Aquatic laser fluorescence analyzer: field evaluation in the northern Gulf of Mexico

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Abstract: The new Aquatic Laser Fluorescence Analyzer (ALFA) provides spectral and temporal measurements of laser-stimulated emission (LSE) for assessment of phytoplankton pigments, community structure, photochemical efficiency (PY), and chromophoric dissolved organic matter (CDOM). The instrument was deployed in the Northern Gulf of Mexico to evaluate the ALFA analytical capabilities across the estuarine-marine gradient. The robust relationships between the pigment fluorescence and independent pigment measurements were used to validate the ALFA analytical algorithms and calibrate the instrument. The maximal PY magnitudes, \( \text{PY}_m = \text{PY}(1-1.35 \times 10^{-4} \text{PAR})^{-1} \), were estimated using the underway measurements of PY and photosynthetically active radiation (PAR). The chlorophyll (Chl) spatial patterns were calculated using the ratio of Chl fluorescence to PY to eliminate the effect of non-photochemical quenching on the underway Chl assessments. These measurements have provided rich information about spatial distributions of Chl, PY, CDOM, and phytoplankton community structure, and demonstrated the utility of the ALFA instrument for oceanographic studies and bio-environmental surveys. The data suggest that the fluorescence measurements with 514 nm excitation can provide informative data for characterization of the CDOM-rich fresh, estuarine, and coastal aquatic environments.

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References and links


1. Introduction

The Advanced Laser Fluorometry (ALF; see the Table of Abbreviations in Appendix [Table 1]) [1] is an analytical technique that addresses the optical complexity of natural aquatic environments to characterize the key fluorescence constituents (phytoplankton pigments, biomass, photophysiology, phycobiliprotein (PBP) containing phytoplankton groups, and CDOM). The original ALF fluorometer is a compact flow-through instrument that combines spectral and temporal LSE measurements using dual-wavelength excitation at 405 and 532 nm. It is used for underway shipboard measurements and analysis of discrete water samples [1–3]. The spectral deconvolution (SDC) analysis of the LSE signatures is used to assess the overlapped spectral bands of aquatic fluorescence constituents. Along with the accurate concentration measurements, the SDC provides quantification of the non-chlorophyll fluorescence background in the spectral area of Chl fluorescence for more accurate assessments of phytoplankton photo-physiological parameter, PY [1,4]. The ALF measurements of variable fluorescence can be used to improve assessments of Chl concentration [2].

The major ALF advantages over other techniques capable of phytoplankton structural characterization are that it provides additional useful information about phytoplankton photophysiology and CDOM, conducts the measurements directly in the sampled water without any preparation and treatment of the sample, can be used for fast automatic real-time analysis in various settings, including high-resolution underway measurements from small vessels and motorboats [1]. On the other hand, the ALF capacity for phytoplankton structural characterization is limited to assessment of the PBP-containing cyanobacteria and cryptophytes, while other techniques used in the oceanography, such as flow cytometry or DNA analysis, can provide more detailed information on phytoplankton community structure and size distributions (e.g., [5]), as well as activity and functional roles of the dominant species (e.g., [6]).

In 2006-2011, the ALF technique was extensively tested in diverse water types (including Chesapeake and Delaware Bays, the California Current, Bering Sea, Gulf of Mexico, and the Amazon River Plume) and proved to be a useful tool for aquatic research and environmental monitoring. It was used for assessment of Chl concentration, phytoplankton biomass and photochemical efficiency (PY), detection and quantification of PBP-containing cyanobacteria and cryptophytes, and measurements of CDOM spatial variability [1–3, 7–9].

These field deployments provided useful information to identify the ALF design solutions and analytical algorithms that needs to be further refined. The new instrument components, including miniature spectrometers and lasers with new emission wavelengths, have become available. These factors facilitated development of the next generation ALF technology. Several new options for the optical design and ALF instrument configuration were developed [10]. A single-laser (514 nm) and three-laser ALF-T instruments comprising new 514 and 375 nm lasers were built and tested in the laboratory and field conditions [4]. These and earlier ALF measurements were analyzed to optimize the spectral excitation for operation in various water types [11].

A new commercial instrument, the Aquatic Laser Fluorescence Analyzer [12] (ALFA), has been recently developed on this basis in collaboration between the Lamont-Doherty Earth Observatory of Columbia University and Western Environmental Technology Laboratories,
Inc. (WET Labs). The benchtop ALFA instrument operationally implements the key design solutions, measurement protocols and analytical methods of the ALF technique. It provides spectrally and temporally resolved LSE measurements for assessment of phytoplankton pigment biomass, PBP-containing phytoplankton groups, PY, and CDOM. It can be used for laboratory and field flow-through measurements with an external water supply, and for analysis of water samples using a miniature peristaltic pump installed on the front panel of the instrument case [Fig. 1]. In the dual-laser instrument configuration, ALFA-405/514, the alternate laser excitation at 405 and 514 nm is used for characterization of the fluorescence constituents in a broad range of water types, including fresh, estuarine, coastal and offshore waters [11].

![ALFA instrument setup](image1.jpg)

**Fig. 1.** Left: The ALFA instrument setup for the underway flow-through measurements and discrete sample analysis (R/V Pelican; Apr. 2012). Right: A map of ALFA underway measurements and sampling stations in the Northern Gulf of Mexico in April 2012.

In this article, we present results of the initial field evaluation of the ALFA-405/514 instrument and its analytical capabilities during a research cruise of the R/V Pelican in the Northern Gulf of Mexico in April 2012. The new ALFA instrument was used in conjunction with traditional oceanographic instruments (Turner 10AU fluorometer) and analytical techniques (high performance liquid chromatography (HPLC)) to provide additional, often missing information about spatial variability in phytoplankton biomass, photochemical efficiency, community structure, and CDOM abundance across the estuarine-marine gradient in the coastal zone influenced by the Mississippi-Atchafalaya River system. The measurements were conducted in collaboration between the research teams of Columbia University, Texas A&M University (Galveston campus), and WET Labs.

The Mississippi-Atchafalaya River system is the end-point for the largest drainage system in the United States, moving water from parts or all of 31 states [13,14]. While many studies focused on the relationship between phytoplankton biomass and productivity to nutrients in the Northern Gulf of Mexico (e.g., [13–15]), few studies have investigated phytoplankton physiology [16–19] and community composition (e.g., [19–21]). This is partly due to the challenges associated with traditional approaches such as microscopic taxonomy or HPLC, which are very time/labor consuming and require professionally trained specialists. The ALF and other new optical techniques can provide a means for field studies and environmental surveys to improve our understanding of aquatic biogeochemical processes and advance our capacity to monitor state of health of aquatic ecosystems.

2. ALFA measurements

The ALFA-405/514 measurement protocols are similar to those used in the ALF-T instrument [3]. During the in-line underway measurements, the ALFA instrument continuously repeats the basic measurement cycle including:

1. the broadband (400-800 nm) LSE spectral measurement with the 405 nm excitation,
2. the broadband (500-800 nm) LSE spectral measurement with the 514 nm excitation,
(3) the pump-during-probe (PDP [1,4,22]) temporal measurement of the LSE induction over 200 μs with the 405 nm excitation,

(4) the PDP temporal measurement of the LSE induction over 200 μs with the 514 nm excitation.

The discrete sample analysis begins with two PDP induction measurements using the 405 and 514 nm excitation, followed by the LSE spectral measurements with the alternate 405 and 514 nm excitation. The spectral integration time is automatically adjusted in 0.1-3 s range to optimize the signal:noise ratio and avoid saturation of the CCD sensor. The PDP induction curves are averaged over the preset number of excitation shots to ensure representative measurements in the sample volume and improve the signal. The gain and input range of the PDP sensor are automatically adjusted to optimize the PDP measurements.

The raw spectra are corrected for the instrument spectral response immediately after the measurements as described in [4] to obtain the instrument-independent spectral data. The real-time SDC analysis of the overlapped spectral bands is incorporated in the ALFA software for assessment of Chl concentration, PBP pigments and CDOM. The intensities of fluorescence bands are normalized to water Raman scattering (WR) to improve the accuracy of fluorescence assessments [1,23,24]. The PY values (i.e., the quantum yield of photochemistry in photosystem II) are also derived in real time from the spectrally corrected PDP induction measurements using a bio-physical model as described in [1,4].

The ALFA measurements described herein were conducted in April 24-30, 2012. The ALFA instrument was installed in the laboratory of the R/V Pelican [Fig. 1] and used for analysis of discrete water samples collected at various depths at the stations, and for high-resolution underway flow-through measurements between the sampling stations. A map in Fig. 1 shows locations of the sampling stations (marked with letters) and the transect lines of the underway ALFA measurements between the stations. Overall, 8353 underway measurement cycles were conducted by the ALFA instrument over the total distance of 601.6 km, resulting in the average spatial resolution of 72 m. For the underway operation, the instrument was connected to the shipboard uncontaminated system that provided continuous sampling of sub-surface water at 2.5 m depth. The ALFA was controlled by a notebook computer using software developed under the LabView environment (National Instruments). A compact GPS unit (MAP76Cx, Garmin) with an external marine antenna (Model 010-12017-00, Garmin) was connected to the instrument computer to provide coordinate and time information during the underway ALFA measurements. The PAR sensor, model SQ-110 (Apogee Instruments), was installed on the upper deck and connected to the ALFA instrument to import the PAR data during the ALFA underway measurements.

Examples of the ALFA measurements in diverse water types are shown in Fig. 2, displaying the screenshots of the ALFA user interface during the measurements. The measurement locations are marked with letters A, B, C, and D, respectively, on the map in Fig. 1. The measured LSE spectra with 405 and 514 nm excitation are displayed with blue and green dots in the left and middle panels, respectively; the golden lines represent the SDC best fits based on the linear scaling of the spectral components [1,4]. The apparent changes in intensity of the CDOM and Chl fluorescence (CF) (blue and green bands) vs. WR peak (red) reflect variations in the concentration of these constituents in fresh (A), coastal (B), and offshore (C and D) waters. A substantial variability in phycoerythrin (PE) fluorescence of green-water cyanobacteria (yellow peak at 578 nm) vs. WR is also evident in the LSE spectra measured with 514 nm excitation (middle panels). The PE fluorescence bands of eukaryotic cryptophytes (590 nm) and blue-water cyanobacteria (565 nm) were also detected by the SDC analysis of the green-stimulated LSE spectra measured offshore (panels C514 and D514). The fluorescence band of phycocyanin present in photosynthetic apparatus of these PBP-containing phytoplankton groups [1,3] can be seen in the green-stimulated spectra at 644 nm (middle column in Fig. 2). Consistently with the earlier observations (e.g., [1,4,11,25,26]), the
spectral overlap of emission bands of various aquatic constituents is evident in a broad spectral range, particularly in the green-stimulated spectra. The real-time SDC analysis is incorporated in the ALFA software to quantify the intensities of spectral bands of the fluorescence constituents and WR scattering [1,4].

Fig. 2. ALFA measurements using laser excitation at 405 nm (left column) and 514 nm (middle column: LSE spectra; right column: PDP induction). The locations of measurements are marked as A, B, C, and D, respectively, in Fig. 1.

The temporal PDP measurements of the LSE induction in the CF spectral area (~680 nm) using 514 nm excitation are displayed in the right column of Fig. 2. The “0” time marks the beginning of the excitation flash; the blue horizontal line visualizes the intensity of the non-Chl fluorescence background calculated from the SDC analysis of the spectra displayed in the middle column of Fig. 2. This background is subtracted from the LSE induction before best fitting with the biophysical model (white line) to assess the PY value [1,4].
Fig. 3. A,B: The regression relationships between Chl concentration (C_{chl}) in water samples and Chl fluorescence normalized to WR (F_{chl}) measured with 405 and 514 nm excitation. C,D: The regression relationships between C_{chl} in water samples and F_{chl} normalized to phytoplankton photochemical efficiency (PY) measured with 405 and 514 nm excitation (data for the same set of samples as in panels A and B).

3. Fluorescence analysis of Chl concentration and PBP-containing phytoplankton

The results of linear regression between the measurements of Chl concentration (C_{chl}) in pigment extracts [10] and the ALFA measurements of CF normalized to WR (F_{chl} [1,4],) are displayed in Figs. 3(A) and 3(B). The F_{chl}^{405} and F_{chl}^{514} values were calculated by the ALFA software using SDC analysis of the LSE spectra measured with 405 and 514 nm excitation, respectively, in the dark-adapted water samples collected at the locations marked with letters on the map in Fig. 1. The slope value in Fig. 3(A) is close to the magnitudes earlier derived in the estuarine and coastal waters using 405 nm excitation [1,2], but ~2-fold greater than the slope obtained in the California Current waters [4, 11]. The slope value in Fig. 3(B) is also ~2-fold higher than observed in the California Current. We assume that these differences could be caused by the distinct composition of oceanic phytoplankton in the California Current, resulting in their higher CF yield.

Figs. 3(C) and 3(D) show the linear regression relationships between C_{chl} and the fluorescence parameter F_{chl}/PY (i.e., the ratio of F_{chl} and PY values) measured using the 405 and 514 nm excitation, respectively. As shown in [2], the use of the F_{chl}/PY units for assessment of Chl concentration from the daytime underway fluorescence measurements may be beneficial, because it appears to be invariant with regard to the solar-induced non-photochemical quenching (NPQ) that may adversely affect the accuracy of C_{chl} assessments based on the F_{chl} underway measurements. As evident from Figs. 3(C) and 3(D), the F_{chl}/PY values have shown the robust regression relationships with the independent measurements of Chl concentration. The slope value in Fig. 3(C) is close to the slope magnitude measured in the estuarine waters of the Chesapeake and Delaware Bays (1.88 [2]), suggesting that this approach may potentially be used to improve the accuracy of the Chl fluorescence assessment in a broad range of coastal and estuarine water types.
Our recent field measurements in the California Current have demonstrated that the 514 nm laser excitation can be used for assessment of blue-water cyanobacteria and cryptophytes [11], but low concentration of green-water cyanobacteria has not permitted testing the 514 nm excitation for their fluorescence assessment. Fortunately, we found high enough concentration of green-water cyanobacteria in the coastal zone of Northern Gulf of Mexico to conduct the test. Similar to our approach used for blue-water cyanobacteria and cryptophytes [1,11], the test was based on analyzing the relationship between the accessory pigment zeaxanthin and the specific for green-water cyanobacteria PE fluorescence peak at 578 nm [1], normalized to WR scattering ($F_{PE2\text{514}}$).

As shown by earlier studies (e.g., [27,28]), photoacclimation of cyanobacterial photosynthetic apparatus to low-light conditions in deeper water layers may lead to significant increase in the PE cellular quotas with depth. During our recent measurements in the California Current we found that this may result in considerable, up to 10-fold, increase with depth in the cyanobacterial PE fluorescence per unit of their carbon biomass, which complicated fluorescence assessments of cyanobacterial biomass in the euphotic layer [3]. On the other hand, the ALFA data has shown strong linear regression relationship between the cyanobacterial PE fluorescence and their biomass in the subsurface water layer above the Chl maximum ($R^2 = 0.95$ vs. 0.42 for the entire euphotic layer), which was used to build the high-resolution surface distribution of *Synechococcus* carbon biomass using the ALFA underway PE fluorescence measurements [3].

During our ALFA field measurements in Northern Gulf of Mexico, most of the water samples were collected in the 0.3-12 m subsurface layer above the Chl maximum, and the maximum depth of sampling did not exceed 19 m, which was above or below the shallow Chl maximum, depending on the station location. To evaluate the ALFA potential for assessment of cyanobacterial biomass, we analyzed both the entire data set, and the subset of subsurface (0.3-12 m) ALFA measurements of cyanobacteria-specific PE fluorescence vs. HPLC concentration measurements of accessory carotenoid pigment zeaxanthin.

Although zeaxanthin is often used as a cyanobacterial pigment biomarker (e.g. [29]), it is not 100% specific for cyanobacteria. Photosynthetic apparatus of chlorophytes and eustigmatophytes (non-PBP-containing phytoplankton groups) may also contain zeaxanthin. The potential presence of these two groups in substantial and variable proportion vs. cyanobacteria may result in reduced correlation between the cyanobacteria-specific PE fluorescence and less-specific zeaxanthin. In fact, we obtained quite robust linear regression relationship between the ALFA $F_{PE2\text{514}}$ measurements and zeaxanthin concentration in the subsurface water layer (0.3-12 m) above the Chl maximum (Fig. 4(A); $R^2 = 0.86$). This suggests that zeaxanthin was mainly associated with green-water cyanobacteria in the area of study. Indeed, our ALFA underway measurements have shown relatively high concentration of green-water cyanobacteria, and large areas where they comprised a significant fraction of
phytoplankton community (see Section 6). We also obtained reasonably good correlation between FPE2\textsubscript{514} and zeaxanthin for the entire data set (R\textsuperscript{2} = 0.77). Consistent with our earlier observations [3], it was reduced vs. the sub-surface data above the Chl maximum, likely due to the gradual photo-acclimative increase in the PE fluorescence:biomass ratio below the Chl maximum. Further, in a parallel study using HPLC to examine pigment composition of water samples and phytoplankton community analysis during the same cruise [19], chlorophytes and eustigmatophytes comprised ≤ 5% of the community.

Interestingly (and in contrast with our observations in the offshore oceanic waters [1,3]), the FPE\textsubscript{1514} fluorescence (PE peak at 565 nm; normalized to WR) of blue-water cyanobacteria [1], less abundant in the area of study, showed weak correlation with zeaxanthin concentration, which was likely associated with more abundant green-water cyanobacteria. Consistent with such assumption, the regression relationship between zeaxanthin and total cyanobacterial fluorescence FPE\textsubscript{12514} = FPE\textsubscript{1514} + FPE\textsubscript{2514} appeared to be almost as robust for the green-cyanobacteria (R\textsuperscript{2} = 0.82 for the subsurface (0.3-12 m) samples above Chl maximum; Fig. 4(B)). The ALFA measurements of FPE\textsubscript{3514} fluorescence of cryptophytes (PE peak at 590 nm; normalized to WR) also indicated their relatively low abundance vs. green-water cyanobacteria (maximal FPE\textsubscript{3514} ~0.12 vs. 0.56 for FPE\textsubscript{2514}; Fig. 4). Nonetheless the FPE\textsubscript{3514} data showed good correlation with HPLC measurements of their specific pigment biomarker, alloxanthin, for the entire set of samples from 0.3 to 19 m (Fig. 4(C); R\textsuperscript{2} = 0.69). In contrast with cyanobacteria, the subsurface (0.3-12 m) subset of data for cryptophytes did not show higher correlation, suggesting no clear photo-acclimative trends in the FPE\textsubscript{3} cryptophyte fluorescence.

The robust regression relationships between the group-specific PE fluorescence and concentration of pigment biomarkers associated with the PBP-containing phytoplankton demonstrate that the ALFA spectral measurements of fluorescence stimulated at 514 nm can be used for detection, identification, and quantitative assessments of the PBP-containing cryptophytes and cyanobacteria. These and earlier published data (e.g., [1,3]) suggest that the cryptophyte biomass can be estimated with reasonably good accuracy in the entire euphotic layer. For cyanobacteria, it can be also done in the subsurface layer above the Chl maximum [3]. An important practical application is that the underway fluorescence measurements from various moving platforms (ships, small vessels, autonomous surface vehicles) can be used for real-time high-resolution assessments of horizontal distributions of cyanobacterial biomass along with other key aquatic characteristics (Chl, PY, CDOM, etc.; for example, see [3]). The ALF technique can be also employed for long-term monitoring of cyanobacterial biomass and other relevant characteristics in the surface waters in stationary settings (buoys, platforms, etc.). The methodology of fluorescence assessment of cyanobacterial biomass at the euphotic depths below Chl maximum however needs additional research, involving (and accounting for) the photo-acclimative changes in the cyanobacterial fluorescence:biomass ratio.

The complementary chromatic adaptation (CCA) of cyanobacterial photosynthetic apparatus (e.g., [30]) should be considered in this context among other photo-acclimative mechanisms. A detailed discussion of this biophysical phenomenon is beyond the scope of this publication focused on the field evaluation of the new instrument. To summarize, some (not all (e.g., [31])) cyanobacterial strains may exhibit the CCA that results in altering the relationship between light absorption by the phycourobilin (PUB) and phycoerythrobilin (PEB) chromophore due to the prolonged (~days) effect of change in spectral composition of the ambient light. The CCA does not affect the spectral characteristics of PE fluorescence and, therefore, the accuracy of spectral fluorescence detection and identification of the PBP-containing cyanobacteria (e.g [26]). On the other hand, it may alter the cyanobacterial light absorption in the blue-green spectral range, and, therefore, affect the efficiency of fluorescence excitation with a green laser. The CCA may develop with depth in the stratified euphotic layer because of alteration in the spectral composition of the ambient light field. In particular, it could contribute to the increase with depth in the cyanobacterial PE
fluorescence:biomass ratio (e.g., [3]). The robust regression relationships in Figs. 4(A) and 4(B)) suggest no significant CCA effect in the subsurface water layer (0.3-12 m) in the area of study. The underway ALFA measurements of PE fluorescence are analyzed below in Sections 4 and 6 under this assumption.

4. ALFA underway transect measurements

Fig. 5 shows an example of underway measurements with the ALFA instrument during the outbound ship transient between the sampling stations E and D [Fig. 1]. The dots represent 1626 measurements automatically conducted over a distance of 79.1 km, resulting in spatial resolution of 48.6 m per measurement cycle. The measurements began at 6:54 am (Central Standard Time), and were finished at 13:10 pm the day of April 26, 2012 (right to left on Fig. 5). The ALFA underway measurements have provided quite informative data set for characterization of transition from coastal to offshore marine environment. In particular, the gradual decline in CDOM fluorescence ($F_{CDOM}^{514}$; Fig. 5(C)) indicated an outbound decrease in the CDOM content. Even stronger, ~8-fold offshore CF reduction was observed along the transect line ($F_{chl}^{514}$; Fig. 5(A)). Though this decrease was generally consistent with the transition from the high-biomass coastal to low-biomass offshore waters, a potential diel modulation by the solar-induced NPQ might also contribute to the CF transect variability. As discussed in [2], the photoprotective NPQ regulation may affect both CF and PY measurements of sub-surface phytoplankton, which has to be accounted for when interpreting the underway measurements over extended period of time. As evident from Fig. 5(C), the PAR intensity on the water surface increased from 146 μM photons m$^{-2}$ s$^{-1}$ at 6:54 in the morning, when departing station E [Fig. 1], to >2000 μM photons m$^{-2}$ s$^{-1}$ at the end of the transect (station D; Fig. 1). Such significant PAR rise could result in the gradual NPQ development in the subsurface phytoplankton, leading to the NPQ-caused reduction in the PY and CF values [2]. Indeed, the diel increase in PAR along the transect line was accompanied by the gradual, up to 30% decline in PY ($PY_{514}$; Fig. 5(C)). Assuming that the PY reduction was primarily caused by the NPQ development rather than an offshore decrease in phytoplankton photochemical efficiency, we analyzed the correlation between the $PY_{514}$ and PAR values measured along the transect line. The strong linear regression relationship between these variables ($R^2 = 0.83$; Fig. 6(A)) appeared to be consistent with such assumption.

We speculate that (i) the solar-induced NPQ was indeed the main cause of the underway $PY_{514}$ variability along this transect line, and (ii) the maximal potential PY values for dark-adapted (PAR = 0) phytoplankton, $PY_{m}$, would be relatively high (~0.5) all across the transect line. The latter is consistent with the earlier PY measurements in the nutrient-rich coastal and near-shore waters during the spring bloom in the northern Gulf of Mexico and other areas (e.g., [16–19,32]). The CF and PY decrease with increasing PAR was also apparent in the underway ALFA measurements at other locations during this survey, but the spatial/temporal patterns were more complex. It likely reflected a combined effect of both diel NPQ modulation and spatial variability in CF and PY caused by other natural factors, such as nutrient supply and mixing in the water column.

As shown in [2], the NPQ effect on the accuracy of $C_{chl}$ assessments from the underway fluorescence measurements can be eliminated by using the NPQ-invariant CF:PY ratio, instead of the NPQ-affected CF, to calculate the $C_{chl}$ values. Following this approach, we calculated the $C_{chl}$ transect distributions from the ALFA underway measurements [Fig. 7(A)] using the robust regression relationship $C_{chl} = 0.50 \cdot F_{chl}^{514}/PY_{514}$ obtained for the water samples collected at the stations [Fig. 3(D)]. Examples of $C_{chl}$ transect distribution calculated using the $F_{chl}^{514}$ and $PY_{514}$ underway data (Figs. 5(A) and 5(C), respectively) is shown in Fig. 5(C). Note that it indicates a smaller, ~5-fold offshore decline in Chl concentration vs. up to 8-fold decline in CF [Fig. 5(A)], which was also affected by the gradual diel NPQ development during the outbound transect measurements. Based on these $C_{chl}$ estimates, the
offshore waters in the western part of the transect can be qualified as mesotrophic \((0.1 < C_{chl} < 1 \ \mu g \ L^{-1})\), while the coastal waters in the eastern part of the transect were rather eutrophic \((C_{chl} > 1 \ \mu g \ L^{-1})\). Such relationships have been previously documented in this region (e.g., [18,19]). Assuming that the robust relationship between the PY and PAR [Fig. 6(A)] describing the diel NPQ regulation of PY was applicable to the entire area of our survey, we used the ALFA underway measurements of PY and PAR to calculate the spatial distributions of physiological parameter \(PY_m = PY \ (1 - 1.35 \cdot 10^{-4} \ PAR)^{-1}\) that characterized the maximal potential PSII photochemical efficiency of dark-adapted phytoplankton, not affected by the NPQ (see Fig. 7(B)).

The ALF underway measurements of pigment fluorescence can also provide important (though not comprehensive) information about the spatial variability in phytoplankton composition [3,8,9]. In particular, the transect distribution of \(F_{PE1} 514\) fluorescence specific for blue-water cyanobacteria displayed in Figs. 5(A) and 5(B) indicates ~30% increase in their biomass in the offshore portion of the surveyed area. The \(F_{PE2} 514\) and \(F_{PE3} 514\) patterns show the opposite trends, with an overall offshore decrease in green-water cyanobacteria and cryptophytes, respectively (the former were not detected at all in the middle of the transect). The contrast between the coastal and offshore portions of the transect is even more apparent in the \(F_{PE}/F_{chl}\) patterns [Fig. 5(B)] indicating the substantial offshore increase in relative abundance of the PBP-containing photosynthesizing groups in phytoplankton community [1,3]. The particularly striking, 10-fold increase in the \(F_{PE1} 514 \cdot F_{chl}\) ratio was detected for the blue-water cyanobacteria that have high PUB/PEB ratio, making them better adapted for light harvesting in the offshore blue waters ([1] and references therein). The maximal \(F_{PE1} 514 \cdot F_{chl}\) values ~0.3 suggest that the blue-water cyanobacteria were dominant in the offshore phytoplankton community, comprising up to 80% of total phytoplankton biomass [3]. On the
other hand, the relatively low $F_{PE}/F_{Chl}$ magnitudes in coastal waters in the eastern portion of the transect suggests that the elevated Chl concentration observed there [Fig. 5(C)] could be attributed to the increase in biomass of non-PBP-containing eukaryotic phytoplankton.

![Graph](image)

Fig. 6. (A): The regression relationship between the photosynthetically active radiation, PAR, and the ALFA underway measurements of phytoplankton photochemical efficiency, $PY^{514}$, measured underway between the sampling stations D and E (Fig. 1). (B, C): The regression relationships between the ALFA underway measurements of phytoplankton photochemical efficiency, $PY$, and CDOM fluorescence normalized to WR scattering, $F_{CDOM}$, measured with 405 and 514 nm laser excitation. The data for all transects shown in Fig. 1 are displayed.

5. Fluorescence assessments of phytoplankton photochemical efficiency and CDOM using ALFA dual-wavelength excitation

The ALFA field measurements described in this article provided an opportunity to evaluate the analytical capabilities of the ALF instrument configured with a new 514 nm diode laser as source of fluorescence excitation in the diverse water types across the estuarine-marine gradient. As evident from the data displayed in Figs. 3 and 4, the 514 nm laser excitation can serve well for pigment fluorescence assessments of Chl concentration and phytoplankton community structure. An example of transect measurements in Fig. 5 shows that it can also provide reasonably good signal:noise ratio for measuring $PY$ and CDOM fluorescence. In Figs. 6(B) and 6(C) we present the results of correlation analysis of underway measurements of these two parameters along all the transect lines displayed in Fig. 1 in a broad range of water types, including fresh, estuarine, coastal and offshore waters.

The results of measuring $PY$ with 405 and 514 nm excitation in the Northern Gulf of Mexico appeared to be consistent with our measurements in the Ligurian Sea (Mediterranean) [11]: both data sets have shown high correlation ($R^2 = 0.89$ [Fig. 6(B)] vs. 0.98 [11]) between the $PY^{514}$ and $PY^{405}$ values, and very close slope magnitudes (0.85 [Fig. 6(B)] vs. 0.87 [4]). Following the interpretation given in [11], we speculate that the 13-15% higher values of $PY^{514}$ vs. $PY^{405}$ may be caused by the greater photochemical efficiency of cyanobacteria. They comprised significant fraction of phytoplankton community (Fig. 5(B); see also Figs. 7(G) and 7(H) and relevant discussion below), but did not adequately contribute to the bulk $PY^{405}$ measurements since the blue light is low-efficient for excitation of their pigment fluorescence (e.g., [1]).

Similar to the ALF measurements in the Gulf of Mexico area adjacent to the Mississippi River delta [11], our analysis of the underway ALFA measurements in the Northern Gulf of Mexico has revealed a robust linear regression relationship between the $F_{CDOM}^{514}$ and $F_{CDOM}^{405}$ (Fig. 6(C); $R^2 = 1$ vs. 0.95 reported in [1]; the slope value is 1.97 vs. 1.81, respectively). The maximal $F_{CDOM}$ magnitudes were measured in the Atchafalaya River, indicating very high CDOM content. An apparent deviation from the regression trendline for $0.3 < F_{CDOM}^{514} < 0.8$ measured in coastal waters may suggest some structural changes in CDOM components contributing to the CDOM fluorescence.
6. ALFA underway mapping of fluorescence patterns

The ALFA underway measurements have provided rich and informative high-resolution data to characterize spatial distributions of the key fluorescence characteristics in the Northern Gulf of Mexico during the spring phytoplankton bloom in April, 2012 [Fig. 7]. In this section, we briefly discuss these fluorescence patterns to illustrate the analytical capabilities of the ALF technique. A detailed discussion of the oceanographic features of this survey is beyond the scope of this article, and will be given elsewhere. The maps displayed in Fig. 7 were built using ALFA measurements with 514 nm excitation; some distinct features of the fluorescence patterns are marked as 1-6.

The spatial distribution of Chl concentration [Fig. 7(A)] was calculated using the regression equation $C_{\text{chl}} = 0.50 F_{\text{chl}}^{514}/P_{\text{Y}}^{514}$ derived from analyzing discrete water samples at the stations [Fig. 3D]. The NPQ-invariant $F_{\text{chl}}/P_{\text{Y}}$ parameter was used to eliminate the $F_{\text{chl}}$ modulation by the NPQ and improve the accuracy of $C_{\text{chl}}$ assessments (section 4 and [2]). A significant range of $C_{\text{chl}}$ variability, 0.4-12 $\mu$g L$^{-1}$, was detected across the estuarine-marine gradient surveyed in this area. The offshore and coastal locations of $C_{\text{chl}}$ minimum and maximum, respectively, are marked as “4” and “6” in Fig. 7.

Fig. 7(B) shows the spatial distribution of phytoplankton physiological parameter $P_{\text{Y}}$, the NPQ-corrected phytoplankton photochemical efficiency. It was calculated using the ALFA underway measurements of the actual, potentially affected by the NPQ photochemical efficiency, $P_{\text{Y}}$, and PAR as $P_{\text{Y}} = P_{\text{Y}} (1 - 1.35 \cdot 10^{-4} \text{PAR})^{-1}$, assuming that the regression relationship in Fig. 6(A) is applicable to the entire surveyed area. Though different phytoplankton groups may have different NPQ mechanisms, the ChemTax analysis of the HPLC pigment measurements at the stations has shown that diatoms were dominant phytoplankton group (accounting for 40-80% phytoplankton biomass) in the area of study [19], despite the structural variability among other, subdominant phytoplankton groups. Since the ALFA fluorescence measurements reflect the bulk properties of phytoplankton community, the $P_{\text{Y}}$ values and NPQ mechanisms involved were therefore mainly determined by the diatoms. This justifies our assumption regarding the applicability of the regression equation for the NPQ correction of the $P_{\text{Y}}$ measurements. The $P_{\text{Y}}$ values were derived from the ALFA fluorescence induction measurements corrected for the non-Chl spectral background [1,4], particularly significant in high-CDOM fresh and estuarine waters [Fig. 2A].

The $P_{\text{Y}}$ values also exhibited substantial variability across the area, 0.3-0.65. The highest values, $P_{\text{Y}}^{514} \approx 0.6-0.65$, were found in the estuarine waters adjacent to the mouth of the Atchafalaya River. These values are close to the maximal photochemical efficiency, 0.65, measured using single-turnover technique for phytoplankton grown in the optimized laboratory conditions (e.g., [33]). It can be rarely observed in the field (e.g., [34]), which is consistent with earlier observations of high photochemical efficiency of healthy phytoplankton in the nutrient-rich estuarine waters in the Gulf of Mexico [16,17]. In contrast, the lowest $P_{\text{Y}}$ magnitudes, indicating low phytoplankton photochemical efficiency, were found offshore (“5” in Fig. 7), where the reduced Chl concentration [Fig. 7(A)], low CDOM content [Fig. 7(C)], and high fraction of green-water cyanobacteria in phytoplankton population [Fig. 7(H)] were also detected.

The CDOM fluorescence distribution [Fig. 7(C)] also varied in a broad range, showing two distinct areas of very high CDOM content in the Atchafalaya River and its estuary ($F_{\text{CDOM}}^{514} \approx 2$; see also Fig. 6(C)), and more moderate, “coastal” values 0.15 < $F_{\text{CDOM}}^{514} < 0.7$. Fig. 7(C) clearly shows the transition from the CDOM-rich freshwater/estuarine/coastal waters (“1”, “2”, and “3” in Fig. 7) to the relatively low-CDOM marine environments (“4”, “5”, and “6” in Fig. 7). The CDOM patterns appeared to be consistent with spatial distribution of seasurface salinity (not shown).
The fluorescence patterns of the PBP-containing phytoplankton groups [Fig. 7(D) - 7(I)] appeared to be more complex, reflecting both overall changes in phytoplankton biomass across the estuarine-marine gradient and structural changes within the phytoplankton population. Consistent with other ALF measurements in the Gulf of Mexico (and in contrast to our measurements in the oceanic waters ([1,3]), the green-water cyanobacteria appeared to be most abundant among the PBP-containing phytoplankton groups ($F_{PE2}^{514}$ up to 0.8 (“3” in Fig. 7(E))), and showed spatial patterns similar to the distributions of $C_{chl}$ and $F_{CDOM}$ [Figs. 7(A) and 7(C)]. They seemed to be better adapted to the specific bio-environmental conditions of the Gulf of Mexico. The green-water cyanobacteria comprised significant, dominant at some locations fraction of the phytoplankton population, mainly in the central and western offshore areas (“2” and “5” in Fig. 7(H); and patches “3”, “4”). The $F_{PE2}/F_{chl}$ values reaching 0.2-0.3 at “5” suggest that they might comprise up to 80% of total phytoplankton biomass in this offshore, low-productive [Figs. 7(A) and 7(B)] area.

The blue–water cyanobacteria were generally less abundant ($F_{PE1}^{514}$ < 0.1), showing their slightly elevated biomass mainly in the low-CDOM waters (“4”,”6”, “2” in Fig. 7(D)). Despite their relatively low concentration, they appeared dominant in the phytoplankton community in the low-CDOM [Fig. 7(C)] and low-Chl [Fig. 7(A)] offshore waters (“6” in Fig. 7). In contrast with our observations in oceanic environments and Delaware Bay (e.g [1,8,9], the PBP-containing eukaryotic cryptophytes were detected at relatively low concentrations ($F_{PE3}<$0.06 in Fig. 7(F)). Nonetheless, they comprised (along with dominant
blue-water cyanobacteria [Fig. 7(G)] a substantial component of phytoplankton population in the low-Chl, low-CDOM blue-water area “6” [Fig. 7(I)].

7. Conclusions

The new ALFA instrument was used for the underway shipboard measurements and discrete sample analysis to survey phytoplankton and CDOM across the estuarine-marine gradient in the Northern Gulf of Mexico. The total length of the measurement transect lines was 601.6 km; 8353 underway measurement cycles were conducted, resulting in the average spatial resolution of 72 m per cycle. The real-time shipboard fluorescence measurements of the water samples collected at the stations have shown the robust regression relationships with the independent measurements of pigment concentrations that were used for instrument calibration and validation. In particular, the NPQ-invariant ratio of regular and variable Chl fluorescence, $F_{\text{chl}}/F_{\text{PY}}$, was used to build the synoptic-scale map of Chl concentration in the area of survey. The robust linear regression relationship between the phytoplankton photochemical efficiency, PY, and photosynthetically active radiation, PAR, was derived from the transect data. It was used to correct the underway PY measurements for the solar-induced NPQ to assess the spatial patterns of the maximal phytoplankton photochemical efficiency, $P_{\text{Ym}}$. The comparative analysis of fluorescence measurements with excitation at 514 nm vs. 405 nm employed in the earlier instrument configurations suggests that the former can be used for fluorescence characterization of the CDOM-rich fresh, estuarine, and coastal aquatic environments. This justifies the development of compact and affordable, yet informative single-laser fluorometers for fluorescence analysis of natural aquatic environments.

Appendix

Table 1. Abbreviations used in text.

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Term</th>
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<tbody>
<tr>
<td>ALF</td>
<td>Advanced Laser Fluorometry/Fluorometer</td>
</tr>
<tr>
<td>ALFA</td>
<td>Aquatic Laser Fluorescence Analyzer</td>
</tr>
<tr>
<td>CCD</td>
<td>Charge-coupling device</td>
</tr>
<tr>
<td>CDOM</td>
<td>Chromophoric dissolved organic matter (substance)</td>
</tr>
<tr>
<td>Chl</td>
<td>Chlorophyll a (pigment)</td>
</tr>
<tr>
<td>$C_{\text{chl}}$</td>
<td>Chl a concentration, $\mu g , L^{-1}$</td>
</tr>
<tr>
<td>CF</td>
<td>Chlorophyll fluorescence</td>
</tr>
<tr>
<td>$F_{\text{CDOM}}$</td>
<td>CDOM fluorescence intensity normalized to WR intensity</td>
</tr>
<tr>
<td>$F_{\text{chl}}$</td>
<td>CF intensity normalized to WR intensity</td>
</tr>
<tr>
<td>$F_{\text{PE1}}$</td>
<td>PE fluorescence of blue-water cyanobacteria (peak at 565 nm) normalized to WR</td>
</tr>
<tr>
<td>$F_{\text{PE2}}$</td>
<td>PE fluorescence of green-water cyanobacteria (peak at 578 nm) normalized to WR</td>
</tr>
<tr>
<td>$F_{\text{PE3}}$</td>
<td>PE fluorescence of cryptophytes (peak at 590 nm) normalized to WR</td>
</tr>
<tr>
<td>HPLC</td>
<td>High performance liquid chromatography</td>
</tr>
<tr>
<td>LSE</td>
<td>Laser-stimulated emission</td>
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<tr>
<td>NPQ</td>
<td>Non-photochemical quenching</td>
</tr>
<tr>
<td>PAR</td>
<td>Photosynthetically active radiation</td>
</tr>
<tr>
<td>PBP</td>
<td>Phycobiliprotein (pigments)</td>
</tr>
<tr>
<td>PDP</td>
<td>Pump-during-probe (measurement protocol)</td>
</tr>
<tr>
<td>PE</td>
<td>Phycocerythrin (pigment)</td>
</tr>
<tr>
<td>PEB</td>
<td>Phycocyanobilin (chromophore)</td>
</tr>
<tr>
<td>PUB</td>
<td>Phycourobilin (chromophore)</td>
</tr>
<tr>
<td>PY</td>
<td>Photochemical efficiency of phytoplankton photosystem II</td>
</tr>
<tr>
<td>$P_{\text{Ym}}$</td>
<td>Maximal PY magnitude in the absence of NPQ</td>
</tr>
<tr>
<td>$R^2$</td>
<td>Coefficient of determination</td>
</tr>
<tr>
<td>SDC</td>
<td>Spectral deconvolution (method, technique)</td>
</tr>
<tr>
<td>WR</td>
<td>Water Raman scattering</td>
</tr>
<tr>
<td>WET Labs</td>
<td>Western Environmental Technologies, Inc.</td>
</tr>
</tbody>
</table>
Acknowledgments

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