1. INTRODUCTION

Phytoplankton photosynthesis in the ocean accounts for approximately half of the global carbon fixation on Earth (Field et al. 1998). Unlike terrestrial plants, however, phytoplankton are phylogenetically diverse, comprising 3 basic evolutionary lineages.
(Delwiche 1999). The first lineage contains all prokaryotic oxygenic organisms, cyanobacteria, that presumably evolved more than 2.8 billion yr ago and played a fundamental role in forming oxygen, carbon, and nitrogen cycles on the planet. All other oxygen-evolving organisms are eukaryotic and contain 13 phyla with more than 20,000 species (Falkowski et al. 2004a). One group of eukaryotes uses chlorophyll b as a secondary photosynthetic pigment and forms the ‘green lineage’ from which all terrestrial plants descended. The ‘green lineage’ played a key role in the ocean from ca. 700 million until 250 million yr ago (Falkowski et al. 2004a). Since that time, another group of eukaryotes, called the ‘red lineage,’ radiated rapidly and became ecologically dominant in the modern ocean (Falkowski et al. 2004a). The ‘red lineage’ of phytoplankton comprises 12 phyla, including diatoms, dinoflagellates, and haptophytes, which are the most ecologically prominent (Falkowski & Raven 2007). These 3 ‘red’ taxonomic groups use chlorophyll c and a series of taxon-specific carotenoids as secondary pigments. Species within the ‘red’ lineage can form massive phytoplankton blooms in coastal, sub-polar, and polar regions of the modern ocean and are responsible for most carbon fixation and sequestration into the ocean interior.

In natural aquatic ecosystems, phytoplankton community structure and physiology vary both horizontally and vertically on micro-, meso-, and regional spatial scales. Community structure is commonly analyzed using microscopic and/or pigment analyses (Claustre 1994, Mackey et al. 1996, Bidigare et al. 2005, Uitz et al. 2006, 2008, Sathyendranath et al. 2009) or flow cytometry (Chisholm et al. 1988, Olson et al. 1991). While conventional flow cytometers are primarily designed for and largely limited to the analysis of pico- and nano-plankton (Chisholm et al. 1988, Li 1997, Olson et al. 2003), imaging-in-flow systems target larger cells (Sieracki et al. 1998, Davis et al. 2004, 2005, Olson & Sosik 2007, Sosik & Olson 2007). While all of these approaches have the advantage of precision and accuracy, they are invariably time consuming and labor intensive. Sample throughput becomes a limiting factor to determining patterns of distributions of photosynthetic organisms and rates.

Major functional groups of phytoplankton primarily differ and are characterized by their pigment composition and, as a result, by their spectral properties. Following pioneering works (Yentsch & Yentsch 1979, Yentsch & Phinney 1984), spectral-resolved measurements of fluorescence have been proposed and explored as a potential tool for rapid monitoring of variations in community structure (Oldham & Warner 1987, Seppälä & Balode 1997, Babin et al. 2005, MacIntyre et al. 2010, Richardson et al. 2010). Because spectral analysis requires very high signal-to-noise ratios and precision, the previously developed instruments have a lower limit of sensitivity of only ~0.5 to 1.0 mg chlorophyll m⁻³ that precludes their applications in the open ocean. Also, although time series of flow-cytometric sampling potentially allow one to deduce growth rates (Sosik et al. 2003), none of the above techniques provides information about phytoplankton physiology or photosynthetic rates.

Variable fluorescence techniques, such as fast repetition rate (FRR) fluorometry (Kolber et al. 1998) and fluorescence induction and relaxation (FIRE) technique (Gorbunov & Falkowski 2005) have been developed for in situ measurements of phytoplankton biomass and photophysiology in the ocean. These techniques are based on recording, with microsecond time resolution, fluorescent transients induced by a precisely controlled sequence of excitation flashes of light. The induced ‘variable fluorescence’ signals are driven by variations in the physiological status of the photosynthetic apparatus and provide a comprehensive suite of photosynthetic properties of light harvesting, photochemistry in Photosystem II (PSII), and electron transport down to carbon fixation. This analysis provides the basis for stress diagnostic protocols (Falkowski et al. 2004b). The use of FRR and FIRE instruments has provided unprecedented information about the functioning of phytoplankton communities and the impact of environmental factors controlling primary productivity in the ocean (e.g. Behrenfeld et al. 1996, Coale et al. 2004, Falkowski et al. 2004b, Lin et al. 2016) and is becoming increasingly important for most biological oceanographic programs.

In an effort to expand capabilities of variable fluorescence for taxonomic analysis, we examined the combination of variable fluorescence and spectral analyses that offers a potential for simultaneous determination of biomass, photophysiology, and taxonomy in natural aquatic ecosystems.

In this paper, we describe a novel ultra-sensitive multispectral FIRE instrument (a mini-FIRE) and examine analytical capabilities of this technique for rapid determination of phytoplankton functional groups, together with physiological characteristics and photosynthetic rates. For this goal, we examined the spectral dependences of both
fluorescence intensities ($F_m$) and functional absorption cross-sections of PSII ($\sigma_{PSII}$), to reconstruct the basic spectra of each major phylogenetic group of eukaryotes. Model experiments were performed with a set of 25 cultures in a wide range of environmental conditions (illumination and nutrient limitation). Using this framework, we developed and tested algorithms for taxonomic identification based on the spectral properties of $\sigma_{PSII}(\lambda_{ex})$, where $\lambda_{ex}$ is the wavelength of excitation light. This approach was compared to conventional analysis of fluorescence excitation spectra. Our results revealed that the spectral properties of functional cross-sections ($\sigma_{PSII}$) are highly conserved for each functional group and can serve as a better taxonomic indicator than classical fluorescence excitation spectra.

2. MATERIALS AND METHODS

2.1. Mini-FIRe instrument and measurement protocols

Variable fluorescence and photosynthetic characteristics were measured with a new miniaturized FIRe system (mini-FIRe) developed and manufactured by M. Gorbunov (Fig. 1A). This instrument is conceptually similar to previous FRR and FIRe systems (Gorbunov & Falkowski 2005), but exhibits ca. 20-fold higher sensitivity and signal-to-noise ratio, compared to its predecessors. This extreme sensitivity is crucial for sampling in oligotrophic regions, which constitute 30% of the world oceans (Lin et al. 2016). The mini-FIRe instrument also employs excitation at multiple wavelengths ($\lambda_{ex}$) that allows for selective excitation of different functional groups of phytoplankton, as well as for spectrally resolved measurements of functional absorption cross-sections of PSII, $\sigma_{PSII}(\lambda_{ex})$. The instrument used in this study employed 6 excitation channels (435, 450, 470, 500, 530, and 590 nm, with 20 nm bandwidth for each channel). The peak excitation intensity in the sounding volume, i.e. the volume from which the signal is recorded, ranges from ca. 20 000 to 25 000 μmol quanta m$^{-2}$ s$^{-1}$ for blue excitation channels (435, 450, 470, 500 nm) and 40 000 to 60 000 μmol quanta m$^{-2}$ s$^{-1}$ for green and orange excitation. These
higher excitation intensities from the green and orange light-emitting diodes (LEDs, Luxeon Rebel) are achieved by applying supra-optimal electric current (ca. 5x nominal values) in short pulses. The excitation intensities for all spectral channels are optimally adjusted to ensure closure of PSII reaction centers and saturation of fluorescence within ca. 100 μs (i.e. a single photosynthetic turnover) in each excitation channel (Fig. 1B), which is crucially important for accurate retrievals of functional cross-sections and quantum yields (Gorbunov et al. 1999).

The fluorescence signal is collected from a central, 8 mm diameter, portion of the sample volume, isolated by a red band pass interference filter (680 nm, with 20 nm bandwidth), and detected by an avalanche photodiode (SD630-70-72-500, Advanced Photonics). The voltage on the detector is automatically adjusted depending on ambient temperature to ensure a constant gain over a wide temperature range (0 to 30°C). A small portion of the excitation light is recorded by a PIN photodiode as a reference signal. By using an avalanche photodiode with high quantum efficiency (ca. 80%) in the red spectral region and integrating the fluorescence signal in the analog mode, the instrument operates with an extremely high signal-to-noise ratio. The mini-FIRe instrument can accurately measure fluorescence signals in samples as low as 0.01 mg chlorophyll m⁻³, effectively the lowest concentration of chlorophyll recorded anywhere in the upper ocean.

σ_{PSII} is a product of the optical absorption cross-section of PSII (i.e. the size of the PSII antennae) and the quantum yield of photochemistry in PSII (Falkowski et al. 2004b). σ_{PSII} is calculated from the rate of fluorescence rise during the single turnover flash (Fig. 1), as this rate is proportional to the product of σ_{PSII} and the photon flux density of excitation light (Gorbunov & Falkowski 2005). Retrievals of σ_{PSII} in absolute units (Å²) require accurate calibration of the average photon flux density in the sounding volume (Gorbunov & Falkowski 2005). Because the taxonomic analysis relies on quantifying small changes in the spectral shape, accurate calibration of all color channels is crucially important. Such calibration is conducted as part of the standard calibration procedure of the FIRe instrument.

2.2. Strains and culture conditions

The eukaryote algal cultures were grown at 18°C in F/2 medium in a 12:12 h light:dark cycle and white fluorescent light (100 μmol photons m⁻² s⁻¹). These cultures included diatoms (Thalassiosira pseudonana CCMP1335, Thalassiosira weissflogii CCMP1336, Chaetoceros muelleri CCMP1318, Navicula incerta UTEX 2044, Ditylum brightwellii CCMP358, Thalassiosira oceanica CCMP1005, Phaeodactylum tricornutum CCMP632, and Phaeodactylum tricornutum CCMP2559), dinoflagellates (Symbiodinium sp. CCMP-832 and CCMP2467, Amphidinium carterae CCMP-1314, and Prorocentrum micans CCMP1589), haptophytes (Emiliania huxleyi CCMP 373, CCMP 374, CCMP379, CCMP1516, and Gephyrocapsa oceanica AC396), and chlorophytes (Dunaliella tertiolecta CCMP1320, Chlorella vulgaris, Tetraselmis chuii, and Chlamydomonas reinhardtii). The cultures of cyanobacteria (Synechococcus sp. CCMP 1379, CCMP 2370, and PCC 7002) were grown at 25°C in F/2 media. The cyanobacterium Arthrospira maxima was grown at 25°C in BG-11 media. Because cyanobacteria are known to exhibit strong chromatic adaptation in their pigment content and spectral properties, we grew cyanobacteria cultures under white and blue (central wavelength 455 nm, half-bandwidth 30 nm) light from a custom-built LED lamp. The white light was intended to simulate conditions in the upper ocean layer, while the blue light simulated the irradiance regime in open-ocean deep blue waters.

2.3. Nutrient stress experiments

Nutrient stress experiments were conducted on batch cultures growing in low-nitrogen media. First, cultures were pre-acclimated (for 4−5 d) to low-nitrogen F/2 medium, with [NO₃] of 10 μM, which is ca. 90 times lower than in normal F/2. This pre-acclimation allowed the cells to maintain maximum growth rates even in this low-nitrogen medium, although the cellular chlorophyll a (chl a) content reduced several fold (a sign of acclimation to low-nitrogen conditions). We then monitored, in batch cultures, the slowly progressing nutrient stress, as the cells digested the remaining [NO₃] and the nitrogen concentration in the medium slowly reduced down to 0 within 48−72 h. The chl a concentration in the samples during these experiments did not exceed 5−10 mg chl m⁻³ to ensure the slow development of nitrogen stress.

2.4. Growth rate measurements

Instantaneous growth rates (μ) were determined from an increase in cell density (N, cells ml⁻¹) over a short (4 to 6 h) period of time (δt), using an exponential growth equation:
Cell densities were recorded using a Beckman Coulter Multisizer 3 cell counter.

2.5. Fluorescence spectroscopy

Fluorescence excitation spectra were measured on a Jasco FP-8500 spectrofluorometer, by scanning the excitation wavelengths over the visible region (from 400 to 660 nm) and recording chl a fluorescence emission at 690 nm (with 5 nm spectral bandwidth). In any conventional spectrofluorometer, the excitation intensities vary with wavelength, inevitably causing variations in the fraction of dynamically closed reaction centers and resulting in variations in the fluorescence quantum yield during the scan. As a consequence, the measured fluorescence excitation spectra may be severely distorted in photosynthetically active cells. To avoid this artifact, we treated all samples with 3-(3,4-dichlorophenyl)-1,1-dimethylurea (DCMU) and recorded fluorescence excitation spectra at the Fm level, with all reaction centers closed.

We should stress that, for the above reason, conducting measurements on DCMU-treated samples is the only way to correctly measure fluorescence excitation spectra in phytoplankton using conventional spectrofluorometers.

2.6. Field studies

Multi-spectral variable fluorescence data were collected during the annual Western Antarctic Peninsula Long Term Ecological Research (WAP LTER) cruise in the austral summer (January–February) of 2017 aboard the RV ‘Laurence M. Gould.’ The WAP LTER encompasses a grid spanning an area of ~900 × 200 km across the WAP shelf (Montes-Hugo et al. 2009). Samples were collected in Niskin bottles and kept at in situ temperature (0 to 1°C). During the day time, samples were acclimated at low light (ca. 5 μmol quanta m⁻² s⁻¹) for 30–60 min prior to measurements, to ensure recovery of variable fluorescence from non-photochemical quenching and photoinhibition. We should note that low-light (not dark) acclimation was crucial for recovery from photoinhibition, because this recovery is energetically driven by the ΔpH gradient across the cellular thylakoid membrane produced by low-light exposure. Phytoplankton community structure was analyzed using microscopic analysis or HPLC (Montes-Hugo et al. 2009).

2.7. Statistical analysis

Data analyses were performed using the Python programming language and the ‘scikit-learn’ library. The K-means clustering algorithm was employed for the analysis of the spectra. The number of clusters was set to 4 based on a priori information on the number of phytoplankton functional groups. It was expected that due to significant variations between the spectra for different taxonomic groups, the cluster analysis would divide the dataset into 4 subsets. The clustering procedure yielded the centers of these 4 clusters (centroids), which were further compared to the basic spectra obtained by averaging the experimental spectra for each group. Based on the proximity of spectral band shapes, the 4 clusters were attributed to diatoms, dinoflagellates, haptophytes, and green algae.

3. RESULTS

3.1. Spectral dependence of \( \sigma_{PSII}(\lambda) \) as a taxonomic index

The proposed approach for taxonomic analysis relies on using the spectral dependences of \( \sigma_{PSII}(\lambda_{ex}) \) and \( F_{m}(\lambda_{ex}) \). We hypothesized that each taxonomic group exhibits its own phylum-specific spectra and that spectral variations within each group are smaller than the differences between different groups. To test this hypothesis, we analyzed multi-color FIRe measurements of the dependences \( \sigma_{PSII}(\lambda_{ex}) \) and \( F_{m}(\lambda_{ex}) \) for a wide variety of eukaryote cultures.

Fig. 2 demonstrates the \( \sigma_{PSII}(\lambda_{ex}) \) spectral dependences obtained for 4 major taxonomic groups of eukaryotic phytoplankton, including diatoms (7 cultures, Fig. 2A), dinoflagellates (3 cultures, Fig. 2B), green algae (4 cultures, Fig. 2C), and haptophytes (4 cultures, Fig. 2D).

Although the absolute values of \( \sigma_{PSII} \) varied by a factor of ca. 2 within each taxonomic group (Fig. 2), the spectral shape of \( \sigma_{PSII}(\lambda_{ex}) \) was remarkably constrained within each group of eukaryotes. Fig. 3A shows the normalized \( \sigma_{PSII}(\lambda_{ex}) \) spectra averaged over each group. We call these averaged normalized spectra the ‘basic spectra’ for each group. For each pairwise comparison between basic spectra, there is a wavelength region where the divergence between these groups exceeds the dispersion, suggesting that these dependences can potentially be used to discriminate between different groups.
We further compared the spectral dependencies $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ of eukaryotes and prokaryotes. Fig. 4 shows $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ for 4 cyanobacterial cultures, in comparison with the average $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ spectra for 4 major phylogenetic groups of eukaryotes. The cross-sections recorded at a single wavelength, $\sigma_{\text{PSII}}(450\text{ nm})$, were extremely small in cyanobacteria (Fig. 4)—several-fold smaller than in eukaryotes, consistent with previous observations (Suggett et al. 2009, Gorbunov et al. 2011). Moreover, in contrast to eukaryotes, the spectral shape of $\sigma_{\text{PSII}}$ varied dramatically among cyanobacterial species (Fig. 4), reflecting variations in phycobilin content and composition.

Chromatic adaptation to blue light decreased by 20–30% $\sigma_{\text{PSII}}(590\text{ nm})$ relative to $\sigma_{\text{PSII}}$ in the blue-green spectral region in Synechococcus spp. CCMP 1379 and PCC 7002. However, chromatic adaptation had virtually no effect on the spectral shape $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ in Synechococcus spp. CCMP 2370.

### 3.2. Mathematical algorithm for using $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ as a taxonomic index

Based on the observed differences in the basic spectra of major phylogenetic groups of eukaryotes, we developed and tested the following mathematical algorithm for taxonomic identification.

First, we define the spectral slope $S_{ij}$ for the basic spectra as:

$$S_{ij} = (S(\lambda_i) - S(\lambda_j))/(\lambda_i - \lambda_j)$$

where $S$ is $\sigma_{\text{PSII}}$ or $F_m$ for the $\sigma_{\text{PSII}}$-based or $F_m$-based algorithm, respectively; $i, j = 435, 450, \ldots 590$ correspond to the 6 excitation wavelengths used in our instrument. Based on the $S_{ij}$ values, the following classification procedure is proposed (Fig. 3B):

First, green algae are identified by a reduced value of $S_{330:590}$, which is due to low absorption in the green spectral range (530 nm).
Second, diatoms are characterized by an enhanced value of $S_{450:470}$, which is due to a steeper slope in the 450−470 range originating from an interplay between an increased chlorophyll content and a spectral dip caused by carotenoids.

Third, dinoflagellates and haptophytes are discriminated by the $S_{470:530}$ value, whereby $S_{470:530}$ is smaller for haptophytes, as compared to dinoflagellates, due to differences in carotenoid content.

Application of this procedure to our dataset of $\sigma_{\text{PSII}}(\lambda)$ for 18 eukaryote species yielded 90% accuracy of this identification algorithm. This result clearly suggests that the spectral variability of $\sigma_{\text{PSII}}(\lambda)$ among different phylogenetic groups is sufficiently large to be used to functionally discriminate among the various phytoplankton taxa.

With the aim to achieve the maximum precision of the taxonomic identification algorithm, we used the entire dataset of spectral signatures for development and parameterization of this algorithm. At the same time, a common practice in validating such algorithms relies on splitting the data into 2 subsets ('training' and 'test') and using one of these subsets for development and another one for validation. For this reason, we conducted additional analyses by splitting our dataset into 2 independent 'training' and 'test' subsets. Our analysis revealed that by following this 'training-and-test' approach, the algorithm correctly identified 88% of the test data, which was very close to 90% correct identifications achieved for the above algorithm based on the entire dataset.

We further propose an alternative, more refined procedure for taxonomic identification, by introducing a ‘proximity index’ which characterizes proximity of a measured normalized spectrum $\sigma_{\text{PSII}}(\lambda)$ to the basic spectrum of a given phylogenetic group:

$$n_j = \sqrt{\sum_i \left( S_x(\lambda_i) - B_j(\lambda_i) \right)^2}$$

where $S_x$ is the analyzed spectrum and $B_j$ is the basic spectrum of the $j^{th}$ phylogenetic group. After calculating the discrepancies between the measured and all modeled basic spectra, the taxonomic identity of a sample is ascribed to the basic spectrum for which the $\Delta$ value is minimal.

Fig. 3. (A) Normalized $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ spectra (i.e. ‘basic’ spectra) of 4 different phylogenetic groups of eukaryotes (a.u.: arbitrary units; error bars are SD). (B) Schematic representation of the procedure used for phylogenetic group determination based on analysis of the $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ spectral shape. Here, $S_{ij}$ is the spectral slope obtained for the region of the $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ spectrum in the $\lambda_i: \lambda_j$ range (see Section 3.2 for details).

Fig. 4. Spectra $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ for selected cyanobacteria (gray symbols) and the averaged spectra $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ for 4 major phylogenetic groups of eukaryotes (open symbols).
Application of this procedure to our dataset of $\sigma_{\text{PSII}}(\lambda)$ also yielded 90% correct identifications. Hence, our analysis clearly suggests that the spectral shape of $\sigma_{\text{PSII}}$ can be used as a taxonomic index, even for major phyla of the ‘red’ lineage of phytoplankton that have similar spectral properties.

To further evaluate the new multi-color FIRe approach for taxonomic analysis, we conducted an independent cluster analysis of the collected database of spectral signatures. This analysis automatically yielded 4 principal groups of spectra that correctly corresponded to the 4 phylogenetic groups used in our experiments (diatoms, haptophytes, dinoflagellates, and chlorophytes). The 4 basic spectra retrieved with this cluster analysis (Fig. 5B) were insignificantly different from the 4 basic spectra recorded experimentally (Fig. 5A).

### 3.3. Effect of nutrient limitation on the spectral properties of $\sigma_{\text{PSII}}$

Phytoplankton photosynthesis in the global ocean is fundamentally limited by the availability of nutrients, such as nitrogen and iron (Lin et al. 2016). How does nutrient limitation affect the spectral properties of phytoplankton fluorescence, including $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$? How might variations in the nutrient status affect the applications of the spectral-resolved variable fluorescence for taxonomic analysis?

To address these questions, we examined the effect of nitrogen limitation on $\sigma_{\text{PSII}}(\lambda_{\text{ex}})$ in 2 model species, the diatom *Phaeodactylum tricornutum* and the chlorophyte *Dunaliella tertiolecta*. We quantified the extent of nitrogen limitation from the reduction in the quantum yield of photochemistry in PSII ($F_v/F_m$) and growth rates.

In *D. tertiolecta*, nitrogen limitation resulted in a characteristic decrease in growth rates, followed by a decrease in $F_v/F_m$ and an increase in $\sigma_{\text{PSII}}$ (Fig. 6). Under severe nitrogen limitation, when the instantaneous growth rates reduced by ~90%, $F_v/F_m$ decreased by ca. 50% and $\sigma_{\text{PSII}}$ increased by 40% (Fig. 6). These changes in $F_v/F_m$ and $\sigma_{\text{PSII}}$ are typical signatures of nutrient stress in phytoplankton (Kolber et al. 1988, Falkowski et al. 2004b). The decrease in $F_v/F_m$ indicates a reduction in the fraction of active PSII reaction centers. Thereby, the absorbed light energy can be transferred from inactive photosynthetic units to remaining active centers, thus increasing the effective absorption cross-section of these active centers. Also, a decrease in cellular pigment density under nutrient limitation leads to reduction in the package effect, further increasing the effective absorption cross-section.

Nitrogen limitation had no effect on the spectral shape of $\sigma_{\text{PSII}}(\lambda)$ (Fig. 6). For instance, changes in the normalized cross-sections, $\sigma_{\text{PSII}}(\lambda_{\text{ex}})/\sigma_{\text{PSII}}^{\text{max}}$ were not statistically significant ($p > 0.5$) for all FIRe wavelengths, except 435 nm. Nitrogen limitation resulted in a small (3%), but statistically significant ($p < 0.05$) decrease in the ratio $\sigma_{\text{PSII}}(435)/\sigma_{\text{PSII}}(450)$. The wavelength of 435 nm corresponds to the maximum of chl a absorption, and the relative decrease in $\sigma_{\text{PSII}}(435)$ is indicative of a reduction in the cellular chl a content (chlorosis), which is typically observed under nitrogen and iron limitation (Kolber et al. 1988, Greene et al. 1991).

In *P. tricornutum*, nitrogen limitation also resulted in a characteristic decrease in the growth rates, fol-
allowed by a decrease in $F_v/F_m$ and an increase in $\sigma_{PSII}$ (Fig. 6D–F). However, the increase in $\sigma_{PSII}$ was much smaller than in Dunaliella. Furthermore, under severe nitrogen limitation ($\mu < 0.2 \mu_{max}$ and $F_v/F_m < 0.3$), $\sigma_{PSII}$ decreased to the values typical of nutrient-replete cells (Fig. 6). The spectral shape of $\sigma_{PSII}(\lambda)$ was not affected by nitrogen limitation for all wavelengths ($p > 0.5$), except 435 nm (Fig. 6). In \textit{P. tricornutum}, the reduction in $\sigma_{PSII}(435)/\sigma_{PSII}(450)$ under severe nitrogen limitation was more pronounced than in Dunaliella (6 vs. 3%).

### 3.4. Comparing $\sigma_{PSII}(\lambda_{ex})$ and $F_m(\lambda_{ex})$ as taxonomic indexes

We further examined which of the spectral dependencies, $\sigma_{PSII}(\lambda_{ex})$ or $F_m(\lambda_{ex})$, is more accurate for taxonomic analysis. For this goal, we first reconstructed the basic spectra $F_m(\lambda)$ for the 4 major phylogenetic groups of eukaryotes (Fig. 7). These basic spectra were defined using the same database that we used for the $\sigma_{PSII}(\lambda_{ex})$ analysis above. We then applied the same mathematical procedure for $F_m(\lambda_{ex})$ analysis as we did for $\sigma_{PSII}(\lambda_{ex})$. Surprisingly, this $F_m$-based analysis yielded only 76% correct identifications, as compared to 90% for the $\sigma_{PSII}$-based analysis.

Next, we applied hyper-spectral fluorescence excitation spectra measured using a conventional fluorometer for taxonomic identification. Fig. 8 demonstrates the normalized averaged fluorescence excitation spectra for 4 major groups of eukaryotes.

Using the same procedure based on evaluation of proximity to basic spectra (Eq. 3), we obtained 89%
correct identifications. This number is very close to that obtained using the \( \sigma_{\text{PSII}}(\lambda_{\text{ex}}) \) algorithm above. At the same time, it clearly shows that the use of 6 excitation wavelengths was not sufficient for accurate identification using conventional fluorescence excitation spectra (e.g. \( F_{\text{mi}}(\lambda_{\text{ex}}) \)).

3.5. Field evaluation

Field validation of the proposed approach was performed using a dataset collected during the WAP LTER cruise off the WAP. The proximity indexes to the basic spectra of major functional groups were calculated for 150 samples. This analysis resulted in 95% of spectra being identified as diatoms, which was in good agreement with the microscopic examination of samples. Fig. 9 shows the average normalized \( \sigma_{\text{PSII}}(\lambda_{\text{ex}}) \) spectrum obtained for the Antarctic communities, as well as the basic spectrum for diatoms obtained from model experiments on diatoms (Figs. 2 & 3). Our taxonomic algorithm identified a small fraction (5%) of field spectra as being assigned to dinoflagellates. These false identifications were for deep-water (>60 m) samples with low chlorophyll content and thus low signal-to-noise ratio and with an elevated amount of fluorescent pigment degradation products.

4. DISCUSSION

Our research has revealed that, in spite of variability in pigment composition, \( \sigma_{\text{PSII}}(\lambda_{\text{ex}}) \) is remarkably constrained within each major taxonomic group of eukaryotic phytoplankton, including diatoms, haptophytes, dinoflagellates, and chlorophytes (Fig. 2A). Variability in \( \sigma_{\text{PSII}}(\lambda_{\text{ex}}) \) within each group was significantly smaller than the difference between individual groups, thus providing the basis for taxonomic identification algorithms. Although the big difference in spectral properties (i.e. color) between the ‘green’ and ‘red’ lineages of eukaryotic algae is well known, the discovered differences between the 3 major phyla of the ‘red’ lineage (diatoms, haptophytes, dinoflagellates) were unexpected. These ecologically dominant eukaryote groups exhibit similar spectral properties and remain extremely challenging for spectroscopic identification. Our results suggest that the multi-color variable fluorescence analysis offers a promising tool for the combined physiological and taxonomic analysis, including identification of 3 major phyla within the ‘red’ lineage of phytoplankton.

The absolute values of the cross-sections of PSII (e.g. \( \sigma_{\text{PSII}} \) measured at a single wavelength, i.e. 450 nm) vary 4- to 5-fold among species (Figs. 3 & 4; and see Suggett et al. 2009). In natural phytoplankton communities, changes in \( \sigma_{\text{PSII}} \) often reflect changes in the community structure and cell size distribution (Gorbunov et al. 1999). For instance, due to a stronger pigment packaging effect, larger cells tend to have a smaller effective \( \sigma_{\text{PSII}} \) compared to small cells (Gorbunov et al. 1999). Therefore, the magnitude of \( \sigma_{\text{PSII}} \) at one excitation wavelength may carry information about taxonomic identity (Suggett
et al. 2009). Our laboratory research supports this hypothesis, to a first approximation. Among eukaryotes, the highest $\sigma_{PSII}$ values were observed in haptophytes and diatoms, and the lowest values in green algae and dinoflagellates (Fig. 2). In cyanobacteria, $\sigma_{PSII}(450 \text{ nm})$ values were extremely small (Fig. 4): several-fold smaller than in eukaryotes, consistent with previous observations (Suggett et al. 2009, Gorbunov et al. 2011, Simis et al. 2012). Moreover, the spectral band shape of $\sigma_{PSII}$ for cyanobacteria is dramatically different from that of eukaryotes and varies dramatically among cyanobacteria species (Fig. 4; and see Simis et al. 2012). At the same time, big variations in $\sigma_{PSII}$ values were observed within each functional group of both cyanobacteria and eukaryotes (Figs. 3 & 4). Also, changes in environmental conditions, such as nutrient limitation, have a profound effect on $\sigma_{PSII}$ (Fig. 6; and see Kolber et al. 1988, Greene et al. 1991). These results clearly suggest that the absolute values of $\sigma_{PSII}$ at one excitation wavelength are not unambiguously indicative of taxonomic identity.

Many prokaryotic phytoplankton, such as cyanobacteria, and some eukaryotes (cryptophytes) can be identified using spectroscopic analysis, based on characteristic orange fluorescence emission from their unique accessory pigments, phycobilins (phycoerythrins, phycocyanin, and allophycocyanin) (Chisholm et al. 1988, Olson et al. 1991, Lantoine & Neveux 1997, Chekalyuk & Hafez 2008). This fluorescence emission is spectrally separated from chl $a$ fluorescence, although allophycocyanin fluorescence closely overlaps with that of chl $a$. All other eukaryotes only emit chl $a$ fluorescence (e.g. Falkowski & Raven 2007). This chl $a$ emission band is virtually identical in all organisms, which makes fluorescence emission spectroscopy unsuitable for taxonomic analysis of major groups of eukaryotes.

Based on the analysis of spectral properties and physiological characteristics, we have developed and tested an operational protocol for phytoplankton taxonomic identification, based on the proximity of measured spectra to basic (i.e. average) spectra of phylogenetic groups. We have tested this protocol on the collected database of 25 species. As much as 90% of the experimental spectra of eukaryotic cultures were correctly classified, clearly suggesting a high predictable capability of this approach. Additional evidence for the principal difference in the basic spectra between the taxonomic groups comes from our cluster analysis of the collected database that in its main components was very similar to the experimentally obtained basic spectra.

In addition to the spectral shape of $\sigma_{PSII}(\lambda_{ex})$, we also examined the spectral shape of chl $a$ fluorescence yields, $F_m(\lambda_{ex})$, as a potential taxonomic proxy. The shape of $F_m(\lambda_{ex})$ reflects the classical chl $a$ fluorescence excitation spectra (Lakowicz 2006). To a first approximation, the excitation spectra $F_m(\lambda_{ex})$ should be virtually identical to $\sigma_{PSII}(\lambda_{ex})$ because both variables are determined by the flux of excitons to the PSII core. Surprisingly, our research revealed that $\sigma_{PSII}(\lambda_{ex})$ is a better and more specific proxy for taxonomic analysis. For instance, the $\sigma_{PSII}$-based algorithm correctly identified 90% of experimental data, while the $F_m$-based algorithm only identified 77%.

The difference in the spectral shape between $\sigma_{PSII}(\lambda_{ex})$ and $F_m(\lambda_{ex})$ may be caused by the following: First, in intact cells, high intracellular density of photosynthetic pigments (i.e. the ‘pigment packaging effect,’ Geider & Osborne 1987, Berner et al. 1989) decreases the effective intensity of incident light within the cell, thus affecting both $\sigma_{PSII}$ and $F_m$. In addition, the pigment packaging leads to re-absorption of emitted fluorescence, further reducing the measured $F_m$, but not $\sigma_{PSII}$. Therefore, the pigment packaging has a stronger effect on $F_m$ than on $\sigma_{PSII}$. Variations in pigment packaging are most pronounced in large cells and introduce errors into the $F_m$-based algorithm. Second, measurements of fluorescence yields (both $F_o$ and $F_m$, but not $F_i$) are sensitive to artifacts of blank correction (Bibby et al. 2008). Because $\sigma_{PSII}$ is calculated from the variable fluorescence component (Falkowski et al. 2004b), $\sigma_{PSII}(\lambda_{ex})$ is not subject to errors in blank correction, at least in FIRe instruments.

Our approach has been implemented on the basis of a novel mini-FIRe system, which allows for simultaneous measurements of photophysiological parameters and spectrally-resolved fluorescence yields, $F_o(\lambda_{ex})$ and $F_m(\lambda_{ex})$, and $\sigma_{PSII}(\lambda_{ex})$. The physiological parameters are used for assessment of photosynthetic rates, including primary productivity (Kolber & Falkowski 1993, Hughes et al. 2018a, Ko et al. 2019), and the impact of environmental factors and stressors (Falkowski et al. 2004b, Gorbunov & Falkowski 2005), while spectrally-resolved fluorescence yields are for taxonomic analysis. The mini-FIRe system operates in 2 modes, including discrete sample analysis and a fully automatic on-line mode, using a flow-through cuvette with an anti-fouling cleaning system. These capabilities, in combination with extreme sensitivity, offer new prospects for automated monitoring of phytoplankton communities and primary production in the global ocean.
The combination of photophysiological parameters and taxonomic proxies from the multi-color FIRe measurements may have important implications for improving fluorescence-based methods to measure primary production. The starting point in these methods is the measurement of photosynthetic electron transport rates (Kolber & Falkowski 1993). The conversion of these rates to the rates of carbon fixation relies on the knowledge of electron requirements for carbon fixation (Kolber & Falkowski 1993, Lawrenz et al. 2013). Extensive laboratory and field studies have revealed that the electron requirements in natural phytoplankton communities are controlled by the extent of nutrient limitation and also may vary with taxonomy (Lawrenz et al. 2013, Zhu et al. 2017, Hughes et al. 2018b, Ko et al. 2019). The multi-color variable fluorescence technique might provide crucial information about these 2 controls of the electron requirement for carbon fixation. Although quantitative taxonomic analysis of natural communities using bulk fluorescence signals is still challenging and merits further research and development effort, our results clearly suggest that such analysis can be conducted, with high accuracy, on the level of a single species, for example using single-cell cytometric analysis. This result offers new opportunities for high-throughput cytometric analysis using multi-color fluorometry. Methodology and operational instrumentation for single-cell measurements of variable fluorescence have been developed (Gorbunov et al. 1999, Olson et al. 2000). Incorporation of multi-color protocols to such instrumentation would provide unprecedented capabilities for combined physiological and taxonomic analysis of natural phytoplankton communities in the ocean.

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