicates the 400 m isobath to mark the approximate location of the shelf break.

\[\text{Chl-a}\] waters in the GoM, on GB, along the coast, and within the major estuaries. Nanoplankton were most prevalent in the intermediate \([\text{Chl-a}]\) and SST waters off-
shore. Picoplankton was the dominant size class in the oligotrophic, high SST surface waters of the Gulf of Stream.

Figure 11 shows the same as Figure 10 but for September 2019. Compared with April, in September areas of [Chl-a] > 1 mg m$^{-3}$ did not extend as far off-shore but were mainly confined to near-shore regions of the MAB, the GoM, and GB. There was a strong gradient in [Chl-a] from the highly productive waters along the coast and to the north to the oligotrophic ([Chl-a] < 0.1 mg m$^{-3}$) off-shore waters to the south. SST
Fig. 11. Monthly composite imagery for September 2019: (a) OC-CCI \(Chl-a\), (b) MUR SST, (c) \(F_{micro}\), (d) \(F_{nano}\), (e) \(F_{pico}\), (f) \(C_{micro}\), (g) \(C_{nano}\), and (h) \(C_{pico}\) from the B-NES-SST model, as per the previous figure.

exceeded 20 °C throughout much of the region, with cooler SST observed to the north around GB and within the GoM, coinciding with the higher \(Chl-a\) observed in these areas. Microplankton comprised a smaller fraction of \(Chl-a\) in both near-shore and off-shore waters in September, with nanoplanckton becoming more dominant on the shelf, particularly in the central GoM and areas immediately surrounding GB. Likewise, the contribution of nanoplanckton generally decreased off-shore in September, with picoplankton becoming more dominant in these waters, coinciding with higher SST and lower \(Chl-a\) than was observed in April.

A comparison of \(F_{micro}\) imagery from the B-NES-SST and MY10 algorithms for April 2019 and September 2019 is presented in Figure 12. Considering the \(F_{micro}\) output from the MY10 algorithm is binned to increments of 0.1, the color scale for the B-NES-SST imagery was adjusted to match the scale of the MY10 imagery to facilitate a more visually equitable comparison. The two approaches displayed similarities and differences in the spatial patterns and extent of estimated \(F_{micro}\). For example, both models indicated higher \(F_{micro}\) on GB and in the northern GoM than the surrounding region in September, while in April both models show areas of elevated \(F_{micro}\) extending farther offshore, beyond the shelf break. Pixels exceeding the MY10 thresholds of detection (\([Chl-a] > 1.75 \text{ mg m}^{-3} \) and \(a_{dg}(443) > 0.17\), plotted in white) were located predominantly in shallow regions very close to shore and within major
embayments, with some masked pixels on GB. While the two algorithms showed some similar spatial patterns, there were differences in the magnitude of the estimated $F_{\text{micro}}$. For instance, the MY10-estimated $F_{\text{micro}}$ was higher around GB and areas in the GoM and the northern MAB than the B-NES-SST model in September, and higher

![Image of monthly $F_{\text{micro}}$ imagery from the B-NES-SST and MY10 algorithms for April 2019 (a, b) and September 2019 (c, d). The MY10 algorithm applies a 2-D average filter, masks pixels that exceed defined thresholds of $\text{Chl-a}$ and $a_{dg}(443)$ (plotted in white), and bins $F_{\text{micro}}$ to increments of 0.1 (see Section 825).](image)

**Fig. 12.** Comparison of monthly $F_{\text{micro}}$ imagery from the B-NES-SST and MY10 algorithms for April 2019 (a, b) and September 2019 (c, d). The MY10 algorithm applies a 2-D average filter, masks pixels that exceed defined thresholds of $\text{Chl-a}$ and $a_{dg}(443)$ (plotted in white), and bins $F_{\text{micro}}$ to increments of 0.1 (see Section 825).
2.7.2.2). The color scale for the B-NES-SST imagery was modified to match the output of the MY10 algorithm. The black line indicates the 400 m isobath to mark the approximate location of the shelf break.

within the off-shore feature of elevated [Chl-a] located around 38ºN, 69ºW in April. The MY10 imagery also generally showed a higher degree of spatial variability in $F_{micro}$ compared with that of the B-NES-SST model. This is evident within the central GoM in April, where the MY10 imagery showed areas of $F_{micro}$ spanning the full fractional range (i.e., 0-1), whereas the B-NES-SST imagery showed a much more uniform distribution of $F_{micro}$, ranging only between 0.5 and 0.7.

4. Discussion and conclusions

The focus of this study was the regional refinement and evaluation of PSC algorithms in the NES. Like many similar studies, *in situ* estimates of PSCs derived from HPLC pigment data using the DPA method were used for model re-parameterization and statistical comparisons. While this approach is a popular choice given the relative abundance of HPLC samples compared with other methods for quantifying PSCs *in situ*, it has important limitations. First, DPA is not a direct measure of cell size, but rather an approximation of size structure based on assumptions about the taxonomic groups attributed to different pigments, and the size classes represented by those taxa. In reality, pigments are not perfectly diagnostic, and are known to be shared across multiple taxa in varying concentrations dependent on physiological state (Uitz et al., 2008). Further, taxonomic groups may span multiple size classes in ways that are not fully represented by the DPA equations (Leblanc et
Although proposed modifications to account for some of these biases were incorporated in this study (Devred et al., 2011), the efficacy of this specific approach for characterizing PSCs in the NES region is uncertain and warrants further investigation. Recently, Chase et al., (2020) evaluated the DPA method by comparing pigment-based PSC estimates to coincident measurements of cell size by imaging-in-flow and conventional flow cytometry in the North Atlantic and found that DPA overestimated micro- and picoplankton and underestimated nanoplankton relative to cytometry. They recommended a revised set of DPA equations to better account for the presence of dinoflagellates and diatoms in the nanoplankton, and the presence of $[TChl-b]$ in both pico- and nanoplankton. To reduce uncertainty on this front, continued efforts to inter-compare multiple in situ PSC methods across different oceanic environments will be extremely valuable.

Abundance-based PSC algorithms are attractive for their ease of implementation, using satellite $[Chl-a]$ as the sole input parameter, and have been shown to perform well globally (Brewin et al., 2011; R. J. W. Brewin et al., 2015) and in a variety of oceanic regions (Brito et al., 2015; Di Cicco et al., 2017; Gittings et al., 2019; Sun et al., 2018). Here, previously published abundance-based methods were tested in the NES using both original model parameters and new parameters derived from a region-specific HPLC pigment data set. Our results indicated that regional re-parameterization alone offered minimal statistical improvement compared with other models evaluated, which included both globally and regionally parameterized models. All eight of the abundance-based models tested yielded similar comparison metrics, particularly for the micro- and combined pico- and nanoplankton classes, when
applied to the *in situ* [Chl-a] and compared with the DPA-based size class estimates from the independent *in situ* validation data set. There was more variation in the comparison metrics for separated nano- and picoplankton classes, but in no instance were the re-parameterized models exclusively the best performing, with the exception of the H-NES model for estimating picoplankton, which showed slight improvement relative to the other models when applied to both the *in situ* and satellite [Chl-a]. Based on the regional validation conducted in this study, the OC-CCI [Chl-a] product was chosen as the satellite input for model application and comparison. However, the systematic positive and negative bias of this product at low and high [Chl-a] respectively was directly translated to the $C_{size}$ estimates from the regionalized abundance-based methods, underscoring the importance of the satellite [Chl-a] input on the accuracy and uncertainty abundance-based PSC output.

While re-parameterization alone provided little benefit in terms of improving abundance-based model performance in the NES, the incorporation of remotely sensed SST into the re-parameterization of the B10 model did consistently improve PSC prediction accuracy. When applied to the *in situ* validation data set, the regional SST-dependent B-NES-SST model showed a reduction in MAD of 10-12% for all size fractions with respect to the SST-independent B-NES model. The B-NES-SST model also outperformed the other two SST-dependent models tested (B17-SST and MB20-SST) for all size classes. This result supports what has been shown by previous studies (Brewin et al., 2017b; Moore and Brown, 2020; Sun et al., 2019b; Ward, 2015) - that the addition of SST into abundance-based model frameworks can improve PSC prediction accuracy. The relationships between [Chl-a], SST, and phytoplankton size
structure observed in this study were also in general agreement with the findings of previous studies, with lower SST associated with an increase in the fraction of microplankton and a decrease in the fraction of smaller cells (i.e. pico- and nanoplankton) at similar [Chl-a]. This relationship is not entirely surprising, given long-established connections between temperature, water-column stability, nutrient availability, and phytoplankton community size structure in the marine environment (Bouman et al., 2003; Margalef, 1978). While SST is used as the additional predictor variable in these models, the associated changes in size structure are not necessarily in direct response to changes in SST but rather the result of a combination of co-varying environmental conditions, including light availability, stratification, and nutrient availability. Further in-depth analysis of the interactions between environmental, hydrodynamic, and biogeochemical/optical parameters, similar to that conducted by Mouw et al. (2019), is required to understand which of these variables are the predominant drivers of phytoplankton size variability in the NES region.

Absorption-based algorithms are advantageous over abundance-based methods in that they are rooted in a direct spectral response to phytoplankton cell size, as opposed to relying on indirect statistical connections between [Chl-a] and phytoplankton size structure. This means that they enable detection of changes in PSCs that occur outside of the general biomass-size co-variation relationship and are less prone to change temporally or geographically. When applied directly to the satellite data, the two absorption-based algorithms examined in this study performed with comparable accuracy to the regionalized abundance-based models. The MY10 algorithm showed statistically similar performance to the SST-dependent model,
without the inclusion of additional environmental information. Considering that pigment-based size class estimates from DPA were used for validation, the similarity in performance for the absorption-based algorithms is encouraging given they were not developed or parameterized using DPA, as was the case with the abundance-based methods. This suggests consistency between estimates of PSCs derived from spectral phytoplankton absorption and those determined from HPLC pigment analysis in the NES, as has been previously reported in other regions (Devred et al., 2011). Moreover, as they are not solely reliant on [Chl-a] for deriving size, the satellite-estimated $C_{size}$ from the absorption-based approaches did not suffer from the same bias at high and low [Chl-a] inherent in the OC-CCI [Chl-a]. Nevertheless, as is the case with abundance-based approaches, any biases in satellite $a_{ph}(\lambda)$ or $a_{dg}(\lambda)$ used as input will be directly translated to the PSC output. Despite their strengths, the absorption-based methods were unable to characterize areas where $a_{ph}(\lambda)$ retrievals were poor (negative $a_{ph}(555)$ in the case of the CO2 model) or, in the case of the MY10 model, where [Chl-a] and $a_{dg}(443)$ were outside of set thresholds. These issues will be more prevalent in optically complex waters such as the NES and may be a drawback to the use of these algorithms in regions of high biomass or large relative optical contribution of CDOM and NAP.

While less common than abundance-based and absorption-based methods and outside of the focus of this study, approaches that exploit satellite backscattering data to derive PSCs have also been developed (Kostadinov et al., 2016, 2009; Montes-Hugo et al., 2008). These "backscattering-based" methods share many of the same advantages of absorption-based algorithms, as they utilize the direct spectral
response of in-water constituents and are less reliant on empirical relationships, making them less susceptible to ecological shifts in a changing ocean. However, current approaches estimate PSCs under the assumption that the particle assemblage is totally biogenic in nature, an assumption that generally holds in the open ocean but can be violated in coastal systems where inorganic particles are introduced through processes such as river discharge and tidal mixing (Kostadinov et al., 2016). Nonetheless, backscattering information provides an important tool for remotely estimating the size structure of phytoplankton communities and its use in combination with methods based on other optical and environmental data has potential to greatly improve PSC retrievals.

The PSC algorithms and products evaluated in this study may be useful for the validation or assimilation of regional ecosystem or biogeochemical models (IOCCG, 2020). However, given the uncertainties associated with the pigment-based size class estimates used for algorithm assessment, as well as the different inputs and outputs between methods, it is difficult to make a definitive determination of which approach is the best choice for such applications. The most suitable method may be dependent on the specifics of the intended application or the questions to be addressed. For instance, biogeochemical models that produce chlorophyll-based phytoplankton size estimates may prefer to compare to abundance-based algorithm outputs, while models that include optics may prefer to compare to output from absorption-based methods – each enabling a more direct comparison depending on the underlying algorithm/model frameworks and outputs being compared.
In the near future, satellite ocean color remote sensing is moving toward more advanced radiometric instruments with hyperspectral capability and enhanced spatial and temporal resolution (Cetinić et al., 2018). The increased spectral information afforded by these upcoming sensors is anticipated to greatly improve our ability to accurately separate the absorption attributed to different optically significant in-water constituents (i.e., CDOM, NAP, phytoplankton) and retrieve information on phytoplankton community composition and size structure. This improved capability will be particularly relevant to optically complex waters, including coastal and continental shelf regions like the NES ecosystem. Thus, existing absorption-based PSC models may potentially become more robust, and newer methods that exploit the full range of available spectral information will continue to be developed. Furthermore, to the extent that satellite [Chl-a] estimates improve as a result of the increase in spectral resolution, abundance-based approaches may continue to be an effective option for estimating PSCs, especially when combined with SST or other ecologically relevant environmental parameters. While not considered in this work, the integration of high-resolution spectral information with environmental data readily attainable from remote sensing should be considered in future PSC algorithm development efforts.

6. Appendix A

Statistical metrics comparing the in situ pigment-based estimates of $F_{size}$ and $C_{size}$ from the parameterization and validation data sets with estimates from the abundance-based models shown in Figure 6. Statistical calculations were performed in linear space for $F_{size}$ and log$_{10}$ space for $C_{size}$. Metrics for $C_{size}$ are shown in parentheses.
5. References


Mobley, C.D., Sundman, L.K., 2013. HYDROLIGHT 5.2 ECOLIGHT 5.2 Technical Documentation.


6. List of Figure and Table Captions

Table 1. Symbols and definitions.

Fig.1. Locations of the in situ data and satellite match-ups used in this study. The 200 and 2000 m isobaths from the 2019 General Bathymetric Chart of the Oceans.
(https://www.gebco.net/) are shown for reference. See Table 2 for information on data sources.

**Table 2.** Summary of *in situ* data sources. *N* denotes the number of samples (after QA) and the number in parentheses refers to the number of satellite match-ups. Citations for the individual data sets from SeaBASS are also provided.

**Fig. 2.** Relative frequencies of *in situ* observations from the NES data set used in this study (blue stairs): (a) HPLC-measured [Chl-a] (*N* = 786), (b) *a*$_{ph}$(443) (*N* = 214) and (c) *a*$_{dg}$(443) (*N* = 173), with their respective monthly distributions (d-f, black bars). Global distributions from an OC-CCI v4.2 annual satellite composite for 2018 are overlaid for comparison (red line). Frequencies were normalized by the maximum value.

**Fig. 3.** Comparison of the HPLC-measured [Chl-a] (*C$_{HPLC}$*) and the reconstructed [Chl-a] from the sum of seven diagnostic pigments (*C$_{DP}$*) using the weighting coefficients derived from this study (green circles; $MAD = 0.12, r = 0.98$), the weights derived by Uitz et al. (2006) from a global data set (blue triangles; $MAD = 0.47, r = 0.96$), and no weighting coefficients (magenta squares; $MAD = 0.62, r = 0.96$).

**Table 3.** Diagnostic pigments [P] and their associated taxonomic groups and attributed size classes, along with weighting coefficients [W] obtained from this study
and previous studies. The number of data points and geographical regions of each study are also provided.

Table 4. Parameter values for the abundance-based models of Brewin et al. (2010) (Eqs. 9 and 10) and Hirata et al. (2011) (Eq. 16), obtained from this study and from previous studies.

Fig. 4. Brewin et al. (2010) model parameters (a) $C_{m_{\text{pico,nano}}}^{m_{\text{pico}}}$, and (b) $D_{\text{pico,nano}}/D_{\text{pico}}$ as a function of SST. Circle markers show the median parameter values obtained by performing a running bootstrap fit of the model (Eqs. 9 and 10) to the in situ parameterization data set ($N = 418$) sorted by increasing SST. The SST-parameter relationships were smoothed using a 5-point running average. Dashed lines indicate the SST-independent model (B-NES) parameters obtained when fitting the model to the full parameterization data set.

Fig. 5. Bivariate histograms showing the satellite-to-in situ comparisons of [Chl-$a$], $a_{\text{ph}}(\lambda)$, and $a_{\text{dg}}(\lambda)$, shaded by number of observations: (a) [Chl-$a$] from the standard OC-CCI algorithm, (b) [Chl-$a$] from the regional algorithm of Pan et al. (2010), (c) $a_{\text{ph}}(443)$, (d) $a_{\text{ph}}(490)$, (e) $a_{\text{ph}}(510)$, (f) $a_{\text{dg}}(443)$, (g) $a_{\text{dg}}(490)$, and (h) $a_{\text{dg}}(510)$ from the standard OC-CCI algorithm (QAA_v5). The solid black line is the 1:1 line, dashed black lines indicate the 1:1 line ± 30%, and the red line is the Type-II regression line. $N$ denotes the number of match-ups, MAD denotes the mean absolute difference, $\delta$ denotes the bias, $r$ denotes the correlation coefficient, and $S$ denotes the regression
slope. The $a_{ph}(\lambda)$ and $a_{dg}(\lambda)$ data are shown using the same x- and y-axis range for comparison.

**Fig. 6.** Pigment-based estimates of $F_{\text{size}}$ (a-d) and $C_{\text{size}}$ (e-h) as a function of $C_{\text{HPLC}}$ from the parameterization data set ($N = 418$) with abundance-based models overlaid: re-parameterized B10 model (B-NES, solid black), re-parameterized H11 model (H-NES, dashed black), Brewin et al., 2010 (B10, blue), Brewin et al., 2015a (B15, green), Brewin et al., 2017 (B17, red), Devred et al., 2011 (D11, violet), Hirata et al., 2011 (H11, brown), and Moore and Brown, 2020 (MB20, yellow).

**Fig. 7.** Modeled $F_{\text{size}}$ (a-d) and $C_{\text{size}}$ (e-h) from the B-NES-SST model plotted as a function of [Chl-a], with the color gradient illustrating the changes in the model when model parameters vary as a function of SST (see Section 2.7.1.7, Fig. 4). The black line indicates the SST-independent model, with a single set of model parameters (B-NES).

**Table 5.** Mean absolute difference ($MAD$) and correlation coefficients ($r$) for $F_{\text{size}}$ and $C_{\text{size}}$ (values for $C_{\text{size}}$ shown in parentheses) for the SST-dependent abundance-based models applied to the in situ parameterization and validation data sets. The percent change in the metrics when incorporating SST relative to the SST-independent models is included for reference. Percentages are rounded to the nearest 1%. 

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Fig. 8. Bivariate histograms showing the satellite-to-in situ comparisons of $C_{size}$ estimated from the regionally parameterized B-NES-SST (a-d), B-NES (a-d), and H-NES (i-l) abundance-based models, shaded by number of observations. The solid black line is the 1:1 line, dashed black lines indicate the 1:1 line ± 30%, and the red line is the Type-II regression line. Dashed green lines indicate the maximum chlorophyll concentrations imposed by the B-NES model. $N$ denotes the number of match-ups for each parameter, $MAD$ denotes the mean absolute difference, $\delta$ denotes the bias, $r$ denotes the correlation coefficient, and $S$ denotes the regression slope.

Fig. 9. Bivariate histograms showing the satellite-to-in situ comparisons of $C_{size}$ estimated from the absorption-based algorithms applied in this study: (a) $C_{micro,nano}$ and (b) $C_{pico}$ from the algorithm of Ciotti et al. (2002) and (c) $C_{micro}$ and (d) $C_{pico,nano}$ from the algorithm of Mouw and Yoder (2010). The solid black line is the 1:1 line, dashed black lines indicate the 1:1 line ± 30%, and the red line is the Type-II regression line. $N$ denotes the number of match-ups, $MAD$ denotes the mean absolute difference, $\delta$ denotes the bias, $r$ denotes the correlation coefficient, and $S$ denotes the regression slope.

Fig. 10. Monthly composite imagery for April 2019: (a) OC-CCI [Chl-a], (b) MUR SST, (c) $F_{micro}$, (d) $F_{nano}$, (e) $F_{pico}$, (f) $C_{micro}$, (g) $C_{nano}$, and (h) $C_{pico}$ from the B-NES-SST model. Color scales for $F_{size}$ are adjusted to reflect the range of the model for each size class (see Fig. 8). The black line indicates the 400 m isobath to mark the approximate location of the shelf break.
**Fig. 11.** Monthly composite imagery for September 2019: (a) OC-CCI [Chl-a], (b) MUR SST, (c) $F_{micro}$, (d) $F_{nano}$, (e) $F_{pico}$, (f) $C_{micro}$, (g) $C_{nano}$, and (h) $C_{pico}$ from the B-NES-SST model, as per the previous figure.

**Fig. 12.** Comparison of monthly $F_{micro}$ imagery from the B-NES-SST and MY10 algorithms for April 2019 (a, b) and September 2019 (c, d). The MY10 algorithm applies a 2-D average filter, masks pixels that exceed defined thresholds of [Chl-a] and $a_{dg}(443)$ (plotted in white), and bins $F_{micro}$ to increments of 0.1 (see Section 2.7.2.2). The color scale for the B-NES-SST imagery was modified to match the output of the MY10 algorithm. The black line indicates the 400 m isobath to mark the approximate location of the shelf break.