

Lipid remodeling in phytoplankton exposed to multi-environmental drivers in a mesocosm experiment

Sebastian I. Cantarero¹, Edgar Flores¹, Harry Allbrook¹, Paulina Aguayo^{2,3,4}, Cristian A. Vargas^{3,5}, John E. Tamanaha⁶, J. Bentley C. Scholz⁶, Lennart T. Bach⁷, Carolin R. Löscher^{8,9}, Ulf Riebesell¹⁰, Balaji Rajagopalan¹¹, Nadia Dildar¹, Julio Sepúlveda^{1,5}

¹Department of Geological Sciences and Institute of Arctic and Alpine Research (INSTAAR), University of Colorado Boulder, Boulder, CO 80309, USA

²Departamento de Oceanografía, Universidad de Concepción, Casilla 160-C, Concepción, Chile

³Department of Aquatic System, Faculty of Environmental Sciences & Environmental Sciences Center EULA Chile, Universidad de Concepción, Concepción 4070386, Chile

⁴Institute of Natural Resources, Faculty of Veterinary Medicine and Agronomy, Universidad de Las Américas, Sede Concepcion, Chacabuco 539, Concepcion 3349001, Chile

⁵Millennium Institute of Oceanography (IMO), Universidad de Concepción, Concepción 4070386, Chile

⁶Laboratory for Interdisciplinary Statistical Analysis, Department of Applied Mathematics, University of Colorado, Boulder, Boulder CO, 80309, USA

⁷Institute for Marine and Antarctic Studies, University of Tasmania, Hobart, TAS 7004, Australia

⁸Nordcee, Department of Biology, University of Southern Denmark, Odense, Denmark

⁹Danish Institute for Advanced Study, University of Southern Denmark, Odense, Denmark

¹⁰Marine Biogeochemistry, GEOMAR Helmholtz Centre for Ocean Research Kiel, Düsternbrooker Weg 20, D-24105, Kiel, Germany

¹¹Department of Civil, Environmental, and Architectural Engineering, University of Colorado-Boulder, Boulder, CO, USA

Correspondence to: Sebastian I. Cantarero (sebastian.cantarero@colorado.edu)

Abstract. Lipid remodeling, the modification of cell membrane chemistry via structural rearrangements within the lipid pool of an organism, is a common physiological response amongst all domains of life to alleviate environmental stress and maintain cellular homeostasis. Whereas culture experiments and environmental studies of phytoplankton have demonstrated the plasticity of lipids in response to specific abiotic stressors, few analyses have explored the impacts of multi-environmental stressors at the community-level scale. Here, we study changes in the pool of intact polar lipids (IPLs) of a phytoplanktonic community exposed to multi-environmental stressors during a ~2-month long mesocosm experiment deployed in the eastern tropical South Pacific off the coast of Callao, Perú. We investigate lipid remodeling of IPLs in response to changing nutrient stoichiometries, temperature, pH, and light availability in surface and subsurface water-masses with contrasting redox potentials, using multiple linear regressions, classification and regression trees, and random forest analyses. We observe proportional increases of certain glycolipids (namely mono- and di-galactosyldiacylglycerols; MG and DG, respectively) associated with higher temperatures and oxic conditions, consistent with previous observations of their utility to compensate for thermal stress, and their degradation under oxygen stress. N-bearing (i.e., betaine lipids and phosphatidylethanolamine; BL and PE) and non-N-bearing (i.e., MG, phosphatidylglycerol, and sulfoquinovosyldiacylglycerol; PG, SQ) IPLs are anti-correlated, and have strong positive correlations with nitrogen repleted and depleted conditions, respectively, which suggests a substitution mechanism for N-bearing IPLs under nitrogen limitation. Reduced CO₂(aq) availability and increased pHs are associated with greater proportions of DG and sulfoquinovosyldiacylglycerol (SQ) IPLs, possibly in response to the lower

concentration of $\text{CO}_{2(\text{aq})}$ and the overall lower availability of inorganic carbon for fixation. A higher production of MG in surface waters corresponds well with their established photoprotective and antioxidant mechanisms in thylakoid membranes. The observed statistical relationships between IPL distributions, physicochemical parameters, and the composition of the phytoplankton community suggest evidence of lipid remodeling in response to environmental stressors. These physiological responses may allow phytoplankton to reallocate resources from structural or extrachloroplastic membrane lipids (i.e., phospholipids and betaine lipids) under high-growth conditions, to thylakoid/plastid membrane lipids (i.e., glycolipids and certain phosphatidylglycerols) under growth-limiting conditions. Further investigation of the exact mechanisms controlling the observed trends in lipid distributions are necessary to better understand how membrane reorganization under multi-environmental stressors can affect the pools of cellular C, N, P, and S, as well as their fluxes to higher trophic levels in marine environments subjected to increasing environmental pressure. Our results suggest that future studies addressing the biogeochemical consequences of climate change in the Eastern Tropical South Pacific Ocean must take into consideration the impacts of lipid remodeling in phytoplankton.

1 Introduction

The Eastern Tropical South Pacific (ETSP) is one of the most productive eastern boundary upwelling systems in the world (Chavez and Messié, 2009) and harbors one of the largest oxygen deficient zones (ODZs) (Fuenzalida et al., 2009; Ulloa and Pantoja, 2009; Thamdrup et al., 2012). Global warming has led to the expansion of ODZs over the recent decades and they are expected to continue expanding due to the reduction in oxygen solubility with increasing temperature (Stramma et al., 2008; Stramma et al., 2010; Gilly et al., 2013), as well as because of enhanced ocean stratification and reduced ventilation of the ocean's interior (Keeling et al., 2010). The future behavior of the ETSP upwelling system in a warmer world remains uncertain; increases in wind-induced upwelling intensity and duration (Gutiérrez et al., 2011; Bakun et al., 2010) may increase the supply of nutrients to the surface in coastal regions, whereas enhanced thermal stratification may reduce nutrient supply in the open ocean (Behrenfeld et al., 2006). Furthermore, upwelling regions are prone to highly variable pH (Capone and Hutchins, 2013), and the global ocean will experience a decreasing average pH as more CO_2 accumulates in the atmosphere and is absorbed by the ocean (Jiang et al., 2019). Accordingly, major shifts in marine planktonic community composition, turnover rates (Henson et al., 2021), and adaptations (Irwin et al., 2015) are expected in future scenarios of ocean conditions, which are expected to lead to cascading effects on ocean biogeochemistry and marine ecosystems (Hutchins and Fu, 2017).

Primary productivity in the ETSP is predominantly regulated by the wind-induced upwelling of nutrients, light availability, and Fe limitation (Messié and Chavez, 2015). Thus, changes in the supply of inorganic N along the upwelling region of the ETSP are likely to induce significant shifts in the phytoplankton community composition. Longer upwelling seasons along nearshore environments could further stimulate productivity of fast-growing eukaryotic algae that currently dominate these systems (e.g., diatoms; Messié et al., 2009). However, shorter upwelling seasons or weaker upwelling currents could favor more survivalist or mixotrophic algae, in addition to N-fixing diazotrophs that thrive under widespread nitrogen limitation (Dutkiewicz et al., 2012). The rate of primary productivity in the surface ocean not only affects the supply of sinking organic matter and thus oxygen consumption via microbial respiration in the subsurface (Wyrki, 1962), but also results in a shift of redox potentials that drive substantial losses of bioavailable N under reducing conditions at intermediate depths (Lam et al.,

2009; Wright et al., 2012). Additionally, expected changes in ocean warming and stratification (Huertas et al., 2011; Morán et al., 2010; Yvon-Durocher, 2015), lowered dissolved oxygen concentration (Wu et al., 2012), and decreased pH (Dutkiewicz et al., 2015, Bach et al., 2017), will disrupt phytoplanktonic assemblages differently based on their individual tolerances and physiological plasticity. However, little is known of the physiological adaptations of phytoplankton on a community-level scale in response to multi environmental stressors.

Phytoplankton have been shown to activate several lipid-based physiological mechanisms in response to environmental stimuli (Li-Beisson et al., 2019; Sayanova et al., 2017; Kong et al., 2018). In fact, intact Polar Lipids (IPLs) are a class of membrane lipids characterized by a polar head group typically attached to a glycerol backbone from which aliphatic chains are attached via ester and/or ether bonds (Sturt et al., 2004; Lipp et al., 2008; Schubotz et al., 2009; Van Mooy and Fredricks, 2010). Dominant planktonic lipid classes include phospholipids with a phosphate-bearing polar head group (e.g., phosphatidylcholine PC; phosphatidylethanolamine PE; and Phosphatidylglycerol PG), glycolipids featuring a sugar moiety in the polar head (e.g., monoglycosyldiacylglycerol MG; diglycosyldiacylglycerol DG; sulfoquinovosyldiacylglycerol SQ), and betaine lipids with a quaternary amine positively charged and attached to lipid chains (e.g. diacylglyceryl hydroxymethyl-trimethyl- β -alanine DGTA; diacylglyceryl trimethylhomoserine DGTS; and diacylglycerylcarboxy-N-hydroxymethyl-choline DGCC) (Kato et al., 1996; Rütters et al., 2001; Zink et al., 2003; Suzumura, 2005; Van Mooy et al., 2006). The remodeling of IPLs, the main constituents of cell and organellar membranes, provides numerous physiological adjustments to attenuate environmental stressors impacting phytoplankton (Zienkiewicz et al., 2016). These include nutrient limitation (Van Mooy et al., 2009; Meador et al., 2017; Abida et al., 2015; Wang et al., 2016), homeoviscous regulation in response to changing temperature (Sato et al., 1980; Sinensky, 1974; Neidleman, 1987) and pH (Tatsuzawa and Takizawa 1996; Poerschmann et al., 2004; Guckert and Cooksey, 1990; Jin et al., 2021), or photosynthetic function under varying light availability (Sato et al., 2003; Gašparović et al., 2013; Khotimchenko and Yakovleva, 2005). While IPL distributions in environmental studies are typically used as chemotaxonomic biomarkers that trace the presence and abundance of specific microbial groups (Sturt et al., 2004; Schubotz et al., 2009; Van Mooy and Fredricks, 2010), their distributions have been used in conjunction with additional microbial or geochemical measurements to assess how microbial metabolisms contribute to the chemical environment (Van Mooy et al., 2009; Wakeham et al., 2012; Schubotz et al., 2018; Cantarero et al., 2020). Yet, few studies have explored how multiple environmental drivers impact IPL remodeling at the community level and in time series, and the associated adaptability of phytoplankton to environmental change.

Lab-based culture experiments have been a major step forward in understanding how lipid remodeling may impact a biogeochemical system (as summarized above). However, a significant challenge remains in contextualizing these findings at the community scale. Conversely, observational studies from direct measurements of natural systems are often logistically limited in temporal scale, and consequently, do not fully capture the dynamics and heterogeneity of biogeochemical conditions. Mesocosms are experimental apparatuses at the interface between controlled culture experiments and environmental

observations that allow for the examination of natural systems and entire ecosystems under semi-controlled conditions to explore the impacts of a changing climate and ocean system (Riebesell et al., 2013). Here, we study changes in the composition, diversity, and abundance of phytoplanktonic IPLs in response to changes in the biological, physical, and chemical composition of a marine ecosystem subjected to semi-controlled conditions in a 2-month long mesocosm experiment off the coast of Perú. We investigate the potential for IPL remodeling amongst phytoplankton in response to multiple environmental stressors including nutrient availability, O₂ concentration, pH, temperature, and light availability to highlight adaptation strategies available to phytoplankton in response to a changing ocean system.

2 Methods

2.1 Mesocosm Deployment and Sampling

On February 22, 2017, eight “Kiel Off-Shore Mesocosms for Ocean Simulations” (KOSMOS; Riebesell et al., 2013) were deployed just north of Isla San Lorenzo, 6 km off the Peruvian coastline (12.0555° S, 77.2348° W; Fig. 1; Bach et al., 2020) in waters 30 m deep. Each individual mesocosm consisted of a cylindrical polyurethane bag (2 m diameter, 18.7 m length, 54.4 ± 1.3 m³ volume) suspended in an 8 m tall flotation frame. Water exchange was permitted via nets (mesh size 3mm) for three days before the water mass inside each mesocosm was enclosed and isolated from the surrounding Pacific water by attaching sediment traps at the base (~19 m length total). The enclosing of the mesocosms marked the start (day 0) of the 50-day experiment. As detailed in Bach et al. (2020), the experiment involved several manipulations including the addition of ODZ water to simulate upwelling conditions, salt to maintain water stratification, and the introduction of organisms.

The collection of subsurface waters and their addition to the mesocosms is described with detail in Bach et al. (2020). Briefly, on experiment days 5 and 10, two batches of local subsurface ODZ water were collected from stations with varying nutrient stoichiometries (Table 1; Fig. 1) using deepwater collectors first reported by Taucher et al., (2017). The water mass collected from 30 m deep at station 1 (12.028323° S, 77.223603° W) was characterized by a very low N:P ratio (Table 1), whereas the water collected from 70 m deep at station 3 (12.044333° S, 77.377583° W) was characterized by a higher but still low N:P ratio (Table 1). On experiment day 8, 9 m³ of water was removed from each mesocosm at a depth between 11 and 12 m and then on day 11, 10 m³ of ODZ water was injected into each mesocosm at depths between 14 and 17 m. On experiment day 12, the entire procedure was repeated but this time with 10 m³ removed between 8 and 9 m, and 12 m³ added evenly between 1 and 9 m. To maintain stratification and preserve the low O₂ subsurface layer, a NaCl brine solution was injected evenly into the subsurface of all mesocosms on experiment day 13 (0.067 m³, 10 to 17 m depth) and experiment day 33 (0.046 m³, 12.5 to 17 m depth). On day 14, Peruvian Scallop larvae (*Argopecten purpuratus*) were added (~10,000 individuals m⁻³), and on day 31 Fine Flounder eggs (*Paralichthys adspersus*) were added (~90 individuals m⁻³). However, few scallop larvae and no fish

larvae were found in the mesocosms after the release indicating that their influence on the plankton community was likely small (Bach et al., 2020).

We sampled two integrated water depths of the mesocosms for suspended organic matter and biological and chemical characterization using 5 L integrating water samplers (IWSs; Hydro-Bios, Kiel) equipped with pressure sensors to collect water evenly within a desired depth range. The samples were collected across two integrated intervals of the water column representing surface and subsurface layers, sampling depths were slightly modified over the course of the experiment to accommodate for changes in water stratification and the position of the chemocline (refer to Bach et al., 2020 for further details). The depths applied for surface and subsurface waters were 0–5 and 5–17 m on days 1 and 2, 0–10 and 10–17 m from day 3 to 28, and 0–12.5 and 12.5–17 m from day 29 to 50.

Samples of suspended organic matter for IPL analysis were collected from all 8 mesocosms at both the surface and subsurface sampling depths on non-consecutive days throughout the experiment. See details of changes in water depths above. Due to the labor- and time-intensive nature of IPL analysis, we focused on 4 mesocosms (two from each treatment) at both depths and from 9 different days spanning the 50-day experiment. We filtered 5 L of mesocosm water through pre-combusted, 142 mm Advantec glass fiber filters (GF75142MM) of 0.3 μm pore size. All samples were wrapped in combusted aluminum foil and shipped frozen to the Organic Geochemistry Laboratory at the University of Colorado, Boulder for IPL extraction and analysis.

2.2 Water Column Physicochemistry

Depth profiles of salinity, temperature, O_2 concentration, photosynthetically active radiation (PAR), and chlorophyll a (chl-a) fluorescence were measured through vertical casts using the CTD60M sensor system (Sea & Sun Technology). O_2 concentrations were cross-verified with the Winkler O_2 titration method performed via a micro-Winkler titration method described by Aristegui and Harrison (2002). Seawater pH_T (pH on total scale), was determined spectrophotometrically using m-cresol purple (mCP) indicator dye as described in Carter, et al. (2013); see Chen et al. (2022) for details. See Bach et al. (2020) for additional detailed information of sampling methods for water column physicochemistry.

Samples for inorganic nutrients were filtered immediately upon arrival at the laboratories at the Instituto del Mar del Perú (IMARPE) using 0.45 μm Sterivex filter (Merck). The subsequent analysis was carried out using a continuous flow analyzer (QuAatro AutoAnalyzer, SEAL Analytical) connected to a fluorescence detector (FP-2020, JASCO). The method for analyzing PO_4^{3-} followed the procedure outlined by Murphy and Riley (1962), while $\text{Si}(\text{OH})_4$ was analyzed according to Mullin and Riley (1955). NO_3^- and NO_2^- were quantified through the formation of a pink azo dye as established by Morris and Riley (1963) and additional corrections to all colorimetric methods was achieved with the refractive index method developed by Coverly et al. (2012). Ammonium concentrations were determined fluorometrically following the method of K  rouel and

175 Aminot (1997). Further methodological specifics and the respective limits of detection for each analysis can be found in Bach
176 et al. (2020).

177 **2.3 CHEMTAX Analysis**

178 Pigment samples were flash-frozen in liquid nitrogen directly after filtration and kept frozen on dry ice during transport to
179 Germany for extraction as described by Paul et al. (2015). Concentrations of extracted pigments were measured by reverse-
180 phase high-performance liquid chromatography (HPLC; Barlow et al., 1997) calibrated with commercially available standards.
181 The relative contribution of distinct phytoplankton taxa was calculated with CHEMTAX, a program for calculating the
182 taxonomic composition of phytoplankton populations (Mackey et al., 1996). Input pigment ratios specific to the Peruvian
183 upwelling system, determined by DiTullio et al. (2005) and further described by Meyer et al. (2017), were incorporated in
184 these calculations (see Bach et al., 2020).

185 **2.4 Flow Cytometry**

186 Samples (650 μ L) from each mesocosm were analyzed using an Accuri C6 (BD Biosciences) flow cytometer. The signal
187 strength of the forward light scatter was used to distinguish phytoplankton groups, in addition to the light emission from red
188 fluorescence of chl-a, and the light emission from orange fluorescence of phycoerythrin. Size ranges were constrained via
189 gravity filtration using sequential polycarbonate filters ranging from 0.2 to 8 μ m and the strength of the forward light scatter
190 signal. Additional details of this method can be found in Bach et al., 2017. In this study, we only report *Synechococcus* (0.8-3
191 μ m) counts (cells mL^{-1}) because it is the only phytoplankton group that was consistently selected to exhibit statistically
192 significant trends with IPLs likely due to the non-size fractionated nature of IPL sampling.

193 **2.5 Lipid Extraction and Analysis**

194 Intact polar lipids were extracted from glass fiber filters via a modified version (Wörmer et al., 2015) of the original Bligh and
195 Dyer Extraction method (Bligh and Dyer, 1959). Samples were extracted by ultrasonication a total of five times, with three
196 different extraction mixtures. Two extractions were performed using Dichloromethane:Methanol:Phosphate buffer
197 (aq) [1:2:0.8, v:v:v], adjusted to a pH of $\sim$ 7.4, followed by another two extractions using
198 Dichloromethane:Methanol:Trichloroacetic acid buffer(aq) [1:2:0.8, v:v:v], adjusted to a pH of $\sim$ 2.0. A final extraction was
199 performed with Dichloromethane:Methanol [1:5, v:v]. After each addition, samples were vortexed for 30 s, sonicated for 10
200 min, and then centrifuged for 10 min at 3000 rpm while kept at 10 $^{\circ}$ C. The supernatant of each extraction mixture was then
201 transferred to a separatory funnel where the organic fractions were washed and combined before solvent removal under a
202 gentle N_2 stream. Before analysis, the TLEs were resuspended in dichloromethane:methanol (9:1 v:v) and filtered through a
203 0.45 μ m polytetrafluoroethylene (PTFE) syringe filter.

204

Chromatographic separation and identification of IPLs was achieved using a Thermo Scientific UltiMate 3000 high-performance liquid chromatography (HPLC) interfaced to a Q Exactive Focus Orbitrap-Quadrupole high resolution mass spectrometer (HPLC-HRMS) via heated electrospray ionization (HESI) in positive mode, as described in detail in applications by Cantarero et al. (2020) and Flores et al. (2022). Briefly, a flowrate of 0.4 ml/min was applied to an Aquity BEH Amide column (150 mm, 2.1 mm, 1.7 μ m) using a gradient program first described by Wörmer et al (2013). All filtered TLEs were suspended in Dichloromethane:Methanol (9:1, v/v) prior to injection (10 μ L) on column. The following HESI conditions were applied; auxiliary gas temperature: 425 $^{\circ}$ C, capillary temperature 265 $^{\circ}$ C, spray voltage 3.5 kV, sheath gas flow rate: 35 arbitrary units (AU), auxiliary gas flow 13 AU, S-Lens RF level 55 AU. Samples were analyzed in full scan mode to obtain an untargeted screening (or lipidomic profile) of each sample, in addition to targeted MS/MS mode for compound identification via diagnostic fragmentation patterns (e.g., Sturt et al., 2004; Schubotz et al., 2009; Wakeham et al., 2012). IPLs were identified by their exact masses, polar head groups, the total number of carbon atoms and unsaturations in the core structure, and their retention times. While other studies have analyzed IPLs under both positive and negative ionization modes to determine the composition of individual fatty acid chains in the core lipid structures, we took advantage of the high resolution of the Orbitrap mass spectrometer to focus on the diversity of head group combinations with total carbon atoms and unsaturation only (Cantarero et al., 2020).

Quantification of IPLs was achieved using a combination of an internal standard added (2 μ g) to samples during extraction (C16 PAF, Avanti Lipids), in addition to an external calibration curve consisting of 17 standards representing different IPL classes (see Cantarero et al., 2020 for full details of all internal/external and deuterated standards). The intensity of each individual IPL identified in the HPLC-HESI-HRMS analysis was calibrated to a linear regression between peak areas and known concentrations of the same lipid class (or the most similar molecular structure) across a 5-point dilution series (0.0001–10 ng/ μ L). The detection limit, based on individual calibration curves, was determined to be 0.01 ng on column except for DGTS (0.001 ng) and DG, MG, and SQ classes (0.1 ng). Samples were analyzed across 3 separate analytical periods with weekly calibration curves to account for variation in the ionization efficiency of compounds over time. Overall, HPLC-ESI-HRMS is considered a semi-quantitative method due to changes in ionization efficiency of different IPL standards and environmental analytes. These changes are largely caused by differences in polar head group compared to the acyl chain length and degree of unsaturation (Yang and Han, 2011). Nonetheless, we investigate both relative (%) and absolute IPL abundances to compensate for the current analytical limitations in IPL quantification.

While we report IPL structural variations associated with different head groups and modifications in the core structure (i.e., unsaturation degree and carbon length of diacyl chains), we particularly focus on the former. Additionally, several IPLs thought to be absent in eukaryotic phytoplankton or far more abundant in bacteria and archaea (n = 34 in total) have been removed to facilitate the analysis of trends that are predominantly phytoplanktonic in origin. These compounds include PME/PDME (Phosphatidyl(di)methylethanolamine) and intact GDGTs (glycerol dialkyl glycerol tetraethers) classes. In addition, while we

lack detailed structural information of core individual fatty acids, their combined carbon atoms can be used to deduce short (i.e. < 14 carbon atoms per chain) or odd numbered fatty acid chains typically found in bacteria (Volkman et al., 1989; Volkman et al., 1998; Russell and Nichols, 1999; Jónasdóttir, 2019). Thus, compounds conventionally regarded as bacterial (34 in total) were also removed to minimize their impact on the analysis of predominantly phytoplanktonic IPLs (165 in total). We recognize that while this selection approach reduces the influence of non-eukaryotic lipids, we still cannot rule out an undetermined contribution from heterotrophic bacteria to the IPL pool in this experiment. A summary of these IPL classes and their acronyms is provided in Table 2.

2.6 Multiple Linear Regression

We performed multiple linear regression between 8 relevant environmental factors and 165 unique IPLs. With so many IPL-environmental factor pairs to analyze, we used multiple linear regression (MLR) for quick and easily digestible outputs. In each MLR model, the relative abundance of individual IPL molecules was employed as a response variable, with environmental factors serving as predictor variables. Additionally, phytoplankton abundances were included to account for their linear effects on IPL distributions. The rationale behind this inclusion lies in the understanding that variations in phytoplankton abundance may exert a proportional and predictable impact on the abundance patterns of specific IPLs.

We prioritized linear relationships with IPL relative abundances (% abundances) to emphasize changes in the proportions of phytoplankton lipids, rather than their absolute concentration and contribution to biomass. This approach enables us to distinguish compositional changes in the lipid pool from variability in total biomass production. Additionally, MLRs were constrained to focus on the most abundant IPLs in this system, defined as those constituting more than 2.5% to the total IPL pool. This restriction was implemented to reduce noise associated with low-abundance IPLs and enhance the robustness of analysis.

We chose to investigate linear relationships within individual depths (surface and subsurface) to focus on temporal changes within distinct environments, rather than comparing these two environments directly (see CART and Random Forest methods, below). IPLs and environmental factors were permuted to tabulate the regression coefficient for each IPL-environmental factor pair. Model coefficients were directly comparable due to centering and scaling of environmental and phytoplankton variables (see eq. 1) to linearize the relationship and better align with the model assumptions:

$$Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots + \beta_n X_n + \varepsilon \quad \text{eq. (1)}$$

where, Y is the dependent variable (or response), $X_1, X_2, \dots, X_n$ are the predictor variable (or independent variables), β_0 is the intercept (constant term), $\beta_1, \beta_2, \dots, \beta_n$ are the coefficients representing the magnitude and direction of the relationship between

the predictor variable and the dependent variable, and ε is the error term capturing the variability not explained by the model. Correlations in the MLR analysis were also controlled for false discovery rate following the procedure of Benjamini and Hochberg, (1995) to calculate adjusted p-values and by applying an alpha cutoff of 0.1.

2.7 Classification and Regression Tree (CART) and Random Forest

Classification and Regression Trees (CART; Breiman et al., 1984) are predictive machine learning algorithms that partition and fit data along a predictor axis into homogenous subsets of the dependent variable. Regression trees are used for dependent variables with continuous values, while classification trees for categorical values. Here, we apply regression trees to diagnose the environmental and biological variables that affect IPL concentrations by polar head group class. We limited the size of the tree splits via a “pruning” process, where less significant variable splits are removed as determined by a deviance criterion resulting in the best-fit tree with the least mean squared error. Additionally, predictor variables were selected in conjunction with random forest analyses, which ranks the variable contribution to the model performance.

Random forests are derived from bootstrapping of observational data, and the generation of many decision trees based on each bootstrap sample. We employed a total of 72 samples for the random forest analysis which was run separately for each major IPL class ($n = 7$) to identify the most important environmental predictors ($n = 12$) for each individual class of lipid. The random forest method utilizes bootstrap aggregation to generate and average numerous permutations of an out-of-bag score in predictive performance. This approach offers an effective methodology for analyzing high-dimensional data with limited sample sizes (Biau and Scornet, 2016). This method has been widely adopted across various disciplines within the water sciences (Tyralis, 2019), ecological and species distribution modeling (Luan et al., 2020), as well as in bioinformatics and high-throughput genomics (Chen and Ishwaran, 2012; Boulesteix et al., 2012). For a covariate vector, the predicted estimate of IPL class concentrations is an average of the many randomly configured decision trees with the same distribution. A randomly generated subset of decision trees is used to split each node, enabling reduced variance and correlation amongst individual trees, and ideally improving the accuracy of model predictions (Breiman, 2001). The random forest algorithm also allows for the ranking of predictor variables based on the prediction performance and is reported here as the predictor’s contribution in reducing the root mean square error (RMSE) in the model. For additional details on the random forest algorithm, please see Hastie et al. (2009). All CHEMTAX, flow cytometry, and physicochemical variables were included in the CART and random forest analyses as predictors. Regression tree (CART) splitting criteria are determined by evaluating the sum of squared deviations in all possible splits and selecting those that result in the greatest reduction of residual error. To prevent overfitting in the CART analysis, a pruning procedure is run to remove nodes that contribute little to the model accuracy based on a cost complexity measure. This procedure allows us to simplify the CART results and thus focus our interpretations on the most significant predictors of IPL headgroups only. In the Random Forest model, following the averaged cross validated accuracy estimates, we implemented a cutoff of 5% reduction in RMSE to eliminate variables that do not significantly reduce

the error of the model prediction. This cutoff allows us to focus our interpretation of only variables that contribute significantly to the out of bag predictor performance (For further details, refer to Supplementary Material, Note S1). CART and random forest-based predictions serve as ideal methods to explore environmental drivers of IPL distributions- not only because of their predictive performance- but also because of their non-parametric nature, their diagnosis of variable importance, and their ability to handle non-linear interactions and small sample sizes (Tyrallis et al., 2019). We do not employ these methods to predict the concentration of IPL classes, but rather to identify the primary environmental and biological drivers of change in IPL class concentrations, and potential interactions between environmental conditions and IPL remodeling amongst phytoplankton. Therefore, we focus our interpretations of both the CART and Random Forest analyses on only the most significant predictors and their relative order of importance in the model predictions.

3 Results

3.1 Oxygen and pH_T

Oxygen concentration in surface waters ranged between ~125 and 140 $\mu\text{mol/L}$ before the ODZ water addition in all 4 mesocosms (Fig. 2A). The concentration dropped slightly (~15 $\mu\text{mol/L}$) following the water addition and steadily increased to a maximum between ~185-220 $\mu\text{mol/L}$ by day 28. Day 30 marked a significant drop in oxygen concentration by 30-60 $\mu\text{mol/L}$ in all mesocosms with conditions stabilizing between 140 and 155 $\mu\text{mol/L}$ by day 50. The temporal variability in pH_T in surface waters showed a largely mirrored trend to that of oxygen content. Before ODZ water addition pH_T was 7.9-8.0, which dropped by ~0.2 immediately following the water addition. After a few days of relatively stable pH_T at ~7.8 in all four mesocosms, the pH_T increased significantly between days 18 and 28 reaching ~8.1-8.2. From days 30-40 the pH_T gradually decreased by approximately 0.2 in all mesocosms until increasing again (most notably in mesocosms 7 and 5) between days 42-50 to maxima ranging from 8.1-8.3.

The subsurface waters were more oxygen-depleted compared to the surface and showed an earlier onset of increasing oxygen concentration, beginning immediately after the ODZ water addition. All four mesocosms reached maximum O₂ levels (60-75 $\mu\text{mol/L}$) by day 13 and began decreasing markedly by day 16. The lowest concentrations (~15 $\mu\text{mol/L}$ O₂) were reached between days 30 and 34. Oxygen concentrations recovered slightly in the last 10-16 days of the experiment and were all within 30 ($\pm$ 10) $\mu\text{mol/L}$ of O₂. The subsurface waters, again, showed similar temporal patterns in pH_T as with O₂, except for mesocosm 7. The pH_T was variable (~7.60 $\pm$ 0.05) in the early portion of the experiment (days 10-16) but did not show a dramatic response to the OMZ water addition as was the case in the surface samples. The pH_T did gradually decrease from days 18-30, with pH_T minima in all four mesocosms reached on day 30 (~7.45-7.50). Day 30 also marked the beginning of a significant increase in the pH_T (~0.15-0.20) across all mesocosms with the pH_T steadily increasing to ~7.65-7.70 by day 50. Additional details on the carbonate chemistry of the mesocosms can be found in Chen et al. (2022).

3.2 Nutrient Concentrations

In mesocosm surface waters, the nutrients NH_4^+ , NO_3^- , NO_2^- , PO_4^{3-} , and Si(OH)_4 all showed consistent trends over time, with the highest concentrations occurring either just before (day 10) or immediately after (day 12) the ODZ water addition (Fig. 2B). Nitrogen species ranged between 1 and 3 $\mu\text{mol/L}$ during this early part of the experiment, while Si(OH)_4 ranged between 4 and 7 $\mu\text{mol/L}$, and PO_4^{3-} remained at $\sim 2 \mu\text{mol/L}$. The concentration of all nitrogen species dropped quickly to near minimum values by day 15, and typically remained $<0.5 \mu\text{mol/L}$ for the remainder of the experiment. PO_4^{3-} remained replete ($>1.5 \mu\text{mol/L}$ for the entirety of the experiment). Si(OH)_4 dropped to $\sim 3\text{--}4 \mu\text{mol/L}$ by day 15 and gradually increased in all 4 mesocosms by 1-2 $\mu\text{mol/L}$ until days 36-38, where concentrations gradually dropped to near minimum values of 3-4 $\mu\text{mol/L}$. There were periodic enrichments in NH_4^+ , most notably between days 40-50, as discussed in Bach et al., 2020.

The mesocosm subsurface water nutrients showed similar temporal patterns as the surface but were generally more enriched in all nutrient species. All nitrogen species decreased in the few days following the ODZ water addition, but at a more gradual pace than in the surface. Before and immediately after the water addition, NO_3^- was enriched at 4-6 $\mu\text{mol/L}$ and NH_4^+ between 2.5-4 $\mu\text{mol/L}$. NO_2^- concentrations were $\sim 0.5\text{--}1 \mu\text{mol/L}$ in mesocosms 6 and 7 during this period but remained low ($<0.3 \mu\text{mol/L}$) in mesocosms 5 and 8. Broadly speaking, the nitrogen species all reached minimal values ($<0.1 \mu\text{mol/L}$) by days 18-20, except for in mesocosms 6 and 8, where NO_3^- persisted at significant concentrations (0.2-2.3 $\mu\text{mol/L}$) until days 22 and 24, respectively. The remainder of the experiment was marked by typically depleted concentrations of all nitrogen species ($<0.1 \mu\text{mol/L}$) with occasional spikes in NH_4^+ reaching up to 1 $\mu\text{mol/L}$, particularly towards the end of the experiment. Subsurface PO_4^{3-} concentrations were similar to the surface, remaining around $\sim 2.0 \mu\text{mol/L}$ for most of the experiment which gradually decreased to concentrations of $\sim 1.5 \mu\text{mol/L}$ by the end of the experiment. Si(OH)_4 similarly decreased from maximum concentrations of 4.0-4.5 $\mu\text{mol/L}$ shortly after the ODZ water addition, and gradually decreased by $\sim 1 \mu\text{mol/L}$ over the course of the experiment.

3.3 Total Chlorophyll a

Chl-a concentrations were highly variable throughout the experiment, particularly in subsurface waters (Fig. 2C). Concentrations were generally more elevated at the surface compared to the subsurface, with values ranging $\sim 2.56\text{--}2.96 \mu\text{g/L}$ on day 10. In all mesocosms, the chl-a concentration increased to $\sim 7.94\text{--}14.01 \mu\text{g/L}$ after the ODZ water addition and through day 20. Day 22 showed a significant drop in chl-a concentrations to between 1.35 and 2.89 $\mu\text{g/L}$ in mesocosms 7, 5, and 8. Chl-a concentrations at the surface remained rather constant until days 36-40, where concentrations rapidly increased until maximum concentrations on days 48-50 (14.0-47.3 $\mu\text{g/L}$). In subsurface waters, chl-a concentrations were notably lower than in the surface and ranged between 0.58-0.84 $\mu\text{g/L}$ on day 10. After the ODZ water addition, chl-a concentrations increased slightly until day 14, but did not show consistent distributions amongst the 4 mesocosms afterwards; decreases were observed in mesocosms 6, 5, and 8, but an increase in mesocosm 7. Chl-a concentrations increased rapidly on day 22 in all 4 mesocosms

368 ranging between 4.03 and 8.44 $\mu\text{g/L}$. While remaining highly variable, the concentrations generally increased after day 24
369 with a notably abrupt increase between days 40-44 to near maximum values ranging between 2.58-11.3 $\mu\text{g/L}$.

370 3.4 Phytoplankton Community Composition

371 The CHEMTAX based phytoplankton community compositions demonstrated more variability in phytoplankton assemblages
372 in the subsurface samples than in the surface (Fig. 2C). Before the ODZ water addition on day 10, surface waters in all 4
373 mesocosms showed similar phytoplankton distributions with high relative abundances of Bacillariophyceae, referred to from
374 here on as diatoms (20-45%), Chlorophyceae (15-50%), and Dinophyceae, referred to as dinoflagellates (25-45%). These
375 distributions remained rather similar immediately after the water addition (day 12), but dinoflagellate contributions increased
376 in the following days and ranged from ~20% of the total chl-a pool to up to 75% by day 20. Of note, Cryptophyceae made
377 minor contributions to the chl-a pool aside from days 15-18 in mesocosm 6 where they contributed up to 25%. Dinoflagellates
378 largely dominated the chl-a pool for the remainder of the experiment with moderate blooms of diatoms between days 34-44 in
379 mesocosms 7 and 8.

380
381 Subsurface waters exhibited greater variability in the phytoplankton assemblages and a greater contribution of Chlorophyceae,
382 Cryptophyceae, *Synechococcus*, and diatoms to the chl-a pool than in the surface. Pre-addition waters (day 10) were dominated
383 by diatoms making up >75% of the chl-a pool. In the first few days after the ODZ water addition (days 12-15), the relative
384 abundance of Chlorophyceae increased to 25-45% in mesocosms 6, 5 and 8, whereas Cryptophyceae contributed between 5-
385 15% of the total chl-a. During this period, diatoms continued to dominate mesocosm 7 but decreased gradually from 65 to 15%
386 of the phytoplankton community. Notably, in mesocosm 6, Cryptophyceae contributed a moderate amount to the chl-a pool
387 beginning on day 15 (25%) and gradually increased to 40% by day 20. Mesocosms 7, 5, and 8 increased in Cryptophyceae as
388 well, but were limited to ~10-25%. Similar to surface waters, dinoflagellates dominated the chl-a pool by day 20 and
389 contributed 50-80% of the chl-a pool; however, there was considerably greater variability in dinoflagellate abundance after
390 day 20 in subsurface waters. Chlorophyceae remained a significant contributor for the rest of the experiment typically ranging
391 between 10 and 25% of the phytoplankton relative abundance. In mesocosms 7, 5, and 8 diatoms became the dominant
392 contributor, totaling ~50% of the phytoplankton between days 32 and 44. By the final days of the experiment (days 48-50),
393 dinoflagellates again made up most of the phytoplankton community (60-75%). Pelagophyceae, Prymnesiophyceae, and
394 Cyanophyceae (referred to here as *Synechococcus*), remained minor contributors throughout the entire experiment but showed
395 maximum contributions of <10% in days 10-16. Mesocosm 6 showed a minor contribution (<10%) of *Synechococcus* from
396 days 42-50, and from days 34-42 in mesocosm 7.

397 3.5 IPL Class Distributions

398 The IPL distributions throughout the study can be summarized by the relative abundances of different classes determined by
399 their polar head groups. IPL distributions were broadly consistent between mesocosms and treatments during the experiment

but showed significant differences between surface and subsurface waters (Fig. 2D). Glycolipids such as SQ, DG, and MG typically made up ~50-75% of the total IPL pool in the surface, whereas phospholipids such as PC and PG were dominant in the subsurface (~50-75%). The changes in IPL distribution from the surface samples over the course of the experiment were most apparent between days 10 to 12, 20 to 24, and 24 to 38. Day 10 of the experiment marks the sampling period immediately preceding the ODZ water addition and was the only day with a significant fraction of BLs (betaine lipids) in the total lipid pool (ranging from 25-30% in the surface samples). Mesocosms 6 and 8 showed a similar distribution between days 10 and 12 of the experiment with large contributions of SQs (~50%), BLs (~25-30%), and MGs (~20%), while mesocosms 7 and 5 showed considerably more PCs (~25-40%) at the expense of SQs (<10%). In the 2 weeks following the deep-water addition (days 12 to 24), we saw increasing (albeit variable) relative abundances of PGs (from 5 to 50%) and DGs (from 5 to 25%) in surface waters. MGs made up a considerably larger fraction during this time in mesocosm 6 (up to 50%), but were more moderate in mesocosms 7, 5, and 8 (typically ~25%), yet decreased in all mesocosms to <5% by day 24. The final sample days (38 and 50) showed a resurgence of glycolipid contributions, namely in SQs (25-55%) and MGs (5-25%), with moderate contributions of PGs (5-20%) and DGs (5-20%).

In the subsurface samples before the ODZ water addition, mesocosms 6, 7, and 8 showed moderate contributions of MGs (15-25%) which was lower in mesocosm 5 (<5%). There were higher contributions of SQs in mesocosms 6 and 8 (50% and 35%, respectively) than in 7 and 5 (20-25%). Instead, mesocosms 7 and 5 showed a greater contribution of PCs (45-55%). All 4 mesocosms demonstrated some component of PGs ranging from 5-20% of the total IPL pool. The 2 weeks following the ODZ water addition showed highly variable fluctuations between PCs and PGs as the dominant IPL classes, with the contributions of glycolipids (25-40%) and betaine lipids (<5%) remaining consistent. Of note, day 24 marked a consistently low contribution of PCs, which persisted until the end of the experiment. The final sample days (38 and 50) showed similar contributions between MGs, SQs, and PGs that together dominated the total lipid pool.

3.6 Multiple Linear Regressions

We found statistically significant ($p < 0.05$) linear relationships between the relative abundance of individual IPLs and environmental factors (Figure 3). In surface waters, pH_T showed several significant responses towards DGs, MGs, PGs, PEs and SQs. Classes DG, MG, PE and PG showed negative linear relationships with pH_T . SQs (specifically SQ-32:0), however, showed a strong positive linear response to pH_T . In the subsurface, pH_T only imparted significant linear effects towards SQs with a strong negative linear correlation.

The temperature of surface waters showed predominantly positive regression coefficients with several PG, MG, and DG molecules, with inconsistent correlations found in SQs. Whereas some PEs showed significant linear relationships with temperature, their regression coefficients were near 0. Similarly, in the subsurface only 1 IPL structures (SQ 34:3) showed a significant positive linear response but with a regression coefficient near 0.

Oxygen concentrations showed few linear correlations at the surface and were limited to two BLs (negative) and 1 PG (positive). In the subsurface, the strongest linear correlations were found between oxygen and PGs and SQs (negative).

Nutrient concentrations showed many significant linear responses with regression coefficients of higher magnitude (up to ± 8) and adjusted R^2 values of up to 0.5. The concentration of various forms of inorganic nitrogen had largely positive linear responses to BLs and PCs in both the surface and subsurface, with strongly negative responses in PGs and subsurface MGs. Among the SQs, many individual molecular species, with different fatty acid chains, responded linearly to all forms of inorganic N, the signs of these relationships were negative but in the surface, and generally positive in the subsurface (aside from NO_2^-). DGs showed only one positive linear response in the subsurface. PEs showed little response to inorganic nitrogen concentrations (regression coefficients near 0). PO_4^{3-} concentrations showed only a few significant responses from BLs (negative) and PGs (slightly positive) in the surface. However, in the subsurface, PO_4^{3-} showed strong linear responses (negative) to several MG and PG molecules, and strong positive relationships with several PCs.

Light showed few linear responses amongst IPL relative abundances. At the surface, the strongest relationships were amongst PG (positive) and SQ (mixed signs) and to a slight degree BLs (negative). In the subsurface, only one SQ with a significant linear relationship was noted and with a weak regression coefficient, indicating a range of light saturation with little to no linear effect on IPL distributions.

3.7 CART and Random Forest

All the predictive tree-based models showed improved performance with the inclusion of environmental variables (Figs. 4-7). The CART decision trees iteratively identified the key biological and physicochemical variables producing the best performing model in the prediction of IPL concentrations. The random forest analysis complements these best fit decision trees by calculating the % reduction in the root mean square error (RMSE) associated with each variable. In conjunction, these two analyses highlight the most impactful variables in predicting the concentrations of a given IPL class. Overall, model performance amongst each IPL class can be compared by the strength of the correlation coefficients between observed and predicted concentrations, and by the magnitude of the RMSE (Figs. 4-7).

Amongst several IPL classes (i.e., SQs, DGs, and BLs), pH_T was consistently a significant contributing variable to IPL concentrations, as demonstrated by both the CART and random forest analyses (Fig. 4A-D, and Fig. 5). Notably, pH_T was identified as the most important variable amongst the best fit decision trees for both SQs and DGs, as well as the random forest model for SQs. Oxygen concentration was also frequently identified as a major contributing variable to model performance with MGs, DGs, PEs, PCs, (Figs. 4E-H, 5A-D, 6E-H, and 7A-D, respectively) and moderately important in SQs (Fig. 4A-D).

Temperature was selected as the most important variable in PE predictions (Fig. 6E-H), and a major contributing variable in SQ, MG, BL, PG, and PC predictions (Figs. 4-7). Various forms of biologically available nitrogen were also important in the prediction of MG (Figs. 4E-H), BLs (NH_4^+ ; Figs. 5E-H), PE (NH_4^+ ; Figs. 6E-H), and PG (NH_4^+ and NO_2^- ; Figs. 6A-D). PO_4^{3-} concentrations showed significant contribution to model performance amongst BLs (Figs. 5E-H, denoted as Si:P ratios), SQs, MGs, PGs, and PEs (Figs. 4A-D, 4E-H, 6A-D, and 6E-H, respectively). Finally, light availability demonstrated secondary but significant importance in the prediction of SQs, MGs, PGs, PEs, and PCs (Figs., 4A-D, 4E-H, 6A-D, 6E-H, and 7 respectively).

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Variables indicative of biological abundance were also identified as highly impactful to model performance, with chl-a concentration showing a significant contribution to all lipid classes except BLs (Figs. 4-7). Individual phytoplankton abundances also showed to be important predictive variables such as Cryptophyceae abundances for BLs, dinoflagellate abundances for DGs and SQs, *Synechococcus* abundances for DGs, PCs, PEs, and PGs, and diatoms for PCs.

3.8 Water Treatments

The experiment consisted of applying two different treatments to the mesocosms, aimed at exploring the varying impacts of upwelling of ODZ waters with contrasting geochemical properties. The first treatment saw the introduction of water from a coastal area (station 1) with ODZ waters with a very low N:P ratio (0.1; see Table 1, and Fig 2B) into the mesocosm. The second treatment performed the same process, but with ODZ water from an offshore area (station 3) with a low N:P ratio (1.7; Table 1, and Fig 2B). Despite the different chemical signatures of the added water-masses, the resulting nutrient stoichiometries within the mesocosms were similar in between both treatments, likely due to both dilution effects and the time passed between water collection and addition (see Bach et al., 2020). We see largely similar responses in the IPL distributions, as well as in other biogeochemical variables between these two treatments; therefore, we largely focus our discussion on the temporal variation and differences between surface and subsurface environments in these analyses.

4 Discussion

4.1 Biological Abundances as Drivers of IPL Distributions

We expect phytoplankton abundances to exert first order control on the production and distribution of IPLs in this experiment. The majority of the detected molecules have been demonstrated to be chemotaxonomic biomarkers of planktonic biomass (Sturt et al., 2004, Schubotz et al., 2009, Wakeham et al., 2012; Van Mooy and Fredricks, 2010; Cantarero et al., 2020). Previous work in the Humboldt Current System shows that the ratio of total IPLs to POC is high at the chlorophyll maximum and the composition of IPLs found in these surface waters are consistent with predominantly phytoplanktonic biomass (Cantarero et al., 2020). In this mesocosm experiment, the depths of the chlorophyll maximum and oxycline are compressed into a 20 m water column which likely drives a greater contribution of phytoplanktonic IPLs in ODZ waters than would be expected in the natural environment. While we suggest that most of the biomass (and IPL content) measured at these high

chlorophyll depths is likely derived from phytoplankton, we cannot completely isolate nor quantify the contribution of bacterial biomass to the total IPL pool.

Most IPL classes demonstrate phytoplankton abundances as a primary or major predictor in the CART and random forest analyses, such as dinoflagellates in the prediction of SQs (Figs. 4A, C), DGs (Figs. 5A, C), and PE (Fig. 6G), *Synechococcus* in the prediction of PGs (Fig. 6A, C), PEs (Fig. 6E, G), and PCs (Fig. 7A, C), and chl-a in the prediction of every IPL class barring BLs (a mostly minor IPL class in this experiment). Chl-a appears most important in highly abundant IPL classes as an indicator of overall photosynthetic productivity, and dinoflagellates dominate the overall phytoplankton biomass for almost the entirety of the experiment post ODZ water addition. *Synechococcus* demonstrates covariance with the total phytoplankton biomass, yet remains a relatively minor phytoplankton class. We suggest that the prevalence of biological sources as IPL predictors in the decision tree analyses generally indicates the variability in total phytoplankton biomass throughout the experiment.

Among phytoplankton, glycolipids MG, DG, and SQ are predominantly found in thylakoid membranes, whereas phospholipids PC and PE, as well as BLs are structural components in the cell membrane lipid bilayer (note PG is found in both; Guschina and Harwood, 2013). It is important to recognize that the relative proportions of IPLs vary in different phytoplankton classes (Harwood and Jones, 1989; Wada and Murata, 2009), and that many of these lipids, in particular phospholipids, can also be derived from heterotrophic bacterial biomass (Popendorf et al., 2011). Thus, the overall community composition is expected to be a major driver of IPL distributions in this system. However, given the prevalence of algal biomass and the steps taken to minimize bacterial contributions to the IPL pool (see section 2.5), we focus our analysis on that of phytoplanktonic dynamics. We recognize that both phytoplankton abundances and the total planktonic community composition play a major role in the distribution of IPLs. Thus, while the data presented here may refine the phylogenetic association between biological sources and IPLs in marine systems, our main aim is to explore the role of multiple environmental forcing as an additional control on IPL distributions. The following sections focus on the evidence for direct environmental influence on IPL remodeling amongst the phytoplankton community and the potential implications of these physiological responses to broader aspects of ocean biogeochemistry.

4.2 Environmental Variables as Drivers of IPL Remodeling

Since community composition can change concurrently and/or in response to environmental conditions, we employed two distinct strategies to isolate the role of lipid remodeling as a physiological response to environmental forcing only. Firstly, the MLRs which subtract the variability explained by phytoplankton abundance (e.g., CHEMTAX results) in pairwise correlations between lipid abundances and environmental variables. Secondly, the decision tree analyses (CART and Random Forest) which rank variables by their impacts on the model performance.

530 Across virtually every major IPL class common to eukaryotic phytoplankton we see evidence of environmental conditions
531 exerting significant control on polar headgroup distributions in the MLRs (Fig. 3). Similarly, nutrient concentrations, pH_T ,
532 temperature, O_2 concentration, and light availability are consistently identified as statistically important variables in the
533 prediction of IPL head groups in both the CART and Random Forest analyses (Figs. 4-7). In addition, high level comparisons
534 in the relative abundances of IPLs and phytoplankton groups (Figs. 3C, D) suggest that certain environmental conditions may
535 be associated with major shifts in IPL distributions.

536

537 An important distinction between the MLRs and the decision trees is that the MLRs are calculated within individual depth
538 environments (surface and subsurface) to explore statistically significant linear relationships between abundant IPL molecules
539 and changing environmental conditions. On the other hand, the decision trees explore the predictive power of physicochemical
540 differences between the surface and subsurface environments on IPL distributions. The results of these two analyses are meant
541 to be complimentary, in that they focus on the differences in environmental conditions between water depths as well as the
542 temporal development of conditions within a given depth over the course of the experiment.

543 4.2.1 Nutrient Availability

544 Nutrient limitation amongst phytoplankton leads to transitions in cellular activity, from the biosynthesis of
545 growth/reproduction cellular components such as cell membranes to energy storing molecules (Guschina and Harwood, 2013,
546 Zienkiewicz et al., 2016). Nitrogen, an essential nutrient in photosynthesis and the biosynthesis of proteins/enzymes and
547 nucleic acids, is typically acquired by marine phytoplankton through inorganic nitrogen species such as NO_3^- and NH_4^+ . Some
548 phytoplankton can also utilize organic nitrogen sources (Bronk et al., 2007), whereas diazotrophic cyanobacteria can fix
549 dinitrogen gas into bioavailable nitrogen. The coastal region of the ETSP is typically considered to be seasonally co-limited
550 by light, N, and Fe (Messié and Chavez, 2015). However, along the Peruvian shelf Fe concentrations are elevated compared
551 to offshore waters (Hutchins et al., 2002; Browning et al., 2018), and is not considered a limiting source in this mesocosm
552 experiment (Bach et al., 2020). In our mesocosm systems, the inorganic N:P ratio ranged between 0.13 and 4.67, with higher
553 inorganic N in subsurface waters compared to the surface, and with a N:P minimum reached by day 20. Bach et al., (2020)
554 noted that a week after the ODZ water addition, increases in the particulate organic carbon to biogenic silica ratios coincided
555 with low inorganic N and high Si(OH)_4 concentrations, suggesting a N-limited system. Thus, we consider our system to be
556 overall nitrogen limited with varying degrees of severity throughout the course of the experiment. This N-limitation is also
557 reflected in the transition from predominantly diatoms, Chlorophyceae, and Cryptophyceae to a dominance of mixotrophic
558 dinoflagellates approximately 4-6 days after the initial ODZ water treatment (Bach et al., 2020). Such shifts are consistent with
559 the ecological advantage that dinoflagellates exhibit under N-limiting conditions as they can extract nitrogen from the dissolved
560 organic nitrogen (DON) pool (Kudela et al., 2010) as well as from heterotrophy (Smalley et al., 2003).

561

562 In our study, all four mesocosms experienced limitations in inorganic nitrogen (as low as 0.24 $\mu\text{mol/L}$) and consistently high
563 concentrations of PO_4^{3-} (ranging from 1.3 to 2.3 $\mu\text{mol/L}$) throughout the entire experiment. Random forest analysis shows that
564 the inorganic N concentration is an important predictor in the abundance of BL, PG, and MG (Figs. 5G, 6A, and 4E,
565 respectively). The MLRs also indicate that many individual molecules from nearly every IPL headgroup have significant linear
566 correlations with inorganic N concentrations (Fig. 3). Of note, the distributions of several abundant PCs and BLs with N in the
567 headgroup structure are consistently positively correlated with inorganic N species. Whereas other non-N-bearing IPLs are
568 generally negatively correlated with inorganic N concentration (PG, SQ and MG) meaning that they are proportionally more
569 abundant under more severe N-limitation.

570
571 Under P limitation, phytoplankton are known to substitute non-phosphorus-containing glycolipids for phospholipids and
572 reallocate the liberated P for other cellular demands (Van Mooy et al., 2009). We hypothesize that non-N-containing
573 glycolipids and phospholipids may similarly be substituted for IPLs such as PCs and BLs as a mechanism for alleviating
574 cellular N demand in low inorganic N conditions. Both PC and BL are found in extra-chloroplast membranes (Kumari et al.,
575 2013), whereas IPLs found in thylakoid membranes such as MG, PG, and SQ are essential to the photosynthetic machinery.
576 Indeed, the average ratio of total IPLs/chl-a is up to three times higher at depth than in surface waters (Fig. S5), possibly
577 pointing towards a reduced proportion of membrane lipids among phytoplankton subject to environmental stressors such as
578 nutrient limitation (likely in addition to oxygen availability, temperature, and light levels). We note that at least a fraction of
579 this trend could also be explained by the contribution of IPLs from heterotrophic bacteria.

580
581 More generally, nutrient limitation can cause phytoplankton to accumulate highly concentrated stores of energy in the form of
582 triacylglycerols (TAGs) through the activation of multiple biosynthetic pathways (Zienkiewicz et al., 2016). These include
583 synthesis via acyl units donated from phospholipids via the PDAT (phospholipid:diacylglyceroltransferase) enzyme (Dahlqvist
584 et al., 2000), or other chloroplast membranes, as demonstrated in the homologous enzyme Cr-PDAT (Yoon et al., 2012), as
585 well as several DGAT(diacylglycerol:acyl-CoA acyltransferases; Li et al., 2012; Li-Beisson et al., 2019). These enzymes
586 represent significant pathways for TAG accumulation (Popko et al., 2016; Gu et al., 2021), are sensitive to N availability
587 (Yoon et al., 2012; Li et al., 2012), and their encoding genes have so far been identified in green algae, diatoms, and heterokonts
588 (Zienkiewicz et al. 2016). Because our dataset does not include TAG production, further work on this aspect could reveal
589 whether the proportional changes in dominant phytoplanktonic IPLs correlated to N availability are also associated with TAG
590 synthesis, and if so, determine whether recycling of membrane lipids is a significant contributor to these observed community-
591 level distributions.

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4.2.2 pH_T and Inorganic Carbon Availability

Coastal upwelling zones are characterized by low pH_T subsurface waters associated with ODZs, where high fluxes of organic substrates sustain enhanced microbial respiration and the accumulation of CO₂ (Capone and Hutchins, 2013). Thus, we explore evidence for membrane lipid remodeling amongst phytoplankton as a physiological response to varying pH_T. We see evidence of pH_T imparting a potential control on the composition of IPL head groups, particularly amongst SQs and DGs, as noted by the high importance rankings in both the CART and Random Forest analyses (Figs. 4A,C and 5A,C). This likely represents the relatively high abundance of these glycolipids in the surface samples where the pH_T is 0.2-0.6 higher than at depth. The MLRs show several negative correlations between MG, PG, and SQ molecules containing unsaturated or polyunsaturated fatty acids with pH_T (Fig. 3). The observed increased proportion of unsaturated IPLs at lower pH_T is most apparent in surface waters where the variability in pH_T is greatest (± 0.2). It has been suggested that lower pH_T can induce greater proportions of saturated fatty acids as a mechanism to reduce membrane fluidity and prevent high proton concentrations in the cytoplasm (Tatsuzawa et al., 1996). However, this response remains limited to more extreme pH_T ranges (e.g., ~ 1 -10), suggesting that other environmental factors (e.g. nutrient availability) are overprinting the potential impacts of pH on fatty acid saturation.

Rather than a direct consequence of modest changes in pH_T on algal membrane fluidity, the observed changes in fatty acid profiles may be in part a response to the available forms of inorganic carbon for photosynthesis. Following the ODZ water addition, DIC (dissolved inorganic carbon) and pCO₂ rapidly declined within a few days (Chen et al., 2022) due to high productivity. In natural waters, pCO₂ maxima occur in ODZ waters where respiration rates are high (Vargas et al., 2021). Lower pCO₂ at the surface may be a limiting factor for photosynthesis and growth; higher pH_T in marine settings indicates a reliance on active transport of HCO₃⁻ for carbon fixation as opposed to a passive diffusion of CO₂ (Azov et al., 1982; Moazami-Goudarzi et al., 2012). Enrichment of pCO₂ has also been observed to induce an increased proportion of unsaturated fatty acids in microalgae (Morales et al., 2021), which may explain the negative correlation between the proportion of certain unsaturated glycolipids (as well as PGs) and pH_T (high pCO₂; see Fig. 3).

As mentioned above, phytoplankton can employ membrane lipids as substrates for TAG accumulation in phytoplankton under environmental stress. While nutrient limitation is often considered the primary regulator of TAG production, culture experiments also point toward the importance of pH and inorganic carbon availability (Guckert and Cooksey, 1990; Gardner et al., 2011), with different responses between a model diatom (*Phaeodactylum tricornutum*) and chlorophytes (CHLOR1 and *Scenedesmus sp.* WC-1), which is potentially related to individual carbon concentrating mechanisms (Gardner et al., 2012). For instance, under nitrogen/phosphorus limitation, TAG production can be promoted if the supply of inorganic carbon is abundant (Peng et al., 2014). TAGs can also be synthesized and accumulated when inorganic carbon is limited by invoking the recycling of IPLs such as glycolipids and phospholipids (Peng et al., 2014). Thus, the higher proportions of SQs and DGs in the high pH/low CO₂(aq) surface waters of our experiment could be in part related to their recycling under CO₂ limitation.

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Overall, our results are consistent with other experimental data in that pH_T impacts the distribution of IPL head groups amongst certain phytoplankton groups. For instance, the relatively high glycolipid abundances in surface samples with higher pH_T (most notably SQ and DG, see Fig. S1) are likely not related to direct effects on membrane fluidity, but rather to the availability of inorganic carbon and its effect on the recycling of phospholipids (as well as MGs), potentially for TAG synthesis. While our results are consistent with the so far observed IPL substrates for TAG synthesis (Dahlqvist et al., 2000; Yoon et al., 2012), additional analyses of TAG concentration and their fatty acid compositions amongst the phytoplanktonic community would aid in tracing the extent of membrane lipid degradation as a source of acyl units, as well as to trace the location of these biosynthetic pathways in the cell. Such work would aid in determining the extent to which TAG synthesis and IPL recycling relegates phytoplankton classes to certain depths of the water column based on the combined effects of nutrient availability and pH_T (amongst other variables; e.g. light, temperature, O_2 .)

639 **4.2.3 Oxygen**

640 Despite the harvesting of light energy during photosynthesis, photosynthetic organisms also rely on respiration for growth and
641 free radical scavenging under both light and dark conditions (Raven and Beardall, 2003). Specifically, the availability of O_2
642 (particularly during the dark phases or in low-light environments) influences the activation of metabolic responses, such as
643 fermentative metabolism and acetate utilization (Yang et al., 2015). Differences in dark respiration rates relative to light-
644 saturated photosynthesis among algae may confer advantages under varying oxygen availability (Geider and Osborne, 1989).
645 Thus, the ability of some organisms to perform lipid remodeling in response to oxygen stress may partially shape the
646 composition of the phytoplankton community.

647

648 In our analysis, the Random Forest models indicate a significant impact of O_2 concentration in the prediction of nearly all IPL
649 class abundances (Figs. 4-7). This pattern appears to be largely driven by differences in IPL distributions between surface
650 (oxygenated) and subsurface (hypoxic; $< \sim 1.4$ ml/L as defined by Naqvi et al., 2010) waters. Glycolipids (MG, DG, and SQ)
651 make up on average 28% more of the IPL pool in the oxygenated surface waters (~ 125 - 220 $\mu\text{mol/L O}_2$) compared to oxygen
652 deficient subsurface waters (~ 15 - 75 $\mu\text{mol/L O}_2$). The MLRs, however, show no significant relationships between individual
653 glycolipid relative abundances and O_2 concentration in surface samples (Fig. 3); this is possibly due to surface waters remaining
654 well oxygenated throughout the experiment. While several IPL moieties found in BLs, SQs, and PGs demonstrate mostly
655 negative linear relationships (Fig. 3), we suggest that the most prominent and consistent relationships are driven by a major
656 shift from oxic to hypoxic conditions (i.e., surface vs subsurface) as opposed to a sensitivity to variable O_2 concentrations.

657

658 Anaerobiosis amongst green algae has been demonstrated to impact lipid production, with significant reductions (by nearly
659 50%) in polar lipid content and concomitant increases in fatty acids (Singh and Kumar, 1992). Gombos and Murata (1991)

found that the cyanobacterium *Prochlorothrix hollandica* experienced a significant reduction in the relative abundance of MGs that coincided with moderate increases in SQs, DGs, and PGs under low oxygen conditions. Furthermore, culture experiments of anaerobically grown *Chlamydomonas reinhardtii* resulted in both decreased membrane lipid yields (most notably amongst MGDGs and DGDGs; by > 50%) and an accumulation of TAGs (Hemschemeier et al., 2013). It has been noted that oxygen stress appears to induce the degradation of fatty acids (~30% reduction under dark/anaerobic conditions), mostly amongst unsaturated fatty acids commonly found in MGDG and DGDG membrane lipids (16:4 and 18:3) used for TAG assembly (Liu, 2014; Hemschemeier et al., 2013). Glycolipids (i.e., MG and DG) appear to serve as important substrates for TAG production under low oxygen conditions; however, beta-oxidation of fatty acids requires oxygen to contribute to the degradation of acyl groups, potentially explaining why membrane lipid degradation is attenuated under more severe hypoxia (Liu, 2014). This physiological response to low oxygen conditions in subsurface waters may explain the relatively high abundance of glycolipids in well oxygenated surface waters.

Dinoflagellates have been shown to exhibit particularly high dark respiration to light-saturated photosynthetic rates as compared to diatoms, Chlorophyceae, or most notably cyanobacteria (Geider and Osborne, 1989), possibly pointing towards a greater sensitivity to O₂ concentrations. In addition to other environmental conditions, the greater relative abundance of Chlorophyceae, diatoms, and Cryptophyceae in the oxygen-deficient subsurface waters may reflect reduced respiration rates amongst these algae. Differences in the proportion of glycolipids MG and DG amongst different algae, and their relative ability to recycle them under oxygen stress, may play a prominent role in their individual tolerances to oxygen limitation.

Higher proportions of glycolipids in surface waters may also be due to enhanced rates of microbial degradation under oxic conditions, which may be 2-4 times faster than under anoxic conditions, as tested in microcosm experiments (Ding and Sun, 2005). Relatively labile glycolipids can accumulate in the dissolved organic carbon pool (Gašparović et al., 2013). This observation aligns with the slower breakdown of SQs compared to phospholipids observed in IPL degradation experiments under aerobic conditions (Brandsma, 2011). This accumulation process, however, is unlikely in regions of the water column with large numbers of active living cells, and highly active bacterial degradation. In fact, the distributions of IPLs across the ODZ of the ETSP indicate minor contributions of exported IPLs to greater depths, suggesting high surface recycling (Cantarero et al., 2020). However, specific experimental observations encompassing oxygen gradients ranging from well-oxygenated to fully anoxic conditions are necessary to derive more robust conclusions.

4.2.4 Temperature

Phytoplankton have been shown to respond variably to high growth temperatures depending on their individual tolerances (Huertas et al., 2011). Photosynthesis is considered the most heat-sensitive cellular function in photoautotrophs (Berry and Björkman, 1980). In this section we discuss the potential lipidomic responses to heat stress within the IPL distributions of the

693 phytoplanktonic community. Temperature fluctuations affect membrane fluidity, a phenomenon commonly controlled by fatty
694 acid desaturases (Sakamoto and Murata 2002) that catalyze the production of unsaturated/saturated fatty acids to
695 increase/decrease membrane fluidity, respectively (Lyon and Mock 2014). We did not see evidence for temperature effects on
696 the degree of unsaturation in our data (see Fig. S2). This is likely due to the overall narrow temperature range observed during
697 the experiment (17.3 - 21.6 °C), which mirrors the natural variability observed in Callao (average monthly ranges ~16.6-19.6
698 °C from 2017 to 2019 (Masuda et al., 2023).

699

700 Despite the restricted temperature range and its lack of impact on the unsaturation degree of core lipids, the random forest
701 analyses identified temperature as a significant variable in predicting all IPL classes based on their polar head groups aside
702 from DG and PE (Figs. 4-7). This suggests that the response to temperature may vary among different IPLs classes. The MLRs
703 indicate consistently positive relationships between several glycolipids and temperature, suggesting a potential physiological
704 compensation via membrane compositions for higher temperatures. Gašparović et al. (2013) noted an accumulation of
705 glycolipids at temperatures >19 °C in the northern Adriatic Sea, particularly from cyanobacterial synthesis of MGs. The
706 sensitivity of photoautotrophs to thermal stress was also explored by Yang et al. (2006), who showed that DGs and MGs both
707 increase the thermal stability of photosystem II, while phospholipids significantly decrease it. Experiments with a wild-type
708 and mutant *Chlamydomonas reinhardtii* have shown that SQs are an essential component of thylakoid membranes to maintain
709 stability under heat stress (Sato et al., 2003), although at considerably more extreme temperatures (41 °C). Heat stress has also
710 been linked to the production of TAGs (Elsayed et al., 2017; Fakhry and El Maghraby, 2015), which can draw acyl units from
711 degraded membrane lipids (Holm et al., 2022).

712

713 Interestingly, the relative abundance of DGs in our experiment shows the most prominent (R^2 of 0.35-0.44), and statistically
714 significant ($p < 0.05$), linear relationship with temperature (see Fig. S3A). While temperature was not identified as an important
715 variable to the prediction of DGs in either decision tree analysis, this may be due to other covariates, such as pH and O_2 ,
716 masking the effect of temperature. Our results indicate that phytoplankton may either produce DGs in greater abundance to
717 alleviate thermal instability in photosystem II, or preferentially degrade other thylakoid membranes (i.e., PGs, SQs, or MGs)
718 in response to heat stress, leaving the remaining IPL pool relatively enriched in DGs. While several individual MG and SQ
719 molecules did demonstrate linear responses to temperature (Fig. 3), other stressors such as N availability, pH, and light levels
720 may confound the effects of temperature in the overall abundance of these lipid classes.

721 4.2.5 Light Availability

722 Bach et al. (2020) considered the overall productivity in these mesocosm experiments to be co-limited by N and light
723 availability, which have been identified as key limiting factors in eastern boundary upwelling systems (Messié and Chavez,
724 2015). In our experiment, initial high biomass productivity led to self-shading effects that significantly reduced the PAR (Bach
725 et al., 2020). While the maximum photon flux densities measured at noon over the course of the experiment were ~500 – 600

$\mu\text{mol m}^{-2}\text{s}^{-1}$, only ~5-22% and < 4% were measured in surface (~2 m) and subsurface (17 m) waters, respectively, from all four mesocosms. While both depth-integrated samples in our experiment are likely from light-limited planktonic communities, most PAR values (> 15% i.e., 75-132 $\mu\text{mol m}^{-2}\text{s}^{-1}$) in surface waters still demonstrate sufficiently high levels that can affect lipid production and accumulation (Gonçalves et al., 2013) especially under the combined effects of N-deprivation (Yeesang and Cheirsilp, 2011; Jiang et al., 2011).

Light levels are identified by the Random Forest analyses to be a significant variable in the prediction of MGs, SQs, PGs, and PCs (Figs. 4, 6, and 7). While few studies have explored the direct effects of irradiance levels on IPL headgroup distributions, experimental cultures of *Tichocarpus crinitus* found increased IPL production of SQs, PGs, and PCs amongst shade-grown algae (Khotimchenko and Yakovleva, 2005). In *Tichocarpus crinitus* cultures, MGs, one of the most abundant thylakoid lipids, show decreased abundances under low light conditions (Khotimchenko and Yakovleva, 2005). Our results show that both MGs and SQs are present in greater relative and absolute abundances in the surface waters where light is less limited or potentially inhibitive at times, suggesting an impact on thylakoid lipid compositions to maintain efficient photosynthetic rates under changing light conditions. MGs have been shown to provide important photoprotective and antioxidant mechanisms in diatoms (Wilhelm et al., 2014), potentially explaining their higher abundance in the surface waters and the significance of light in the Random Forest models. While the exact function of SQs in thylakoid membranes remains unclear, they have been observed to act as an antagonist to the aggregating action of MGs (Goss et al., 2009), leading to a disaggregating effect on light harvesting complexes (Wilhelm et al., 2014). The latter potentially reflects planktonic responses to variable PAR (5-22%) in the surface via adjustment of the proportions of SQs and MGs in the thylakoids (Gabruk et al., 2017; Nakajima et al., 2018).

The PC lipid class, primarily found in extrachloroplast membranes (Mimouni et al., 2018), consistently exhibits greater relative abundance in the subsurface compared to surface waters. We suggest that light levels at depth may play a role (albeit a secondary one, see Fig. S3B) in non-chloroplast IPL production when algal cells are configured for rapid growth/reproduction. Conversely, under higher light as well as under the combined effects of DIC and N limitation at the surface, algal cells are more prone to survival responses such as TAG production at the expense of IPLs. PG can be found in extrachloroplast membranes but is also an essential component of photosystems I and II, with important roles in electron transport processes (Sakurai et al., 2003; Wada and Murata, 2009; Kobayashi et al., 2017). The overall abundance of PGs contributing to the total IPL pool may be influenced by varying production of certain non-chloroplast membranes under lower light conditions, and other thylakoid membranes under high light conditions, thus possibly explaining the highly variable overall contribution of PGs to the IPL pool. Additional analyses of specific fatty acid structures from intact PGs may illuminate sources within the cell, i.e., thylakoid PGs and extrachloroplastic PGs.

Light intensity has also been shown to have contradictory impacts on fatty acid composition, with some analyses observing increased unsaturation under high light intensity (Liu et al., 2012; Pal et al., 2011). Others observed that excess light energy

induces synthesis of saturated (SFAs) or monounsaturated (MUFAs) fatty acids in several algal species to prevent photochemical damage, or the synthesis of polyunsaturated fatty acids (PUFA) for the maintenance of photosynthetic membranes under low light conditions (Khoeyi et al., 2012, Fabregas et al., 2004, Sukenik et al., 1993; Orcutt and Patterson, 1974). However, we see limited evidence in this experiment of a direct correlation between light intensities and fatty acid composition in the multiple linear regressions (Fig. 3). This may be related to differences in optimal irradiance levels between algal groups overlapping to confound any clear patterns.

Given that the normalized PAR in surface waters remains <25%, and that light appears as a variable of significant but secondary importance in the random forest models, our results suggest that under these experimental conditions, low or variable light availability may amplify the effects of other environmental stressors (i.e., nitrogen availability, oxygen concentration, temperature, and inorganic carbon availability). The effects of self-shading from high biomass production early in the experiment, when greater nutrient availability was high, likely contributed to the markedly different IPL distributions observed between surface and subsurface waters. As a result, the fluctuations in PAR may play a role in the observed variations in the relative proportions of prevalent thylakoid membrane IPLs, including MGs, SQs, and PGs.

4.3 Implications of IPL Remodeling in Phytoplankton Membranes

We expect the ongoing changing conditions in the ETSP, such as rising temperatures, ocean acidification, expanding ODZs, and shifts in upwelling-driven nutrient supply, to instigate multiple physiological responses amongst phytoplankton communities. IPL remodeling and the reallocation of resources among algae are likely to have cascading effects on the composition of phytoplanktonic communities, the nutritional value of algal biomass to higher trophic levels, and the cycling of organic carbon and nitrogen in the upper ocean. Here, we explore the potential consequences of these physiological responses to environmental conditions and outline future areas of investigation to better constrain the impacts of IPL remodeling on marine biogeochemistry.

4.3.1 Relative adaptability of Phytoplankton Classes to environmental change

IPL classes are either partitioned to the thylakoid membranes (i.e., MG, DG, SQ, and PG) in variable distributions to maintain photosynthetic function, or to extraplastidic membranes under growth/reproductive phases (BL, PC, PE, and PG). As environmental conditions change, the ability of phytoplankton to adjust these IPL compositions may be an important driver controlling the phytoplankton community structure. Within a few days of the ODZ water addition, the mesocosms became oligotrophic, which coincided with a major shift in phytoplankton groups, from predominantly diatoms, Chlorophyceae, and Cryptophyceae to a dominance of ‘survivalist’ mixotrophic dinoflagellates. An obvious advantage of mixotrophic

dinoflagellates to the largely N-limited conditions of this experiment is their ability to extract N from the DON pool, and a lower dependence on light compared to the other previously mentioned phytoplankton groups.

To contextualize the impacts of IPL remodeling on the phytoplanktonic community, we performed an additional random forest analysis to identify the most important IPLs (both individual moieties and polar head group totals) in the prediction of individual phytoplankton classes. We applied this analysis to identify what IPL remodeling processes may be more readily available to each major phytoplankton class in this experiment (Fig. 8). While all the phytoplankton classes are likely to produce each IPL class to varying extents, the differences in their relative distributions may provide insight into differences in phytoplankton adaptability under changing environmental conditions. Dinoflagellates are heavily associated with high abundances of the glycolipids SQ and DG, with far less apparent dependence on phospholipids and MGs (Fig. 8). The dominance of dinoflagellate biomass is attributed to the severe scarcity of inorganic nitrogen and relatively low light intensity due to self-shading effects (Bach et al., 2020). Dinoflagellates may also take advantage of the recycling of N-bearing PCs and BLs to alleviate nitrogen limitation or to produce energy storing TAGs. The high proportion of DGs and SQs and recycling of MGs may also prove advantageous under high pH/low pCO₂ conditions in surface waters, alleviating pH stressors, or the energy investment in active HCO₃⁻ transport for photosynthesis. These IPL remodeling strategies may have played a role in the dominance of dinoflagellates under the post-upwelling experimental conditions.

In the early phases of the experiment and when inorganic N is readily available, diatoms (and to a lesser extent Cryptophyceae and Chlorophyceae) dominate the water column biomass. Each of these phytoplankton classes (as well as *Synechococcus*) indicates a greater dependence of the N-bearing PCs and/or BLs, potentially signaling a reduced capacity to scavenge these extrachloroplastic IPLs within the cell for N, or as substrates for TAG production. Notably, these phytoplankton consistently make up a greater proportion of the total chl-a pool in subsurface waters, suggesting an ability to accommodate the lower O₂ concentrations and pH in exchange for greater N availability (see summary in Fig. 8). Some of this adaptability may be related to the recycling of MGs for additional acyl units under hypoxia (most likely amongst Chlorophyceae and *Synechococcus*), and/or the generally higher relative production of structural membrane lipids under low light growth conditions (see impact of light discussion above).

In the Eastern Tropical South Pacific (ETSP), upwelling events exhibit a distinctive trend of increasing frequency and intensity, setting it apart from other eastern boundary systems (Abrahams et al., 2021; Oyarzún and Brierley, 2019). This phenomenon brings about important changes in environmental conditions, including a decrease in sea surface temperature (~0.37°C, average from 1982 to 2019, Abrahams et al., 2021) alongside oxygen reduction and reduced pH in the coastal zones of the region (Pitcher et al., 2021). Simultaneously, there is an observed boost in primary productivity (Gutierrez et al., 2011; Tretkoff, 2011). The impact of strong upwelling, followed by rapid thermal stratification when upwelling subsides, could signify abrupt changes that affect the community structure of phytoplankton (Gutierrez et al., 2011). Such environmental changes favor

species better adapted to stress conditions, such as mixotrophic dinoflagellates and silicoflagellates, at the expense of diatom populations (Hallegraeff, 2010), which could translate to a decrease in highly nutritious saturated fatty acids for higher trophic levels (Hausse et al., 2012). Furthermore, a potential reduction in diatom biomass and diversity could have consequences on the ocean biological carbon pump (Tréguer et al., 2018).

Under the ongoing and projected scenarios of climate change, our mesocosm experiment demonstrates how shifts in the phytoplankton community translate into changes in the lipid composition of their cell membranes. This adds weight to the school of thought that lipid remodeling amongst the phytoplankton community could have repercussions at higher trophic levels, as recently discussed by Holm et al. (2022), and on ocean biogeochemistry as we discuss below.

4.3.2 Cycling of Carbon, Nitrogen, and Sulfur

Combined, neutral (TAGs) and polar lipids (IPLs) make up on the order of 17.6% to 34.7% of the cell by dry weight under N-replete and N-deprived conditions, respectively (Morales et al., 2021), with other environmental stressors able to compound the accumulation of neutral lipids (Zienkiewicz et al., 2016). This represents a considerable portion of algal cellular carbon, and total water column carbon stocks that supply the transfer of fatty acids to higher trophic levels (Twining et al., 2020). Trophic transfer of fatty acids to grazing zooplankton may be affected by the relative contribution of polar and neutral lipids, as they can have significantly different turnover rates (Burian et al., 2020). This suggests that the relative concentrations of TAGs/IPLs in phytoplanktonic communities may impact the rates of fatty acid remineralization or uptake into higher trophic levels. Additionally, while our analysis did not focus on individual fatty acid structures, other experiments investigating the effects of increasing pCO₂ and nitrogen deficiency have demonstrated a reduction in the relative production of essential fatty acids resulting in negative reproductive effects on primary consumers (Meyers et al., 2019). In this context, our IPL results reveal modifications (among polar head groups, carbon numbers, and unsaturations) in response to divergent environmental conditions. However, further investigation of the relative concentrations and turnover rates between TAGs and IPLs, in addition to the production of essential fatty acids, are critical next steps in assessing the impacts of IPL remodeling on trophic transfer processes.

While IPLs are not a major source of cellular N compared to proteins, the highly labile nature of these molecules after cell death may be a potential source of rapidly recycled N via bacterial hydrolysis. On average, N-bearing IPLs such as PCs or BLs, consist of ~2% N by molecular weight (estimated based on a typical 700g/mol betaine lipid). Based on their abundance, they could contribute as much as 10.7 nmol/L of inorganic N in the highly N depleted surface waters of this experiment. Inorganic N species are exceedingly low, and often below the limit of detection (< ~0.1 μmol/L, see methods for details), particularly in the surface samples for most of the experiment after the OMZ water addition. Thus, it is possible that the rapid degradation and recycling of N from IPLs could prove to be an additional supply of N under extremely limiting conditions.

858 Given the diel variability in TAG production via eukaryotic phytoplankton and the potential impacts on energy and carbon
859 fluxes (Becker et al., 2018), further analysis is needed to determine at what temporal scale IPLs may be recycled for TAG
860 synthesis. While Becker et al. (2018) did not find evidence of diacylglycerol transferase (DGAT) activity related to diel cycles,
861 additional investigations of other transferases or lipases relevant to the degradation of IPLs (e.g., PDAT or PGD) are critical.

862
863 IPLs may also be an important intermediate in the surface ocean sulfur cycle. Sulfur metabolites produced by phytoplankton
864 (Durham et al., 2019) play central roles in microbial food webs by functioning in metabolism, contributing to membrane
865 structure, supporting osmotic and redox balance, and acting as allelochemicals and signaling molecules (Moran and Durham,
866 2019). SQs are highly abundant IPLs, particularly in surface waters, and can constitute up to 60 nmol S/L, potentially
867 contributing to a significant proportion of the dissolved organic S in this region (ranging from ~100-225 nmol/L; Lennartz et
868 al., 2019). The ability to catabolize SQs is widespread in heterotrophic bacteria and provides a highly ubiquitous source of
869 sulfur (and carbon) substrates in the synthesis of other essential sulfur metabolites (Speciale et al., 2016). As such, SQs are
870 considered major contributors to the global biosulfur cycle (Goddard-Borger and Williams, 2017) and their considerable
871 production in phytoplankton membranes may be augmented by changes in temperature, light availability (i.e., shading effects),
872 and pH (see discussion above).

873
874 Lipids are also sources of dissolved organic carbon (DOC) following hydrolysis by bacterial extracellular enzymes (Myklestad,
875 2000). Lipids have been observed as the most highly aged component of DOC and are considerably longer lived than lipids
876 found in suspended POC (Loh et al., 2004). However, much of the DOC pool remains uncharacterized (Nebbioso and Piccolo,
877 2013; Hansell and Carlson, 2014) and little is known regarding the specific processes forming refractory DOC fractions
878 (Hansell, 2013). Yet, preliminary estimates of the total production of relatively refractory DOC (>100-year lifetime) suggest
879 potential significance to global biogeochemistry (Legendre et al., 2015). Furthermore, it is yet unclear as to how DOC cycling
880 may be affected by changing ocean conditions (Wagner et al., 2020). Understanding the degradation pathways involved with
881 abundant lipid pools and how they may contribute to the DOC pool in the surface ocean is of crucial importance to constraining
882 the fluxes of refractory organic carbon pools.

883

884 **5 Conclusions**

885 We investigated the potential for phytoplankton to evoke lipid remodeling in response to multiple environmental stressors
886 during a two-month mesocosm experiment in the Eastern Tropical South Pacific Ocean off the coast of Peru. Our study is
887 motivated by the need of understanding how the expected changes in oceanographic conditions due to climate forcing in this
888 area will impact the phytoplanktonic community driving primary productivity in the surface ocean, the chemistry of their cell
889 membranes, and thus the chemical composition of particulate organic matter. A key aspect of our efforts was to statistically

separate the relative roles of phytoplankton community composition and lipid remodeling in response to multiple environmental stressors on the composition of the environmental algal lipidome over time. By using a combination of multiple linear regressions, classification and regression trees, and random forest analyses, we report evidence of multiple environmental variables potentially imparting controls on IPL head groups, many of which are known to have specific functions in the chloroplast. The main takeaways of our study include:

- IPLs are effective predictors of changes in the prevailing sources of phytoplankton biomass throughout the experiment.
- The proportion of glycolipids (MG and SQ) increases at higher temperatures, possibly to maintain thermal stability of the photosynthetic apparatus in response to thermal stress.
- The relationship between light levels and the distribution of MGs and PGs possibly invokes photoprotective and antioxidant mechanisms provided by MGs, as well as the role of PGs in electron transport processes in photosystems I and II.
- Differences in the abundance of glycolipids between surface and subsurface waters appear to covary with oxygen availability. We expect this difference to be driven by the degradation of MGs and DGs under anaerobiosis or oxygen stress.
- The reduction in N-bearing IPLs (i.e. BLs and PE) and the higher abundance of non-N-bearing IPLs (i.e., PG, SQ and MG) under more N-limited conditions suggests a possible IPL substitution mechanism to compensate for N limitation.
- Whereas our study does not include the analysis of TAGs, our results are consistent with broader work suggesting that IPLs may be used as a substrate for the generation of acyl chains in TAG production in response to environmental stressors such as N limitation, variable pH or inorganic carbon availability, hypoxia, and varying light levels. However, such mechanism in this mesocosm experiment remains to be tested in future work.
- The cellular adaptations listed above likely contribute to the observed shifts in cellular content from structural or extrachloroplastic membrane lipids (i.e., PCs, PEs, BLs, and certain PGs) under high growth conditions, to thylakoid/plastid membrane lipids (i.e., MGs, DGs, SQs, and certain PGs) while exposed to environmental stressors.

We hypothesize that these remodeling processes may involve major shifts in the elemental stoichiometry of the cell, and thus alter the fluxes of C, N, and S to higher trophic levels, signaling potential impacts on the broader cycling of these biorelevant elements under different scenarios of future environmental change. Therefore, we suggest that future studies addressing the biogeochemical consequences of climate change in the Eastern Tropical South Pacific Ocean must include the role of lipid remodeling in phytoplankton.

Data availability

The environmental study data are available at: <https://doi.org/10.1594/PANGAEA.923395> (Bach et al., 2020). Biomarkers metadata that generates and supports the findings of this study and R code are available in: <https://github.com/Guachan/IPLs-KOSMOS/tree/0.1.0> (<https://doi.org/10.5281/zenodo.10408453>, Cantarero, 2023).

Author contribution

JS and SC designed the study. JS funded the study. JS and CV funded sample collection during the mesocosm experiment. UL and LTB designed, funded, organized, and carried out the mesocosm experiment. PA, JS, LTB, and UR carried out sample collection. SIC led laboratory and analytical work with the assistance of ND and under the supervision of JS. SEI carried out data analysis. SIC, JET, BCS, and BR performed statistical analyzes. SIC wrote the manuscript with major editorial contributions from EF, HA, and JS and with comments from all co-authors.

Competing interests

The authors declare that they have no conflict of interest. At least one of the (co-)authors is a member of the editorial board of Biogeosciences.

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1420 **Tables**

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1422 **Table 1: Concentration of inorganic nutrients (μmol/L), nutrient stoichiometry, and pH_T of the two batches of ODZ**
1423 **water collected and added to the mesocosm experiment.**

	Si(OH) ₄	PO ₄ ³⁻	NO ₂ ⁻	NH ₄ ⁺	NO ₃ ⁻	N:P:Si	pH _T	Depth (m)
Station 1	17.4	2.6	0	0.3	0	0.1:1.0:6.7	7.47	30
Station 3	19.6	2.5	2.9	0.3	1.1	1.7:1.0:7.8	7.49	70

1426 **Table 2: IPL classes included in this study and their respective acronyms.**

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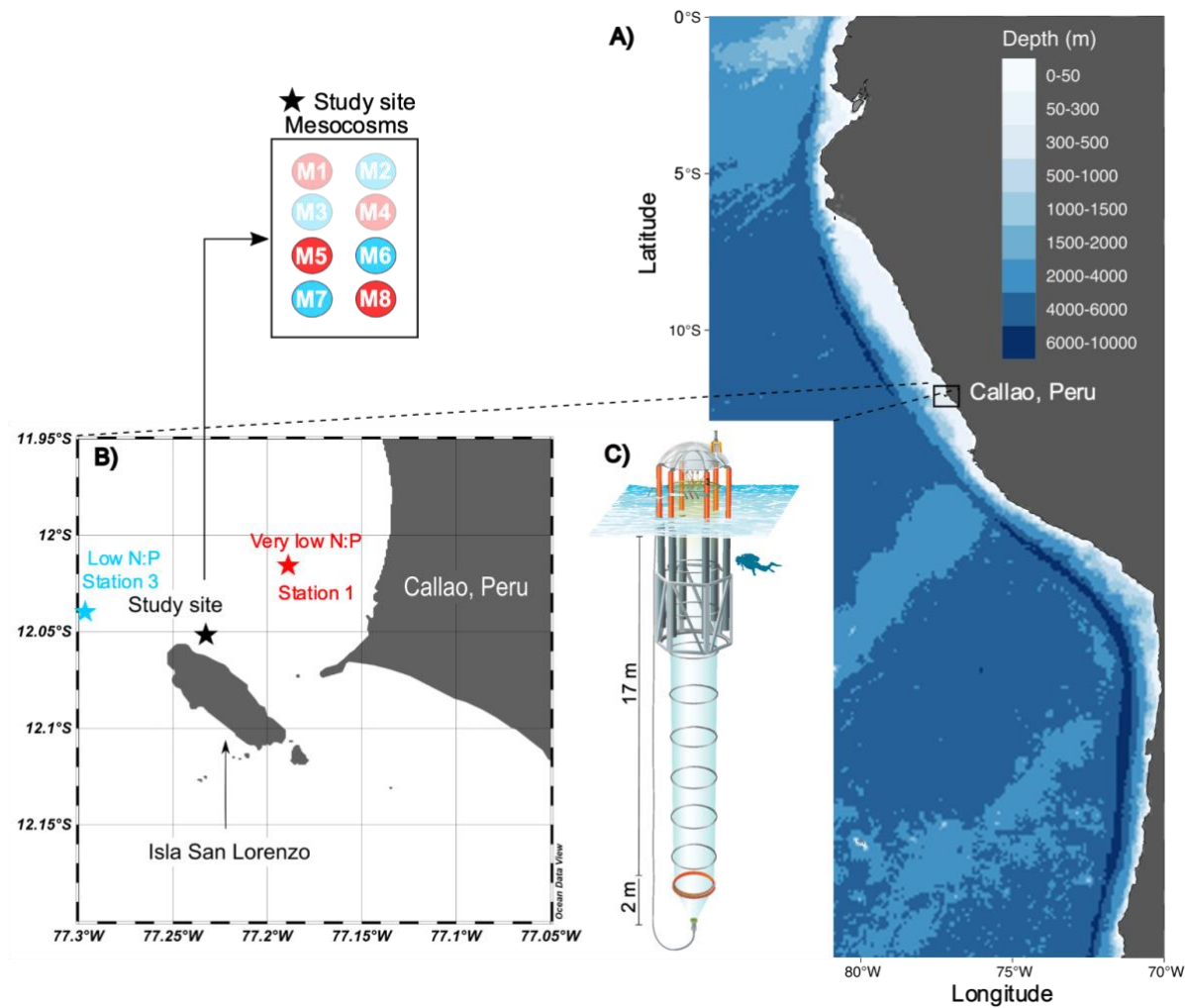
	IPL Headgroups	Acronym	IPL Classes
	Sulfoquinovosyl	SQ	Glycolipid
	Monogalactosyldiacylglycerol	MG	Glycolipid
	Digalactosyldiacylglycerol	DG	Glycolipid
	Phosphatidylglycerol	PG	Phospholipid
	Phosphatidylethanolamine	PE	Phospholipid
	Phosphatidylcholine	PC	Phospholipid
	Diacylglyceryl trimethylhomoserine	DGTS	Betaine Lipid
	Diacylglyceryl hydroxymethyl-trimethyl-beta-alanine	DGTA	Betaine Lipid
1428	Diacylglyceryl carboxyhydroxymethylcholine	DGCC	Betaine Lipid

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1431 **Figures**

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1435 **Figure 1: The KOSMOS study site. (A) Overview map indicating the location of the study region in La Punta, Callao, Perú. (B)**
1436 **Detailed map of the study site with the mesocosm arrangement (M1-8). Red (very low N:P) and blue (low N:P) colors indicate the**
1437 **different stations for ODZ water collection (1 and 3; star symbols) and replicates of the two different water treatments (numbers in**
1438 **circles in insert). (C) Diagram of a mesocosm unit with underwater bag dimensions. Modified figure from Chen et al. (2022).**

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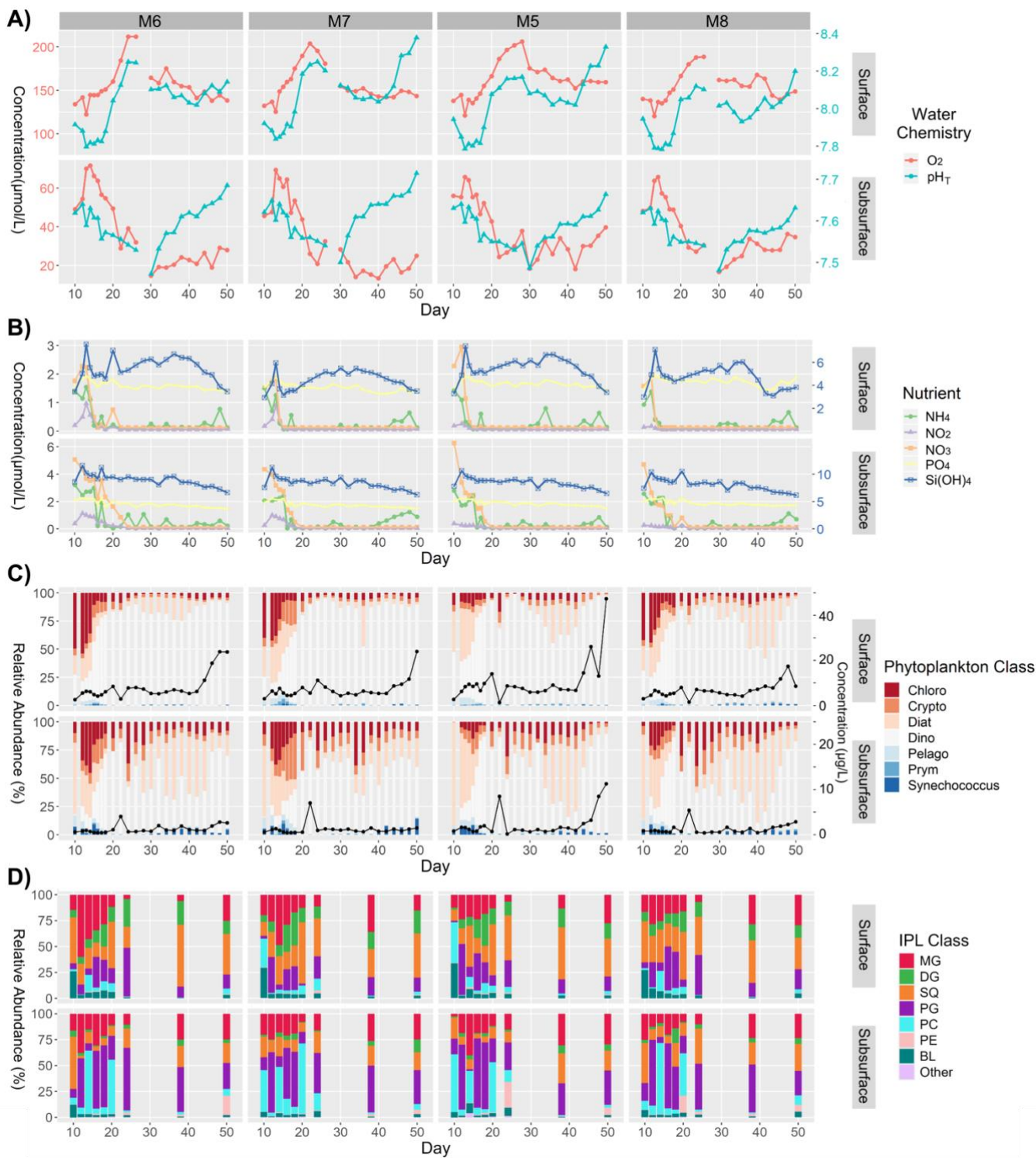
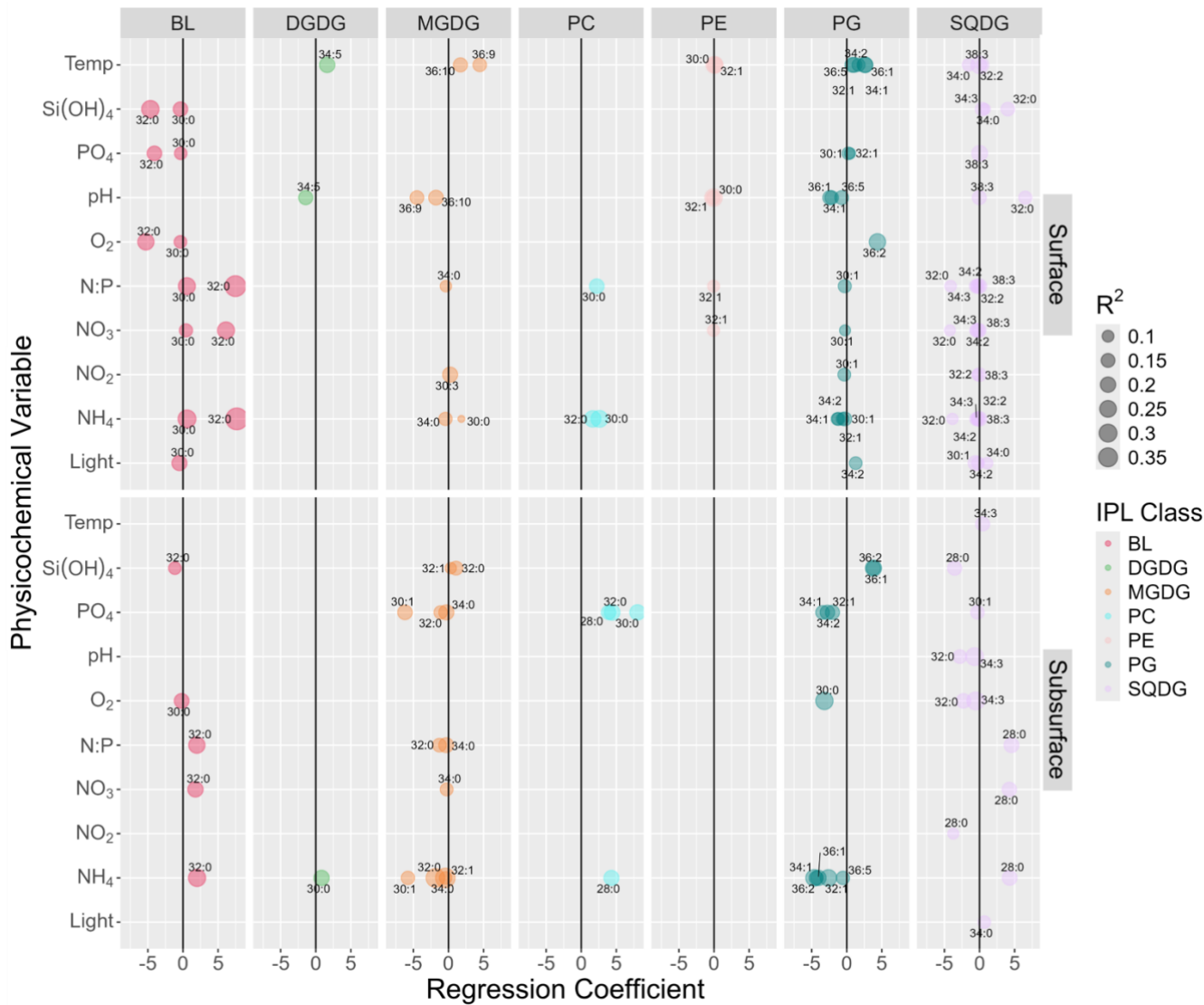


Figure 2: Summary of major physicochemical, biological, and lipidomic measurements in surface and subsurface waters of mesocosms analyzed in this study from the time of deep-water injection to the end of the experiment. A) Concentration of O₂ (μmol/L; left Y axis), and pH_T (total scale; right Y axis). B) Concentration of inorganic N (left Y axis), P (left Y axis), and Si(OH)₄ (right Y axis) expressed in μmol/L. C) Relative abundance (%) of phytoplankton classes (left Y axis) as well as total chl-a concentrations expressed in μg/L (black line; right Y axis). D) Relative abundance (%) of major IPL classes based on headgroup contributions to the total IPL pool.

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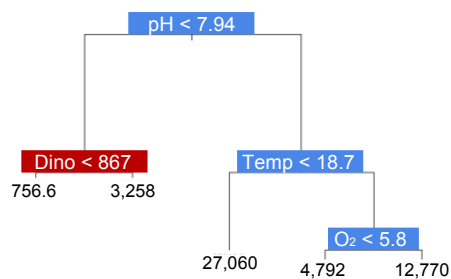


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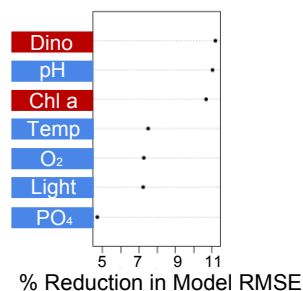
Figure 3: Summary of multiple linear regression analysis between the abundances of major IPL molecules (>2.5% of total IPL pool) and physicochemical parameters showing only statistically significant ($p < 0.05$) linear responses after controlling for false discovery rate using a 0.1 alpha cutoff on adjusted p-values. The size of circles indicates the magnitude of the linear regression adjusted R^2 . The upper and lower panels represent surface and subsurface waters, respectively. Numbers next to circles indicate the total number of carbon atoms and double bonds in core fatty acid chains.

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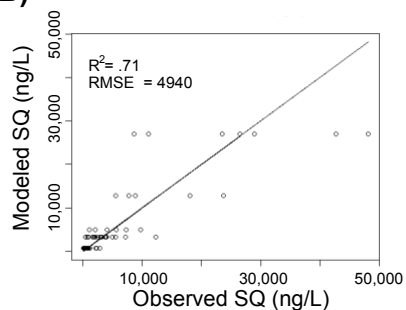
A) Pruned Tree SQ



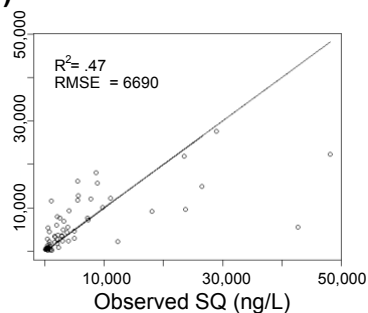
C) Random Forest SQ



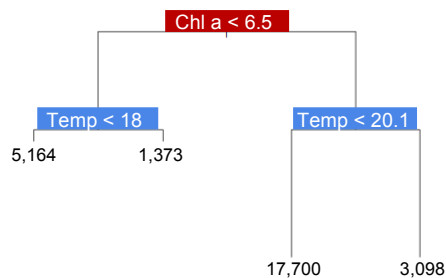
B)



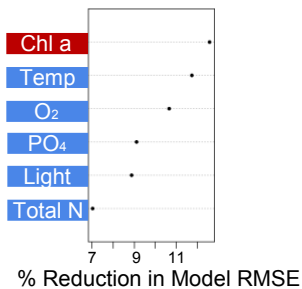
D)



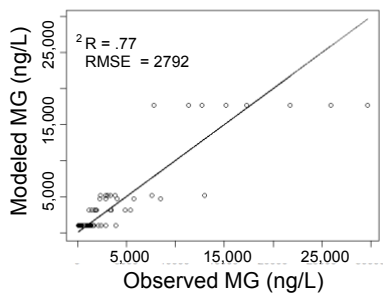
E) Pruned Tree MG



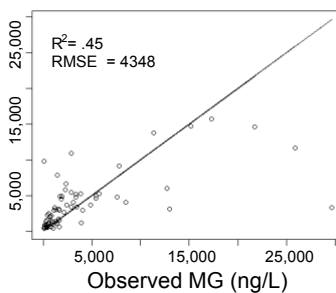
G) Random Forest MG



F)

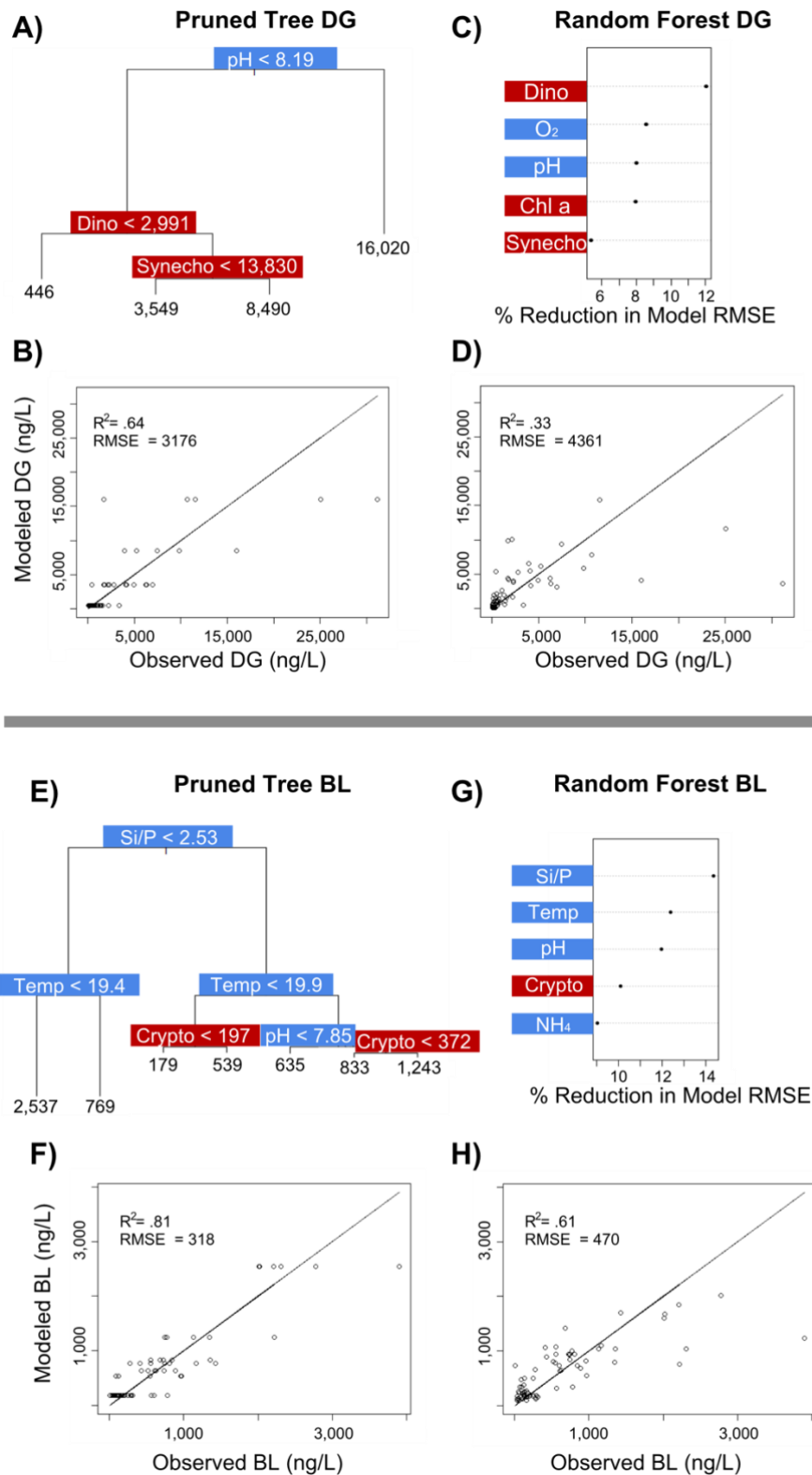


H)



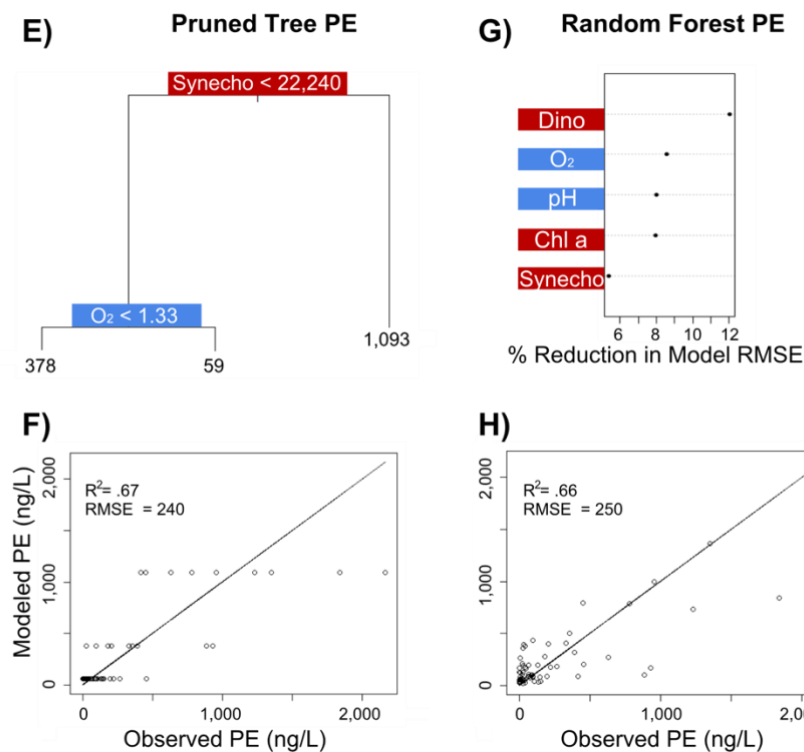
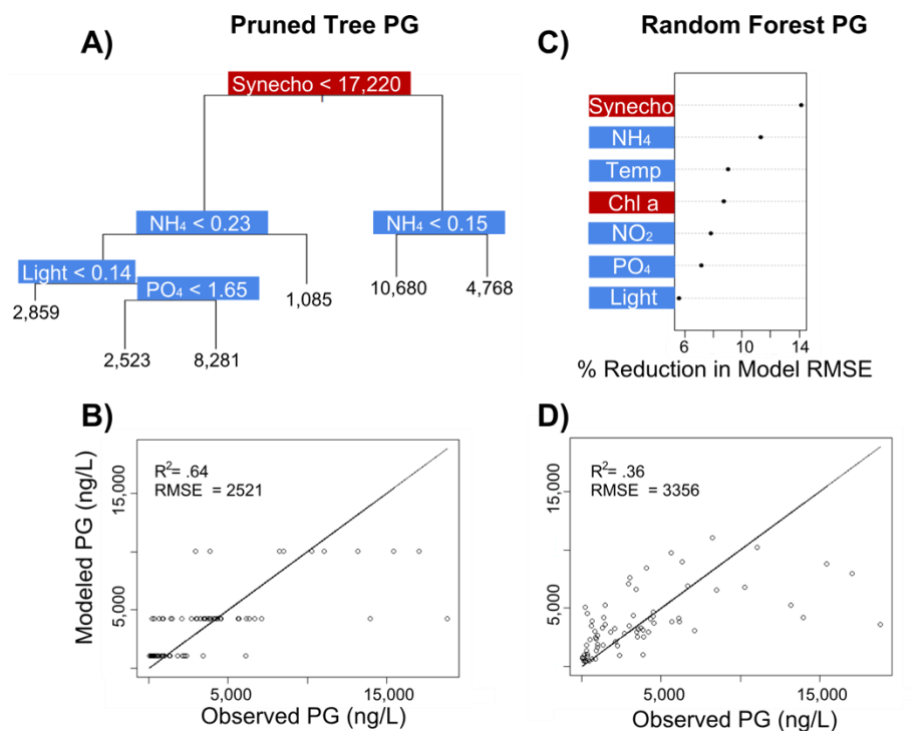
1455 **Figure 4: Classification and Regression Tree (CART) and Random Forest analyses of selected IPL classes (Top: SQ; Bottom: MG).**
1456 **A and E indicate a summary of primary predictors in the best fit CART (i.e. Pruned Tree), whereas B and F show model**
1457 **performance via adjusted R² and RMSE. C and G display Random Forest variable importance in the prediction of IPL classes as**
1458 **defined by % reduction in RMSE, whereas D and H show model performance via adjusted R² and RMSE. Environmental variables**
1459 **are depicted in blue, whereas biological variables are shown in red for both analyses.**

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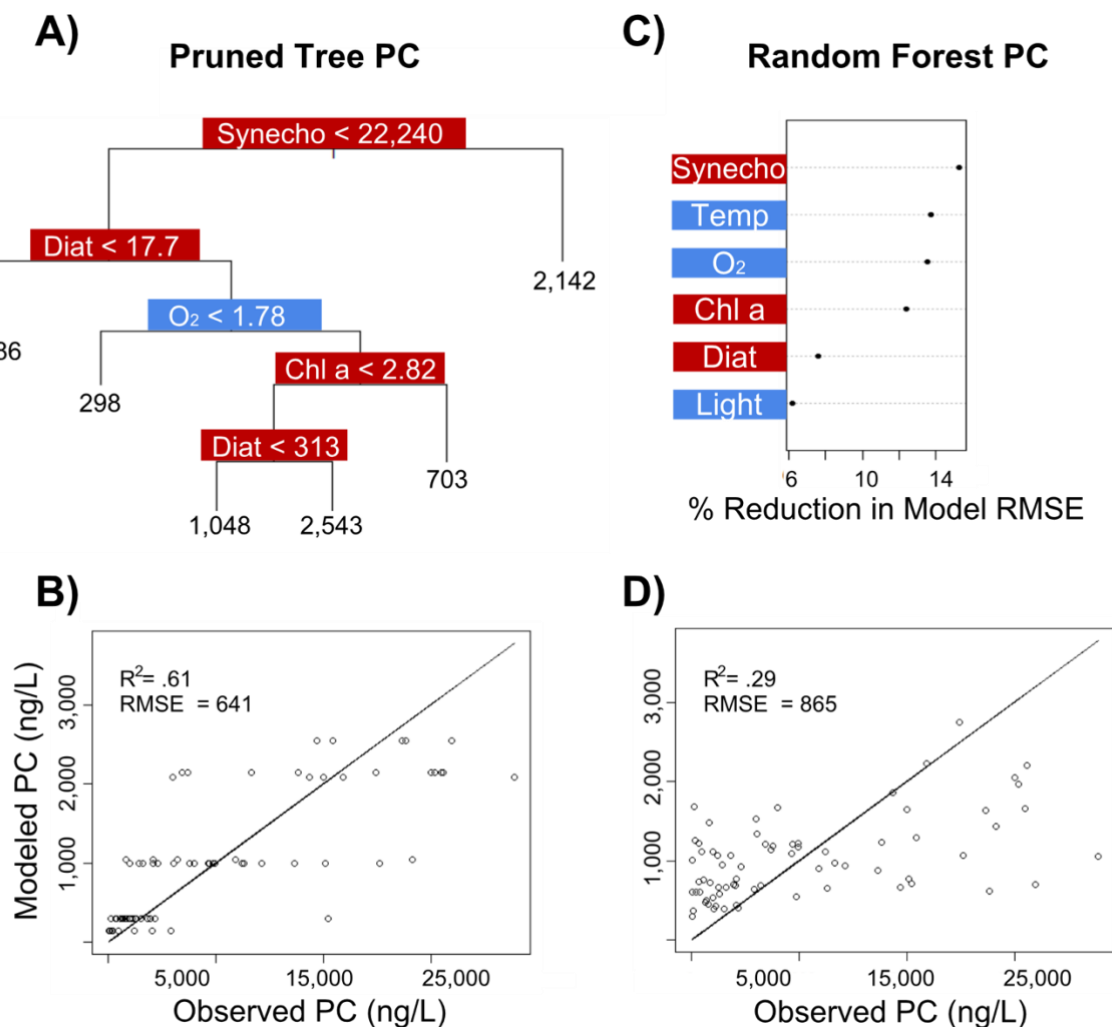
1462 **Figure 5: Classification and Regression Tree (CART) and Random Forest analyses of selected IPL classes (Top: DG; Bottom: BL).**
1463 **A and E indicate a summary of primary predictors in the best fit CART (i.e. Pruned Tree), whereas B and F show model**
1464 **performance via adjusted R² and RMSE. C and G display Random Forest variable importance in the prediction of IPL classes as**
1465 **defined by % reduction in RMSE, whereas D and H show model performance via adjusted R² and RMSE. Environmental variables**
1466 **are depicted in blue, whereas biological variables are shown in red for both analyses.**

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1469 **Figure 6: Classification and Regression Tree (CART) and Random Forest analyses of selected IPL classes (Top: PG; Bottom: PE).**
 1470 **A and E indicate a summary of primary predictors in the best fit CART (i.e. Pruned Tree), whereas B and F show model**
 1471 **performance via adjusted R^2 and RMSE. C and G display Random Forest variable importance in the prediction of IPL classes as**
 1472 **defined by % reduction in RMSE, whereas D and H show model performance via adjusted R^2 and RMSE. Environmental variables**
 1473 **are depicted in blue, whereas biological variables are shown in red for both analyses.**

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1478 **Figure 7: Classification and Regression Tree (CART) and Random Forest analyses of selected IPL classes (PC). A indicate a**
 1479 **summary of primary predictors in the best fit CART (i.e. Pruned Tree), whereas B shows model performance via adjusted R^2 and**
 1480 **RMSE. C and G display Random Forest variable importance in the prediction of IPL classes as defined by % reduction in RMSE,**
 1481 **whereas D show model performance via adjusted R^2 and RMSE. Environmental variables are depicted in blue, whereas biological**
 1482 **variables are shown in red for both analyses.**

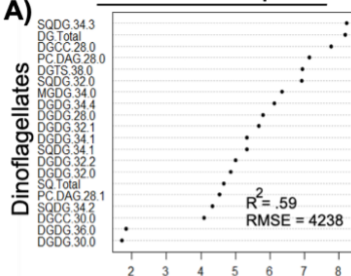
1483

Predictive Lipids

Dominant IPL Classes

Potential IPL Remodeling

A)



SQ, DG

Recycling of PC and BL under N-limitation

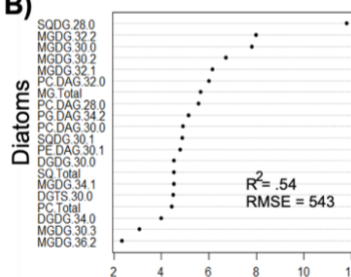
Recycling of MG under high pH/low pCO₂, or anaerobiosis

Increased proportion of DG and SQ under thermal stress or high pH/low pCO₂

Increased proportion of SQ under shaded conditions

Recycling of MG and PG under high growth conditions

B)



MG, SQ, PC

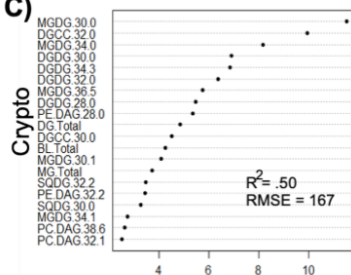
Recycling of BL under N-limitation

Increased proportion of SQ under thermal stress, shaded conditions, and high pH/low pCO₂

Increased proportion of MG for photoprotective and antioxidant mechanisms

Recycling of PG under high growth conditions

C)



BL, DG, MG

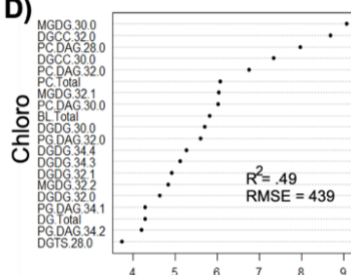
Recycling of PC under N-limitation

Enhanced DG under thermal stress

Enhanced MG for photoprotective and antioxidant mechanisms

Recycling of SQ and PG under high growth conditions

D)



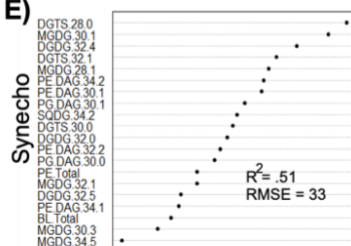
PC, BL, DG

Recycling of MG under high pH/low pCO₂ or anaerobiosis

Enhanced DG under thermal stress

Recycling of SQ, MG, and PG under high growth conditions

E)



PE, BL

Recycling of PC under N-limitation

Recycling of MG under high pH/low pCO₂ or anaerobiosis

Increased proportion of SQ under shaded conditions

Recycling of SQ, MG, and PG under high growth conditions

% Reduction in Model RMSE

1485

1486 **Figure 8: Random Forest-based identification of the 15 most important individual IPLs in the prediction of phytoplankton groups.**
1487 **Dominant IPL classes and the potential remodeling advantages for each group are summarized. The nomenclature of IPL molecules**
1488 **follows that of the main text, starting with the type of headgroup, followed by the number of carbon atoms in fatty acid chains, and**
1489 **the total number of double bonds in the fatty acid chains. The root mean square error (RMSE) of random forest models is defined**
1490 **as ng/L of chl-a, and coefficient of determination (R^2) provided for each model fit.**