Phenology of particle size distributions and primary productivity in the North Pacific subtropical gyre (Station ALOHA)

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Abstract The particle size distribution (PSD) is a critical aspect of the oceanic ecosystem. Local variability in the PSD can be indicative of shifts in microbial community structure and reveal patterns in cell growth and loss. The PSD also plays a central role in particle export by influencing settling speed. Satellite-based models of primary productivity (PP) often rely on aspects of photophysiology that are directly related to community size structure. In an effort to better understand how variability in particle size relates to PP in an oligotrophic ecosystem, we collected laser diffraction-based depth profiles of the PSD and pigment-based classifications of phytoplankton functional types (PFTs) on an approximately monthly basis at the Hawaii Ocean Time-series Station ALOHA, in the North Pacific subtropical gyre. We found a relatively stable PSD in the upper water column. However, clear seasonality is apparent in the vertical distribution of distinct particle size classes. Neither laser diffraction-based estimations of relative particle size nor pigment-based PFTs were found to be significantly related to the rate of ¹⁴C-based PP in the light-saturated upper euphotic zone. This finding indicates that satellite retrievals of particle size, based on particle scattering or ocean color, would not improve parameterizations of present-day bio-optical PP models for this region. However, at depths of 100–125 m where irradiance exerts strong control on PP, we do observe a significant linear relationship between PP and the estimated carbon content of 2–20 μm particles.

1. Introduction

In an elegant text on the scattering of light by marine particles, Jonasz and Fournier (2011) state that “one could argue that seawater is nothing else but suspended particles, whose sizes range from molecules through fish and whales.” Undeniably, seawater carries a rich load of particles including colloids (0.001–1 μm), organisms of various morphologies (autotrophs and heterotrophs, 0.001–1000 μm), viruses, detrital material (nanometer–millimeter), and multicellular organisms (>100 μm). The size distribution of these particles follows a classic pattern—the concentration of particles rapidly decreases with increasing particle size (Jerlov, 1976; Sheldon et al., 1972). In effect, this tells us that the larger the particle is, the rarer it is. From a biological perspective, changes in the particle size distribution largely reflect the time-variant growth and loss of organisms of various sizes: for example as a result of the seasonal succession from smaller dinoflagellates to larger diatoms in the coastal ocean (Smayda and Trainer, 2010), as episodic blooms of large nitrogen-fixing organisms (e.g., Trichodesmium spp.) and diatom-diazotroph associations occur in the open ocean, (Letelier and Karl, 1996; Scharek et al., 1999) or as a result of size-selective protist grazing on small bacterioplankton (Epstein and Shiari, 1992; Hagström et al., 1986).

Planktonic organisms, including both heterotrophs and autotrophs, contribute substantially to total oceanic particle loads and are the dominant particulate component in the scattering and absorption of light in the surface waters of the ocean (Field et al., 1998; Jonasz and Fournier, 2011; Stramski et al., 2001). Phytoplankton, the autotrophic fraction of the plankton residing in the euphotic zone, contribute roughly half of the net photosynthetic carbon fixation on the planet. The cell sizes of individual phytoplankton, which can range from roughly 0.5 to 1000 μm, impact surface area to volume ratios and hence nutrient use efficiency as well as the chlorophyll-specific absorption of light and the quantum efficiency of photosynthesis (Bricaud et al., 2004;
Sathyendranath et al., 1987). Each of these size-specific differences can lead to variation in community photosynthetic rates. Accordingly, a relatively nascent effort is being made to evaluate the extent to which the relative size distribution of phytoplankton communities drives observed variability in the rate of net primary productivity (NPP, see review by Nair et al. [2008]) and ultimately export [e.g., Guidi et al., 2009]. The implicit but often unstated assumption is that larger cells lead to higher growth rates and enhanced productivity [Chisholm, 1992; Schlesinger et al., 1981]. These relationships are readily apparent when looking across large gradients in particle loads and productivity, for example, from upwelling regimes to the oligotrophic gyres, but are less clear within regimes or within taxa [Chisholm, 1992; Marañón et al., 2001].

Derivations of community structure and hence particle size structure are increasingly being seen as a means of potentially constraining variability in productivity, as they reflect snapshots of phytoplankton species composition or phytoplankton functional types [Claustre et al., 2005; Kostadinov et al., 2010; Uitz et al., 2008]. An example of how shifts in phytoplankton community structure (and the PSD) relate to NPP in oligotrophic regions is found in the work of Ondrusek et al. [2001]. These authors developed a primary production model in the subtropical North Pacific Ocean using in situ measurements of chlorophyll, the quantum yield of photosynthesis, and chlorophyll-specific absorption coefficients. Although the model captured the mean productivity of this system, it was only able to account for 50% of the variability in measured production rates. The authors concluded that “understanding community shifts from small prokaryote dominated systems to large eukaryote dominated systems appears to be one of the key elements to improving the performance” of bio-optical models in the open ocean [Ondrusek et al., 2001]. Similarly, Li et al. [2011] report that while less abundant, “larger” phytoplankton (>2 μm) in the North Pacific subtropical gyre (NPSG) appear more efficient at carbon fixation than smaller-celled organisms. The authors then hypothesize that shifts in community structure toward these “larger” cells would lead to enhanced productivity in this system. Contrarily, increased abundance of small cells has also been linked to variability in NPP in other oligotrophic regions. Lomas et al. [2010] report coherence between an enigmatic increase of the cyanobacterium Synechococcus between 1996 and 2007 and enhanced NPP in the Sargasso Sea of the North Atlantic. The authors hypothesized that this increase was due to the ability of Synechococcus to enhance growth rates and accumulate biomass in response to nanomolar pulses of nitrate. This finding indicates that shifts in photosynthetic or nutrient use efficiency may be more important to variability in NPP than particle size alone [Chisholm, 1992].

Several studies also report poor relationships between paired measurements of relative particle size and primary productivity [Hayward and Venrick, 1982; Marañón et al., 2003; Marañón et al., 2007; Marañón et al., 2001]. Marañón et al. [2003] in particular conclude that the lack of a relationship between changes in chlorophyll size-fractions and NPP in the oligotrophic Atlantic imply that “microbial communities in oligotrophic regimes respond to environmental forcing with significant changes in primary productivity that are not associated with trophic shifts.” Taken together, it is not clear whether the available data from oligotrophic ocean regimes fully support a reliable relationship between either the PSD or PFTs and NPP. Positive enhancements in productivity may be more closely related to efficiency of energy capture or uncoupling of growth and loss terms rather than changes in community structure.

These conflicting conclusions regarding the role of phytoplankton community composition as a driver of primary productivity are potentially limited by methodological constraints inherent to measurement of both productivity [Peterson, 1980 for known biases in the 14C tracer method] and estimations of community composition. For example, an increase in one class of cells (e.g., Synechococcus) does not necessarily imply a shift in the mean PSD. Additionally, physical filtration-based size-separation may have artifacts (e.g., absorption of dissolved organics onto filters, membrane clogging, inefficient trapping of cells, or cell breakage) [as examples see Gosli and Moran [1999]; Sørensen et al. [2013]]. There are a number of other approaches to the characterization of phytoplankton community structure that may prove useful in further elucidating potential linkages between PSD and NPP, albeit none are without disadvantage. For instance, several studies [Bricaud et al., 2004; Kostadinov et al., 2010; Uitz et al., 2008; Vidussi et al., 2001] have used high-performance liquid chromatography (HPLC) of diagnostic pigments to estimate the relative proportions of phytoplankton functional types: picophytoplankton (0.2–2.0 μm), nanophytoplankton (2–20 μm), and microphytoplankton (>20 μm). These groupings are based on taxonomical classifications of Sieburth et al. [1978]. The primary advantage of this approach is that pigmented particles reflect a diverse consortium of living phytoplankton with no influence from heterotrophic or detrital material. Disadvantages are that filtration is required, diagnostic pigments may be shared by various phytoplankton groups, pigment concentrations per cell may vary with light history and exposure, and size...
classes are somewhat arbitrary given the known size continuum of phytoplankton classes [Marañón et al., 2007]. Nonetheless, Briceaud et al. [2004] report that the relative proportions of these pigment-based classes are tightly linked to variation in the chlorophyll-specific absorption coefficient (\(a^t, m^2\text{mg chlorophyll}^{-1}\)) and the efficiency of light absorption. Given that the general formulation of many bio-optical models for the estimation of marine primary productivity relies on \(a^t\), this finding has direct implications for the role of community structure on NPP, albeit the relationship between pigment based PFTs and NPP has not been thoroughly explored in open ocean regimes.

A second approach to measure the PSD is via laser diffraction, where a laser beam illuminates a sample volume containing particles and an inversion of the volume scattering function at small forward angles is performed to retrieve the in situ particle size distribution [Agrawal et al., 1991]. A commercial in situ laser diffractionometer (LISST-100X, Laser In Situ Scatterometer/Transmissometer, Sequoia Scientific Inc., hereafter simply LISST) has been utilized in both oceanic and lake environments to study particle dynamics [Barone et al., 2015; Gartner et al., 2001; Mikkelsen and Pejrup, 2001; Serra et al., 2001]. The principle of operation is straightforward: a collimated laser beam is projected through a sample volume having a 5 cm path length. The angular scattering distribution between 0.1° and 20° at 670 nm is measured at a silicon detector with 32 log-spaced rings and the scattering signal is inverted assuming some particle shape [Agrawal et al., 1991; Agrawal et al., 2008]. There is no mechanical aperture, so forward scattering at each angle is collected simultaneously and the size distribution of particles within 1.25 and 250 \(\mu\text{m}\) (equivalent spherical diameter, ESD) is rapidly assessed in a high-throughput fashion (1 Hz). The LISST has been rigorously tested in the laboratory with suspensions of beads, phytoplankton cultures, and various irregularly shaped particles (e.g., sieved sediments) [Agrawal et al., 2008; Karp-Boss et al., 2007]. In side-by-side tests, LISST estimations of the particle size distribution, volume concentration, and mean particle size agree well with results from the Coulter counter [Reynolds et al., 2010], microscopy [Groundwater et al., 2012], and a digital camera (silhouette photography with a LED flash) [Mikkelsen et al., 2005]. The primary disadvantages of laser diffraction are that retrieval of the PSD requires knowledge of a static inversion function which may not adequately reflect the population, individual particles are not analyzed directly, and the method cannot discriminate between living and particulate detrital material.

In order to improve our understanding of the relationship between particle size and NPP in the oligotrophic NPSG, we have investigated particle and productivity dynamics in a region that has been sampled by the Hawaii Ocean Time-series (HOT) program at roughly monthly intervals since 1988 [Karl and Lukas, 1996]. The location of sampling is 100 km north of Oahu, Hawaii at Station ALOHA (A Long-Term Oligotrophic Habitat Assessment; 22° 45’N, 158° 00’W), in approximately 4750 m of water. This region was chosen because of the rich biogeochemical data available and as a representative of oligotrophic ocean regimes. Additionally, key transitions in the upper ocean physical forcing [Karl and Lukas, 1996] and in situ measurements of \(^{14}\text{C}-\)based primary productivity at Station ALOHA are available for December 1988 to December 2012; LISST measurements of the PSD are available from September 2009 to April 2014.

2. Methods

To evaluate shifts in phytoplankton communities, we use a combination of pigment-based approximations of phytoplankton functional types and laser diffraction-based particle size distributions. Time-variant shifts in these independent metrics are compared to rate determinations of primary productivity. HPLC data and in situ measurements of \(^{14}\text{C}-\)based primary productivity at Station ALOHA are available for December 1988 to December 2012; LISST measurements of the PSD are available from September 2009 to April 2014.

2.1. Discrete Samples and Methods

As a part of the HOT program core measurement set, seawater is collected and filtered onto 25 mm diameter glass fiber filters (Whatman GF/F) for pigment analysis by HPLC. Samples are analyzed via protocols...
not assume strict size classes for PFTs even though the prefixes (e.g., “pico”) used here do imply size. Data

tation ALOHA (Letelier et al., 2002) on the slope of the PSD in surface waters (<20 m) and to exclude depths well below the 1% sur-
face light level (95–130 m) [Letelier et al., 2004] where net photosynthetic carbon assimilation is minimal. Selection of a 20 m threshold for bubble injection is also guided by the observation that wind speeds at Station ALOHA (<12 m s⁻¹, Woods Hole/HOT mooring meteorological data, http://www.soest.hawaii.edu/

Table 1. Diagnostic Pigments, Representative Algal Classes for Station ALOHA, and General Size Bins From the Algorithms by Bricaud et al. [2004] and Uitz et al. [2006].

| Diagnostic Pigment | Algal Class | Grouping |
|--------------------|-------------|----------|
| Zeaxanthin         | Primarily cyanobacteria | Pico     |
| Chlorophyll b      | Prochlorococcus | Pico     |
| Alloxanthin        | Cryptophyceae | Nano     |
| 19′-butanoyloxyfucoxanthin (19′BF) | Haptophyceae and Pelagophyceae | Nano     |
| 19′-hexanoyloxyfucoxanthin (19′HF) | Primarily Haptophyceae | Nano     |
| Fucoxanthin        | Primarily Bacillariophyceae (diatoms) | Micro    |
| Peridinin          | Dinophyceae | Micro    |

*We do not assume strict delineation of size between the classes of picoplankton (“pico”), nanoplanckton (“nano”), or microplankton (“micro”); however, we do assume that these pigments correspond to distinct phytoplankton functional types defined by the corresponding algal classes.

When applying these formulations for Station ALOHA, the following must be considered: (1) peridinin concentrations are uniformly low (<7 ng L⁻¹) at this location such that calculations of the fraction of microplankton are largely driven by fucoxanthin levels (0–70 ng L⁻¹ range) and hence diatoms; (2) while there are Chl b containing eukaryotes (e.g., chlorophytes and prasinophytes), Chl b at Station ALOHA is primarily derived from Prochlorococcus spp. [Andersen et al., 1996]; (3) alloxanthin concentrations are rarely detectable at Station ALOHA and so nanoplanckton calculations are driven by biomarkers for pelagophytes and prymnesiophytes, and (4) the size classes are rough approximations; in fact this approach groups picocaryophytes into the nanoplancton and small diatoms (<20 μm) as microphytoplankton. For this reason, we do not assume strict size classes for FPTs even though the prefixes (e.g., “pico”) used here do imply size. Data have been compared to in situ ¹⁴C-based measurements of dawn to dusk primary productivity rates available for this same time period; methods for the deployment and processing of the HOT ¹⁴C array are described in Letelier et al. [1996].

2.2. LISST Deployment

The LISST was deployed on a bio-optical package approximately 1.5 m³ in size. The instrument was mounted horizontally with open water flow through the optical path and lowered at a constant descent rate of 10 m min⁻¹ to a depth of ~200 m using the ship’s winch. The package was deployed on an approximately monthly basis between September 2009 and April 2014 for a total of 42 cruises; 2–3 night casts were performed during each cruise. In the present work, data were only used from (1) downcasts so that the instrument was seeing undisturbed water and (2) profiles collected at night in order to avoid light contamination that would otherwise lead to erroneous concentrations of particles in the smallest size bins [Andrews et al., 2011]. From these casts, only data from depths below 20 m and above 175 m were considered. The rationale for the last criterion is to avoid the potential influence of entrained air bubbles [Zhang et al., 2002] on the slope of the PSD in surface waters (<20 m) and to exclude depths well below the 1% surface light level (95–130 m) [Letelier et al., 2004] where net photosynthetic carbon assimilation is minimal. Selection of a 20 m threshold for bubble injection is also guided by the observation that wind speeds at Station ALOHA (<12 m s⁻¹, Woods Hole/HOT mooring meteorological data, http://www.soest.hawaii.edu/
whots/) correspond to a bubble penetration depth of <15 m [Vagle et al., 2010]. These considerations (avoidance of ambient light and contamination of the scattering signal by bubbles) are absolutely necessary for proper interpretation of LISST data in oceanic settings [also see Barone et al., 2015].

2.3. LISST Description, Processing Methods, and Rationale

Before the light scattering distribution measured by the LISST is inverted to obtain the particulate volume distribution, the signal must be corrected for background scattering due to pure water and any imperfections of the optics that would cause instrument drift (referred to as the "zscat" by the manufacturer). In the oligotrophic waters at Station ALOHA, we found that the mean LISST raw scattering signal in deep water (~200 m, raw counts = 32 ± 5 over 32 rings and 42 cruises) was equivalent to our on-deck background measurements using deionized, reverse osmosis water from the ship’s system (raw counts = 34 ± 5). The p values of a two-sample t test ranged from 0.2 to 0.9; deep water and clean water backgrounds were not significantly different. For consistency, we used the mean of the deepest 2 m of the water column profile as our background (maximal depths of downcast ranged from 176 to 230 m); maximal depths achieved were a function of current speed and the extent to which the optical package descended vertically. While these backgrounds represent the lower detection limit of the LISST, they are not particle-free; the concentration of particulate carbon at 200 m at Station ALOHA is ~0.7 μmol L⁻¹. After correction of the raw scattering signal for this background, we verified that the scattering signal in the upper 20–150 m was significantly greater than the background values (~200 m). In >85% of the samples collected, the scattering signal recorded between 20 and 150 m was significantly greater (n = 56 night profiles over 42 cruises; one-tailed t test p < 0.05) than deep water blanks for rings 12–32 (corresponding to equivalent spherical diameters (ESD) of 1.25–40 μm). The percent of retrievals significantly greater than deep water blanks was lowest (~20%) at smaller ring detectors (e.g., at rings 1–6 which correspond to ESD > 100 μm, data not shown). This detection limit is similar to that cited by Barone et al. [2015]. In this same region, using identical deployment procedures and processing routines for data collected in July–September 2013, they found that the signal measured for rings 12–32 was significantly higher than the deep water background signal (202–204 m) for >90% of values in the upper 20–150 m. Thus it is this size (1.25–40 μm) and depth range (20–150 m) where the LISST provides the most reliable signal in our region of study; particles greater than ~110 μm are not generally above detection limits.

Sequoia Scientific provides two kernel matrices for the scattering inversion. The standard kernel is calculated using Mie theory as a composite of several indices of refraction, and is designed to produce accurate inversion results over a broad range of particle types (ranging from organic to inorganic). Sequoia also provides an empirically derived “randomly shaped” matrix that was developed from light scattering measurements of sieved mineral grains [Agrawal et al., 2008]. The random shape matrix was developed to address artifacts that were observed in data collected on populations of natural particles, where a spurious increase in the small size bins was observed [Agrawal et al., 2008; Mikkelsen and Pejrup, 2001]. We tested both of the inversion kernels provided by Sequoia Scientific as well as a kernel matrix recommended for phytoplankton assemblages by Andrews et al. [2010], where the index of refraction (n) is assumed to be a constant value of 1.17. After the inversion, the data are corrected for the difference in laser power between the factory calibration and the in situ data and an instrument-specific correction factor is applied to obtain the calibrated particle volume concentration, in units of volume particles per volume of water. Average particle volume distributions were then computed at 2 m intervals for each downcast. A 2 m bin size encompasses on average 240 LISST scans and corresponds to a scanned volume of 0.4 L. All data processing was done in MATLAB using manufacturer-supplied code for the background scattering correction (getscat.m), the scattering inversion (invert.p using 15 iterations), and the laser power and concentration corrections (vdcorr.m).

The shape of the PSD is commonly described by the exponent (also known as the PSD slope or Junge slope) of a power-law fit to the relationship between particle number concentration and diameter, the so-called power-law model. The general form of this model is

\[ N(D) = N_0 \left( \frac{D}{D_0} \right)^{-\xi} \]  

(5)

where \( N(D) \) is the particle concentration at a given diameter (units of particles L⁻¹ μm⁻¹), \( N_0 \) is the differential particle concentration at \( D_0 \), \( D_0 \) is a reference diameter, and \( \xi \) is the exponent of the distribution. In this work, the PSD slope was estimated from the change to particle concentration normalized by the width of
For purposes of comparison to other studies [e.g., Buonassisi and Dierssen, 2010; Kostadinov et al., 2012; Reynolds et al., 2010], we have applied a linear least squares fit to log-transformed data; the size range was restricted to 2.63–109.24 μm (median diameters); the reference diameter, \( D_0 \) was 2.63 μm, the smallest size bin used in our fit. The rationale for restriction of the size range is to improve the fit of the power law parameterization, and avoid the largest and smallest bins which are subject to the greatest instrument and/or data processing artifacts, for example, scattering at density gradients [Buonassisi and Dierssen, 2010; Kostadinov et al., 2012; Styles, 2006]. As Barone et al. [2015] discuss, the PSD slope should be interpreted with caution given that (1) there is a somewhat arbitrary need to exclude outer and inner rings which are most sensitive to errors and contamination from the PSD fit, and (2) the common means for fitting these spectra are biased [Clauset et al., 2009]; for example, nonlinear fitting procedures are statistically preferable to the near universal utilization of linear regression of log-normalized data that pervade the LISST-specific literature.

While the fit parameters are shown, there is no a priori reason to assume that the PSD at Station ALOHA follows a power law. Accordingly, we examined the normalized bias of the PSD to assess differences in the shape of the PSD produced by the inversion kernel. The normalized bias (NB) is simply a description of how the PSD deviates from a modeled (or predicted) PSD, in this case the power-fit to the PSD data. NB (%) is calculated as follows:

\[
NB(D_i) = \frac{P(D_i) - O(D_i)}{O(D_i)} \times 100
\]

where \( P(D_i) \) is the predicted particle concentration at a given diameter \( D_i \) and \( O(D_i) \) is the observed particle concentration at the same diameter. We estimated the median NB at each 2 m depth bin with all available profiles in the data set. Figure 1 shows the particulate spectra and the NB for all tested inversions. The shape of the PSD generated using the random inversion shows larger deviations from a power law than the spherical inversion, particularly for small particles (< 3 μm). The inversion matrix suggested by Andrews et al. [2010] generated significantly lower particle concentrations, particularly above 40 μm.

As a means of describing changes in the PSD over depth and time, we also used a weighted average particle size, \( D_{\text{avg}} \) (units of μm), shown in Slade et al. [2011] as

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**Figure 1.** The mean PSD derived from a spherical inversion (a), random inversion (c) and \( n = 1.17 \) (e). Data are binned to 2 m depth bins, and each sample represents the median of all available LISST profiles at that depth. The black line indicates the mean power law fit. The normalized bias of a power law fit is shown for the spherical (b), random (d) and \( n = 1.17 \) (f) inversions as a function of size (x-axis) and depth (color scale).
where, $A(D_i)$ is the areal size distribution in suspended cross-sectional area per volume (m$^2$ m$^{-3}$) for each LISST size class $i$, with mean diameter $D_i$. The areal size distribution is calculated from the volume size...
distribution \( \langle D \rangle \) by assuming spherical geometry: \( A(D) = \frac{3}{2} V(D) D^{1.5} \). The use of areal size distribution in the calculation of \( D_{\text{avg}} \) makes intuitive sense because cross-sectional area is what the LISST actually detects. It is important to note that, unlike the PSD slope, the average particle size estimate makes no assumptions about the shape of the PSD.

To simplify the description of size spectra, we have adopted a version of the particle classification scheme proposed by Sieburth et al. [1978] for the separation of planktonic organisms in picoplankton (0.2–2 \( \mu \text{m} \)), nanoplankton (2–20 \( \mu \text{m} \)), and microplankton (20–200 \( \mu \text{m} \)). We calculate the binned particle volume in the effective size ranges of 1.25–2.05 \( \mu \text{m} \) (the smallest size bin the LISST is capable of resolving), 2.05–20.86 \( \mu \text{m} \), and 20.86–109.25 \( \mu \text{m} \) (the upper limit is restricted to avoid scattering artifacts introduced by density gradients) [Styles, 2006] and to reflect our finding that particles >\( > 100 \mu \text{m} \) are often not detectable. These classes are referred to as 1.25–2 \( \mu \text{m} \), 2–20 \( \mu \text{m} \), and 20–100 \( \mu \text{m} \), respectively. Clearly, this operational definition excludes a large fraction of picoplankton (<1.25 \( \mu \text{m} \)) and larger aggregates or organisms that fall out of this size range. Nonetheless, trends in a component of picoeukaryote populations (nominally 0.2–3.0 \( \mu \text{m} \) in diameter) [Worden and Not, 2008], diatoms (~2–200 \( \mu \text{m} \)) [Hasle, 1996], nanophytoplankton such as prymnesiophytes \( >5 \mu \text{m} \) [Vernick, 1982], and detrital material (1.25–100 \( \mu \text{m} \)) should be captured by this approach.

Finally, to transform particle abundance \( N(D) \) into particulate carbon concentrations, we used the relationship derived by Menden-Deuer and Lessard [2000] for non-diatom protistan plankton. This transformation was developed using cultures of cyanobacteria, dinoflagellates, and prymnesiophytes and is expressed as follows:

\[
C(D) = 0.216 \times V(D)^{0.935} \times N(D) \times 8.3 \times 10^{-8}
\]  

(8)

where \( C \) is carbon concentration (\( \mu \text{mol C L}^{-1} \)), \( V \) is the volume in \( \mu \text{m}^3 \) of a spherical particle at a given median ring diameter \( D \), \( N(D) \) is the particle abundance in cells \( \text{L}^{-1} \), here not normalized to the bin-width and the scaling factor \( (8.3 \times 10^{-8}) \) converts from pg C to \( \mu \text{mol C} \). Just as we have done for particle volume, we have binned carbon content into three classes: 1.25–2 \( \mu \text{m} \), 2–20 \( \mu \text{m} \), and 20–100 \( \mu \text{m} \), respectively. We define the total particle carbon (TPC in \( \mu \text{mol C L}^{-1} \)) as the
The sum of the carbon concentration in the 1.25–109.25 \( \mu \text{m} \) range. LISST-derived TPC does not include contributions from heterotrophic and photosynthetic bacteria less than \( \sim 1.25 \ \mu \text{m} \) nor is it impacted by dissolved organic carbon absorption onto glass fiber filters measurements [see Moran et al., 1999]. Nevertheless, Barone

| Inversion | Particle Abundance (# L\(^{-1}\)) | TPV (\(\mu\text{L} \ \text{L}^{-1}\)) | TPC (\(\mu\text{mol C L}^{-1}\)) | Eukaryotic Phytoplankton (# L\(^{-1}\)) | Particulate Carbon (\(\mu\text{mol C L}^{-1}\)) |
|-----------|-------------------------------|-----------------------|-----------------------------|---------------------------------|-----------------------------|
| 25m       |                               |                       |                             |                                 |                             |
| Spherical | \(1.3 \times 10^6 \pm 3.6 \times 10^3\) | 0.065 ± 0.033         | 0.82 ± 0.37                 |                                 |                             |
| Random    | \(2.0 \times 10^6 \pm 7.9 \times 10^3\) | 0.052 ± 0.035         | 0.59 ± 0.44                 |                                 |                             |
| n = 1.17  | \(1.9 \times 10^6 \pm 3.2 \times 10^3\) | 1.8 \times 10^{-4} ± 1.3 \times 10^{-4} | 0.002 ± 0.001               |                                 |                             |

| 125m       |                               |                       |                             |                                 |                             |
| Spherical | \(1.3 \times 10^6 \pm 6.3 \times 10^5\) | 0.032 ± 0.017         | 0.41 ± 0.18                 | \(9.1 \times 10^5 \pm 3.1 \times 10^5\) | 1.3 ± 0.3                   |
| Random    | \(1.7 \times 10^5 \pm 8.6 \times 10^4\) | 0.024 ± 0.016         | 0.27 ± 0.16                 | \(8.6 \times 10^{-5} \pm 5.3 \times 10^{-5}\) | 0.001 ± 0.001               |
| n = 1.17  | \(1.4 \times 10^5 \pm 0.6 \times 10^3\) | 6.6 \times 10^{-2} ± 5.3 \times 10^{-5} | 0.001 ± 0.001               |                                 |                             |

*For reference, we report the climatological average ± standard deviation of the abundance of eukaryotic phytoplankton, which is a fraction of the total particle abundance and particulate carbon for September 2009 to December 2013, the period of data available which overlap LISST deployments.

Figure 4. Relationship between NPP measured between 25–125 m, (mg m\(^{-2}\) d\(^{-1}\)) and (a) the average particle diameter (\(D_{\text{avg}}\)) and (b) the absolute value of exponent of a power law fit (\(n\)) of the particle size distribution. Colors correspond to sampling depth. The mean depth profile of \(D_{\text{avg}}\) and \(n\) for winter (November–January, blue), spring (February–May, green), summer (June–August, red), and fall (September–October, orange) are shown in Figures 4c and 4d, respectively. While shown, values at less than 20 m are not considered due to the potential influence of bubbles (<20 m). In the 20–175 m strata, the mean ± the standard deviation for \(n\) was 4.2 ± 0.7, whereas the \(D_{\text{avg}}\) was 13.5 ± 5.7 \(\mu\text{m}\) (n = 63 casts binned to 2 m resolution for a period spanning September 2009 to April 2014). Larger values of \(n\) indicate greater contributions by small particles; smaller values of \(n\) indicate greater contributions by large particles.
et al. [2015] report that the sum of flow cytometrically derived bacterial carbon and LISST-TPC accounted for 76\% ± 9\% of measured PC between 25 and 75 m and 51\% ± 14\% at 125 m at Station ALOHA. For this reason, we consider LISST TPC to be a reasonable approximation for living and detrital particles between 1.25 and 100 \mu m.

3. Results and Discussion

3.1. Diagnostic Pigments and Productivity

We have applied equations (1)–(4) to HPLC data collected in the upper 125 m of the water column at Station ALOHA and analyzed results from two ecologically relevant depth bins: (1) the upper 45 m which generally encompasses the surface mixed layer and is considered light-saturated (e.g., light levels are above half-saturation irradiances (Eo) for photosynthesis and NPP is largely independent of photosynthetically active radiation at 14C incubation depths (PARz), see Li et al. [2011]); and (2) 100–125 m encompassing the two standard sampling depths nearest the deep chlorophyll maxima which ranged between ~100 and 140 m over the study period. Notably, this deeper bin is light-limited (PARz < Eo) and NPP should be a linear function of light levels [Letelier et al., 2004; Li et al., 2011]. The seasonal cycle of the relative proportions of pico, nano, and microphytoplankton in these depth bins are shown in Figure 2. Picophytoplankton consistently account for greater than 50\% of the total chlorophyll with maxima in the summer-fall months and the largest observed seasonal amplitude (6–7\%, Figure 2a) whereas microphytoplankton contributions peak in spring to summer months and show a more damped seasonal amplitude (2–3\%, Figure 2c). These trends are similar in surface and deep strata. Nanophytoplankton seasonal cycles show opposing trends when these depth strata are compared; maxima occur in spring in the surface and fall at depth with an amplitude of ~5\% in both strata (Figure 2b). Essentially, the euphotic zone of Station ALOHA is a regime dominated by picophytoplankton, with changes in the relative abundance of larger cells (or at least pigments diagnostic of these cells) being more subtle.

Given that the inherent maximal growth rates of diatoms [Geider et al., 1986] (which are presumed to compose the pigment based microphytoplankton pool) are generally thought to exceed those of the predominant picophytoplankton in this system (Prochlorococcus and Synechococcus) [Chisholm, 1992], we might expect enhanced NPP when this class comprises a larger fraction of the standing stock. We do not however find any relationships between absolute rates of NPP and HPLC size fractions (data not shown). To reduce the impact of the seasonal cycle of solar irradiance on pigment concentrations, particularly in the deeper depth bin, we have also examined the relative change in HPLC size fractions and NPP by subtracting the climatological monthly mean from all data (Figure 2). The variability for any one class is on the order of ±10\%, with larger anomalies in pico and nanophytoplankton fractions at depth than in the surface (Figure 2). There are no statistically significant relationships between HPLC-based particle size classes and primary productivity or anomalies thereof (two-tailed t tests, p ≥ 0.05). The largest anomalies in productivity (> 5 mg C m^-3 d^-1 in surface and > 2 mg C m^-3 d^-1 at depth) are in fact associated with “normal” HPLC-based PFTs (Figure 2); however, episodic increases in microphytoplankton are apparent in spring and summer months in the upper 45 m.

To the extent that these pigment ratios represent changes in taxonomic groups rather than physiology, our findings suggest the following: (1) there are clear and stable seasonal cycles in the relative contributions of pico, nano, and microphytoplankton to total chlorophyll, and (2) increases in microplankton are episodically

| Depth (m) | \( D_{avg} (\mu m) \) median | \( D_{avg} (\mu m) \) SD | PSD slope, \( \xi \) median | PSD slope, \( \xi \) SD | R^2 of PSD Fit |
|----------|-------------------------------|--------------------------|-----------------------------|--------------------------|-------------------|
| 25       | 10.69                         | 5.17                     | 4.48                        | 0.91                     | 0.99              |
| 45       | 9.67                          | 6.67                     | 4.49                        | 0.70                     | 0.99              |
| 75       | 9.75                          | 5.14                     | 4.39                        | 0.65                     | 0.99              |
| 100      | 10.52                         | 12.96                    | 4.36                        | 0.82                     | 0.99              |
| 125      | 10.35                         | 7.66                     | 4.32                        | 0.67                     | 0.98              |
| 150      | 11.56                         | 16.71                    | 4.26                        | 1.44                     | 0.98              |
| 175      | 20.21                         | 26.67                    | 3.86                        | 2.04                     | 0.97              |

aData used are mean and SD within ± 2 m of the sampling depths shown.

Note: Table 3. The Median and Standard Deviation (SD) of Two Descriptors of the Particle Size Distribution at Station ALOHA Standard Sampling Depths (Between 20 and 175 m), Average Particle Size and PSD Slope^a

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[Li et al., 2011]
associated with but not necessary for elevated primary productivity in the upper water column at Station ALOHA during the spring-summer time frame. These analyses suggest that shifts in community structure, from small to large or vice versa, do not drive variability in NPP in this regime. If this is the case, then retrieval of phytoplankton functional types from present-day ocean color remote sensing [e.g., Silico-Calzada et al., 2008; Uitz et al., 2009] may be of limited utility in oligotrophic regimes.

A final point about the relationship between pigments and productivity in the surface ocean of the NPSG: photoadaptation is in fact readily apparent in this region. In the upper euphotic zone, absolute concentrations of chlorophyll peak in low-light winter months and decline in high-light summer months as cells adjust to ambient light and nutrient conditions [Letelier et al., 1993; Westberry et al., 2008; Winn et al., 1995]. This seasonality (Figure 3) opposes the cycle of primary productivity which is highest in summer and lowest in winter [Karl et al., 2012; Letelier et al., 1996]. Accordingly there is no relationship between either total chlorophyll a or weighted diagnostic pigments (equation (4)) and NPP in the light-replete upper 45 m at Station ALOHA. Alternately, at the base of the euphotic zone where productivity is tightly regulated by light [Letelier et al., 2004; Li et al., 2011], the monthly climatology of HPLC chlorophyll is significantly related to NPP (linear regression, $R^2 = 0.72$, $p < 0.05$, Figure 3). So while the surface ocean shows pigment and light independent variability in NPP, the base of the euphotic zone in this region exhibits coherent shifts in NPP and total chlorophyll but not specific PFT fractions. Again though, these conclusions are subject to light-dependent shifts in absolute and relative pigment concentrations. We next explore relationships between NPP, mean particle size, and carbon-based particle size classes which are not subject to the effects of photoadaptation.

3.2. Initial Characterization of the Particle Size Distribution at Station ALOHA

Given that laser diffraction is not routinely applied in oligotrophic settings, first we characterized and examined the shape and fit of the PSD for Station ALOHA. The approximation of particle size from forward scatter
relies on mathematical inversion of the observed scattering signal and certain assumptions about particle shape and refractive index. Without explicit knowledge of the bulk suspended particle properties, these assumptions cannot be verified [Graham et al., 2012]. Moreover, there are multiple scenarios that may lead to divergence of in situ particle size distributions from a power function (the Junge distribution); these include largely monospecific blooms, flocculation/aggregation, and instrument artifacts [Chami et al., 2006].

We have processed raw LISST data in MATLAB using three kernel matrices (spherical, random, and assuming a refractive index of 1.17) and found that the default composite sphere kernel provides a better fit to a Junge-type distribution (Figure 1) and more reasonable particle concentrations (Table 2). The n = 1.17 kernel matrix generated lower particle concentrations across all size bins, whereas the random inversion provided similar spectra to the spherical inversion, albeit the predicted concentrations of <5 μm particles was lower by an order of magnitude (Figure 1). We then compare the predicted particle concentrations from these three inversions to the mean concentration of flow cytometry derived abundances of eukaryotes at Station ALOHA at discrete depths (25 and 125 m, Table 2). Notably, these cells include pico- and nanoeukaryotes and span cell diameters of approximately 0.2–20.0 μm [Pasulka et al., 2013]. This comparison shows that both the random and n = 1.17 retrievals of particle abundance underestimate eukaryotes abundances and therefore must also underestimate total phytoplankton concentrations. Similarly, the estimated particulate carbon concentrations for both the random and n = 1.17 inversions are markedly lower than the spherical inversion. Last, we note that the shape of the n = 1.17 inversion spectra is unique and not easily explained by a power law (Figure 1). The random and spherical inversions do however produce highly similar temporal and vertical patterns (data not shown). For these reasons, we have elected to perform all calculations of particle concentrations, volume, and carbon using the spherical inversion; this selection also allows us to more readily compare our results to those of other studies.

Evaluation of power model fits to the log of the particle size distributions also justify using the spherical inversion. The mean slope of the PSD (Figure 4d, Table 3) is well within the range typically observed in oceanic waters, mostly from 3.8 to 4.5 [Reynolds et al., 2010; Stemmann et al., 2008] and R² of the model fit were above 0.97 (p < 0.01) on average (Table 2). Nevertheless, all tested kernels did show consistent divergence from the idealized power law at specific size bins (Figure 1). For the spherical inversion, a power law overestimates the volume of <2 μm particles and particles in the 20–80 μm size range and underestimates particle concentration in the 2–20 μm particle size range. The shape of the NB is similar for the random inversion; however, the magnitudes of deviations are larger. Given that these deviations tend to occur over consistent size ranges, they likely reflect real aspects of the bulk community size structure in our study area. Alternatively, as noted above, the shape of the n = 1.17 inversion showed the largest underestimations of small,
In sum, these analyses indicate that while a power law is a good statistical approximation of the PSD at Station ALOHA, systematic departures are apparent in distinct size classes. It must also be acknowledged that these slopes are influenced by small, out of range particles (<1.25 µm) which are known to contribute to the smallest LISST size bins [Andrews et al., 2011]. For this reason and as a consequence of known biases using log-log fits, these values should only be used for general comparisons of the PSD. In that regard, both D_{avg} (which makes no assumption about the shape of the PSD) and the exponent of the power law suggest a rather stable vertical distribution of mean particle size at Station ALOHA (Figure 4 and Table 3). Moderate increases in mean D_{avg} (Figure 4) are apparent in the upper 45 m in spring (February–May) and these increases extend to the lower euphotic zone in fall (September–October); these increases are not statistically significant due to high variability in D_{avg} (t test, p > 0.7).

3.3. Temporal Variability of Particle Size Classes and Mean Particle Size

Laser scatterometry allows for much higher vertical and temporal resolution sampling than can be offered by any discrete bottle-based measurements (e.g., HPLC) and permits an assessment of the temporal variability of the abundance and volume of particle size classes as well as the relative contributions of size classes to total particle volume. To investigate variability of particle size at Station ALOHA, we first examined changes in particle volume and carbon content over time and then assessed shifts in the average weighted particle diameter (D_{avg}). As described above, particle volume estimated via the LISST was grouped into size bins of roughly 1.25–2.0 µm, 2–20 µm, and 20–100 µm. Particles in the 1.25–2.0 µm size range generally exhibit maxima at depths of 100–140 m with only rare increases within the upper ocean (September 2009 and June 2011, Figures 5a and 6a). Conversely, 2–20 µm particles were maximal in the upper water column, typically within the mixed layer (Figures 5b and 6b) with a shape similar to the depth dependence of NPP (Figure 6c). No persistent depth profile was apparent for particles in the 20–100 µm size class (Figure 5c). There is also apparent seasonality to these particle distributions, specifically for the 1.25–2.0 µm size class which exhibits progressively deeper maximum depths when grouped by season (Figure 6a). From shallow to deep, the centers of the seasonal depth maxima (Figure 6a, calculated as the depth of the maximum value within 90–130m) are as follows: winter (November–January, 102 m, n = 16), spring (February–May, 104 m, n = 13), summer (June–August, 106 m, n = 15), and fall (September–October, 122 m, n = 12). Seasonal differences in 1.25–2.0 µm volume concentrations are less than 5 µm particles.
apparent in the surface mixed layer where only fall profiles differ from other seasons; volume concentrations are $< 1 \times 10^{-3}$ for fall and $\geq 1 \times 10^{-3}$ in all other seasons (Figure 6a). For the 2.0–20 $\mu$m size class, volume concentrations in the upper 45 m are lowest in winter, increase in spring and reach maxima in summer and fall (Figure 6b). At depths typical of the DCM (100–125 m), the lowest 2.0–20 $\mu$m volume concentrations are observed in winter and the highest in spring, with summer and fall values falling in between.

While not comparable to trends seen in HPLC-based size fractions, we find that LISST-based particle size classes do show moderate seasonality in surface waters, particularly for the 2.0–20 $\mu$m size class which increase from winter through summer/fall. Notably, within this size class, particles with an ESD of $\sim 5$ $\mu$m are the most significant contributors to particle volume and carbon content (Figure 7). We cannot determine the identity of these 5 $\mu$m particles; however, this size range is consistent with small eukaryotic nanoplanктон [DuRand et al., 2001]. Differences between HPLC and LISST-based seasonality may be a result of contributions of detrital material to LISST-based particle loads or a consequence of the impact of photoacclimation, for example, changes in pigment per cell, on HPLC PFTs, rather than cell concentrations.

We also observed shifts in the mean particle diameter (Figure 8). In the surface mixed layer (25 m), $D_{\text{avg}}$ ranged from $\sim 5$ to 20 $\mu$m with episodic increases in September 2009, 2012, and in spring and winter months of 2013 (Figure 8a). At the depth horizon of 125 m, typically at or near the chlorophyll maximum at this site, $D_{\text{avg}}$ was more variable, ranging from 5 to 30 $\mu$m with values regularly exceeding $\sim 15$ $\mu$m nearly every fall/winter. These metrics indicate a relatively stable particle size in surface waters at Station ALOHA punctuated by episodic increases in particle diameter with a more regular increase in particle size at depth (100–125 m) in fall/winter months.

Last, we have investigated the relationship between LISST-based size fractions and net primary production. We find no relationship between $D_{\text{avg}}$ or the PSD slope and NPP rates (Figures 4a and 4b). This indicates that changes in NPP are not impacted by the relative mean particle size. We can also ask whether we see

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Figure 8. Box and whisker plot of weighted mean particle diameter ($D_{\text{avg}}$) at the depth horizon of (a) 25 $\pm$ 2 m and (b) 125 $\pm$ 2 m. The climatological median $\pm$ one standard deviation was 10.3 $\pm$ 4.7 $\mu$m at 25 m and 10.9 $\pm$ 7.3 at 125 m. In both plots, the central mark of each box is the median, the edges of the boxes are the 25th and 75th percentile and whiskers extend to the 95th percentile. Outliers are plotted as crosses. Data are derived from 2 to 3 nighttime casts per cruise.
increased NPP as a result of alteration in the concentration of individual particle size classes. In this case, we find that the estimated volume and carbon content of particles within the 2–20 μm fraction shows a positive relationship to NPP (Figure 9b) only in the lower euphotic zone (discrete depths of 100 and 125 m; Model 2 linear regression, $R^2 = 0.25$ and 0.12, respectively, $p < 0.05$, $n = 23$). This relationship is even stronger when both depths are considered ($R^2 = 0.41$, slope = 0.37 d$^{-1}$, Figure 9b). At the other standard depths, there was no significant relationship between LISST-derived carbon content and primary productivity. To further investigate this finding, we have also examined the relationship between NPP and particulate carbon (PC) as determined by high-temperature combustion via the HOT program. Again, we find a linear relationship with PC and NPP at the base of the euphotic zone (Model 2 regression for 100 and 125 m, $R^2 = 0.41$, slope = 0.13 d$^{-1}$, Figure 9d). The vertical profiles of PC and NPP are similar in shape (see Figure 6c for NPP) at Station ALOHA, which would lead one to expect some relationship between PC (a proxy for biomass) and PP when the full euphotic zone is considered, albeit not necessarily at a single depth. Together these data indicate that increases in the 2–20 μm fraction may contribute to growth and biomass accumulation (carbon and chlorophyll-based, see Figures 3 and 9) at the base of the euphotic zone. In general, increases in 2–20 μm carbon and productivity at depth occur in summer-fall months, consistent with the timing of deepening or shoaling of the 1% light level [see Letelier et al. 2004]. In summary, our findings indicate that neither mean particle size nor shifts in the carbon content of any of the three particle size fractions considered in this study are associated with enhanced productivity in the surface ocean of the NPSG. However, in the lower euphotic zone seasonal increases in the proportion of the 2–20 μm size fraction appears to coincide with enhanced NPP, notably during periods when the 1% light level crosses the 100 m

![Figure 9.](image)

**Figure 9.** Relationship between the rate of NPP measured in 12 h in situ incubations at Station ALOHA between September 2009 and December 2012 and the carbon content of particles in the following size classes: (a) 1.25–2.0 μm, (b) 2.0–20.0 μm, (c) 20–100 μm as well as (d) particulate carbon collected by HOT program. LISST data are the mean of values measured within ±2m of the depth of NPP incubations. Colors correspond to depth. Total particulate carbon and particles in the 2–20 μm range show a positive relationship to measured productivity rates, however this relationship is only linearly significant at 100 and 125 m depth horizons (Type II linear regression shown in Figure 9b: slope = 0.36 d$^{-1}$, $R^2 = 0.41$; Type II linear regression shown in Figure 9d: slope = 0.13 d$^{-1}$, $R^2 = 0.43$).
depth horizon. This increase in the carbon content in 2–20 μm particles does not correspond to significant changes in $D_{\text{avg}}$ or the PSD slope, suggesting that shifts in one component of the community may not alter the mean character of the particulate pool.

Subtle changes in particle size classes may translate into significant export production without measurable increases in bulk chlorophyll or NPP. Because the annual mean of new production in this oligotrophic region is approximately 6% of NPP (Karl et al., 2012), even small increases in large cells (e.g., the 2–4% seasonal change in diatom markers indicated by HPLC, Figure 2 or the <5 mg m$^{-3}$ change in 2–20 μm carbon estimated by the LISST at the 100–125 m depth horizon, Figure 9) may drive pulses of organic matter export into the aphotic zone. Diatoms in particular are known to contribute disproportionately to the particulate flux observed in deep sediment traps (Scharek et al., 1999). Intriguingly, Karl et al. (2012) report annually recurring, elevated fluxes of particulate carbon and nitrogen in 2800–4000 m sediment traps between 15 July and 15 August. This pulse is not detected in the shallow traps (150 m) presumably due to differences in trapping efficiency between deep and shallow traps as well as the selective remineralization of particles with depth (Karl et al., 2012); hence we have not made an attempt to correlate the PSD to shallow export fluxes. If this increase in carbon content and NPP also leads to a proportional increase in carbon flux, then the base of the euphotic zone may serve as a source of organic material for the annual recurrent summer export pulse. This hypothesis would require knowledge of the PSD of sediment trap material, which we are currently lacking at Station ALOHA [albeit this relationship has been studied elsewhere, e.g., Durkin et al., 2015; Guidi et al., 2008]. Future studies pairing measurements of the PSD and size-fractionated productivity with sediment traps would be a step toward better understanding the relative contributions of various size classes to net community production and export in this region.

4. Conclusions

In order to better understand how phytoplankton community structure and particle size impact productivity in the open ocean, we have compared pigment-based PFTs and laser diffraction-based estimates of particle size classes to parallel measures of $^{14}$C-based primary productivity conducted by the HOT program. The conclusion we reach is similar to the assessment of Chisholm (1992): “the simplicity of the general relationships serve as a stable backdrop against which the exceptions can shine.” Particle size and productivity do not seem to covary in the upper euphotic zone at Station ALOHA; albeit there are exceptions at the base of the euphotic zone where light levels exert strong control on NPP.

In the upper 45 m, there are clear and stable seasonal cycles in HPLC pigments diagnostic for pico, nano, and microphytoplankton; however, these cycles are not significantly correlated to productivity, in either an absolute or relative sense (Figure 2). The chlorophyll content of the three HPLC-based size fractions and the monthly anomaly thereof show no statistically significant relationship to rates of primary production. Particle volume and carbon content for particles 1.25–110 μm similarly show no relationship to productivity in surface waters (Figure 9). So while there is temporal variation in PFTs and PSDs (Figures 2 and 6), and 8), these shifts do not help explain the variability in upper euphotic zone productivity in the NPSG. This lack of correlation may reflect methodological bias. For example, HPLC-based classifications only approximate changes in phytoplankton community structure; relative proportions of diagnostic pigments are impacted by photoacclimation as well as biomass changes. Laser diffraction only detects a fraction of the true PSD; particles smaller than ~1-2.5 μm are poorly detected although it is these very particles that are responsible for ~50% of productivity in this region (Williams et al., 2008) and particles >100 μm are rare enough to be at the detection limit of the LISST. Of course the $^{14}$C tracer method also comes with biases, e.g., bottle effects, $^{14}$C DOC excretion and re-uptake and dark $^{14}$C uptake (Peterson, 1980). These biases may obscure relationships between particle size and productivity. Alternately, a more resonant hypothesis is that the variability in productivity in the upper 45 m of Station ALOHA is driven by size-independent shifts in phytoplankton physiology. A number of studies have noted the role of short-lived upwelling and mixing events as well as the passage of mesoscale eddies that result in nutrient injections to surface waters that can fuel net growth (Calil et al., 2011; Letelier et al., 2000). Transient nutrient injections can potentially lead to changes in the efficiency of light absorption, or the quantum yield of photosynthesis without affecting community structure and hence particle size (Finkel et al., 2004; Geider et al., 1986). So while changes in particle size may reflect shifts in community structure they do not necessarily imply shifts in net productivity in the
stratified surface waters of the NPSG. Further investigation of the variability of chlorophyll-specific absorption or the quantum yield of photosynthesis in this system is warranted to constrain the variability of PP rates in the surface mixed layer at Station ALOHA.

At the base of the euphotic zone, we find a significant relationship between chlorophyll \( a \) (Figure 3), and carbon content (which is a function of particle volume, equation (7)) in particles with an equivalent spherical diameter between 2 and 20 \( \mu m \) and parallel measures of PP at depths of 100 and 125 m (Figure 9b). This increase does not correspond to an increase in mean descriptors of the PSD: \( D_{50} \) or the PSD slope (Figure 4). So it seems that while one fraction of the particulate spectrum increases, compensatory changes in other fractions preserve the mean character of the PSD. We believe this increase in 2–20 \( \mu m \) particulate matter reflects a real increase in phytoplankton that can partially explain the increases in summer-fall NPP at these depths. The 1% light level at Station ALOHA varies seasonally by \( \sim 30 m \) [Letelier et al., 2004] from \( \sim 120 m \) depth during summer to \( \sim 90 m \) depth during winter. NPP rates at the deepest standard sampling depths measured by the HOT program (100 and 125 m) closely follow light levels; rates are highest in summer when light penetrates further in the water column allowing summer deepening of the nitracline [Letelier et al., 2004]. During these summer periods, we observe an increase in the volume and carbon content of relatively large cells (2–20 \( \mu m \)); particles of \( \sim 5 \mu m \) diameter drive this signal. We hypothesize that the deepening of isolumes and the subsequent utilization of nitrate allows for increase in the abundance of eukaryotes or small diatoms that would be detected in this size range. Notably, while the HPLC pigment-based approach does not show this same relationship between discrete PFTs and NPP, the seasonal cycle of total chlorophyll at 100–125 is positively related to NPP. In summary, while we cannot rule out factors such as an increase in detrital particles or an accumulation of sinking particles at depth in summer, the seasonal deepening of isolumes and increased nitrate availability appear to fuel net growth and biomass accumulation at the base of the euphotic zone. This study in effect reveals a water column partitioned into a light-independent surface mixed layer where changes in phytoplankton physiology drive variability in NPP and a light-dependent region at the base of the euphotic zone where 2–20 \( \mu m \) particles may contribute to observed increases in phytoplankton biomass that drive enhanced NPP.

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