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Gradients in intact polar diacylglycerolipids across the Mediterranean Sea are related to phosphate availability

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Intact polar membrane lipids compose a significant fraction of cellular material in plankton and their synthesis imposes a substantial constraint on planktonic nutrient requirements. As a part of the Biogeochemistry from the Oligotrophic to the Ultraoligtrophic Mediterranean (BOUM) cruise we examined the distribution of several classes of intact polar diacylglycerolipids (IP-DAGs) across the Mediterranean, and found that phospholipid concentration as a percent of total lipids correlated with phosphate concentration. In addition, the ratios of non-phosphorus lipids to phospholipids – sulfoquinovosyldiacylglycerol (SQDG) to phosphatidylglycerol (PG), and betaine lipids to phosphatidylcholine (PC) – were also found to increase from west to east across the Mediterranean. Additionally, microcosm incubations from across the Mediterranean were amended with phosphate and ammonium, and in the course of several days nutrient amendments elicited a shift in the ratios of IP-DAGs. These experiments were used to assess the relative contribution of community shifts and physiological response to the observed change in IP-DAGs across the Mediterranean. The ratio of SQDG to chlorophyll a was also explored as an indicator of phytoplankton response to nitrogen availability. This study is the first to demonstrate the dynamic response of membrane lipid composition to changes in nutrients in a natural, mixed planktonic community.

1 Introduction

Lipids form the membranes of cells and comprise a substantial fraction of planktonic carbon and phosphorus (Wakeham et al., 1997; Van Mooy and Fredricks, 2010). As such, lipids play key roles in the biogeochemical cycles of carbon and phosphorus in the ocean (Suzumura and Ingall 2001), and their synthesis poses a significant cellular nutrient requirement (Van Mooy et al., 2008, 2009). Bacterial and eukaryotic marine membranes are composed mainly of intact polar diacylglycerolipids (IP-DAGs) and these compounds are valuable tools for the study of microbes in the environment

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(Rütters et al., 2002; Sturt et al., 2004). These IP-DAGs are classified by the structure of their headgroups, which are differentiated by containing phosphorus, nitrogen, sulfur, sugars, or some combination of these (Guschina and Harwood, 2006; Kato et al., 1996; Oliver and Colwell, 1973; Sato, 1992; Van Mooy and Fredricks, 2010). The relative abundance of different headgroups within a membrane contributes to the cellular demand for limiting nutrients. Because phosphorus is in low supply in much of the ocean and is in high cellular demand, the constraint imposed by production of phospholipids is nontrivial for marine microbes (Van Mooy et al., 2006, 2008, 2009).

Different groups of plankton contain different arrays of IP-DAG headgroups (Popendorf et al., 2011; Van Mooy et al., 2006), such that a community dominated by phytoplankton could be expected to have a different total IP-DAG composition than a community dominated by heterotrophic bacteria. Some types of plankton have been shown to be capable of substituting non-phosphorus lipids for phospholipids in conditions of phosphorus stress (Benning et al., 1993, 1995; Martin et al., 2010; Minnikin et al., 1974; Van Mooy et al., 2009), adjusting the composition of their membrane as a physiological response to nutrient conditions. This physiological response has been demonstrated in culture, but has not previously been observed in a natural, mixed planktonic community. It has been observed that the ratios of non-phosphorus lipids to phospholipids in the water column were lower in the phosphorus-replete South Pacific and higher in the phosphorus-limited Sargasso Sea (Van Mooy et al., 2009), but we posit that this difference in IP-DAG ratio could be attributed to either differences in planktonic community structure or differences in the physiological adaptation of plankton to low nutrients. Deconvolving the contribution of these two mechanisms would provide deeper insights on the ecological and oceanographic meaning of IP-DAG ratios in the environment.

The Mediterranean Sea is well-suited for a study of the influence of phosphate limitation on IP-DAG distribution, as it is characterized by exceptionally low concentrations of phosphate and high ratios of dissolved N:P (Krom et al., 1991; Moutin and Raimbault, 2002; Pujo-Pay et al., 2011). In the Biogeochemistry from the Oligotrophic to the Ultra-

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oligotrophic Mediterranean (BOUM) cruise the ratio of nitrate plus nitrite to phosphate ranged from 24.5 at the surface in the west to 32.4 in the east (Pujo-Pay et al., 2011), far exceeding the canonical Redfield ratio of 16 and distinguishing the Mediterranean Sea as one of the most phosphorus-depleted bodies of water in the world. The major input of water to the Mediterranean is over the shallow sill of the Strait of Gibraltar, bringing in nutrient-depleted surface water from the Atlantic and establishing the oligotrophic nature of the Sea (Krom et al., 1991). The bathymetry and circulation lead to distinctly different water masses in the eastern and western basins with a gradient of nutrients, primary production, and planktonic community structure from west to east (Moutin and Raimbault, 2002; Moutin et al., 2002). During the summer months intense stratification develops in the Mediterranean, leading to a shallow mixed layer that is downwelled in some areas by anticyclonic eddies (Siokou-Fragou et al., 2010; Moutin et al., 2011). Thus the Mediterranean Sea, as a contained, highly oligotrophic water mass with a longitudinal gradient of phosphate, provides an excellent study site for the response of IP-DAGs to nutrient limitation.

In this study we set out to see if the phosphorus gradient across the Mediterranean Sea (Krom et al., 1991, Pujo-Pay et al., 2011) leads to a shift in the membrane lipids of marine microbes, where we would expect the oligotrophic conditions in the west to correspond with proportionally more abundant phospholipids than the ultraoligotrophic conditions in the east. In addition, we performed microcosm incubations with nutrient amendments to test whether changes in the available nutrients would elicit a measurable change in IP-DAG composition in a natural, mixed planktonic community on short timescales. These controlled incubations allowed us to assess the relative contribution of physiological response and changes in community composition to the observed changes in IP-DAGs.

2.1 Sampling sites

This study was conducted as a part of the Biogeochemistry from the Oligotrophic to the Ultraoligotrophic Mediterranean (BOUM) cruise in June-July of 2008 aboard the R/V *L'Atalante*. Samples were collected for IP-DAG quantification along a transect from the Algero Provencal basin (39.1° N, 5.2° E) to the eastern Levantine basin near Cyprus (33.7° N, 32.7° W; Fig. 1). At 11 stations samples for IP-DAG depth profiles were collected (numbered stations, 1–21). Each depth profile was composed of nine or ten samples taken between 5 m and 250 m. At three additional stations (lettered stations, station A in the Algero Provencal basin, station B in the Ionian basin, and station C in the Levantine basin), located near the center of anticyclonic eddies, multiple depth profiles were collected in close proximity (10 profiles in total). At each of these stations single profiles were taken to greater depths: 2700 m, 2500 m, and 900 m respectively at stations A, B and C. Microcosm incubations with nutrient amendments were also conducted at these stations.

2.2 Microcosm incubation set up and sampling

Water for the microcosm incubation was collected with Niskin bottles at 8 m depth. Each microcosm incubation consisted of triplicate 20 L carboys for each of four conditions: control (C) no nutrient addition; nitrogen addition (N) with 1600 nM NH $_4$ Cl at stations A and B, 3200 nM at station C; phosphorus addition (P) with 100 nM KH $_2$ PO $_4$ at all stations; nitrogen and phosphorus addition (NP) with 1600 nM NH $_4$ Cl, 100 nM KH $_2$ PO $_4$ at stations A and B, 3200 nM NH $_4$ Cl, 100 nM KH $_2$ PO $_4$ at station C. Carboys were incubated in on-deck incubators flushed with surface seawater to maintain ambient temperature, and outfitted with screens to reduce incident light by ca. 50 %. Samples were taken daily for a range of biogeochemical parameters (see Tanaka et al., 2011 for full results). Samples were taken for IP-DAGs at the initiation of the experi-

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ments and termination, at station A at the end of three days, at stations B and C at the end of four days.

2.3 IP-DAG sample collection and extraction

Samples for depth profiles were collected using Niskin bottles mounted on a rosette equipped with sensors for conductivity, temperature, pressure, oxygen, photosynthetically available radiation (PAR), and chlorophyll fluorescence. All samples for IP-DAGs were 1 L, transferred from CTD casts (depth profiles) or incubation carboys (microcosm) to filtration using polycarbonate bottles. Seawater was filtered onto 47 mm 0.2 µm poresize hydrophilic Durapore filters (Millipore) using gentle vacuum filtration (<200 mm Hg). Filters were folded in half, cell side in, wrapped in combusted aluminum foil and stored in liquid nitrogen until extraction. Extraction was conducted using a modified Bligh and Dyer solvent extraction method with phosphatidylethanolamine-N-(2,4-dinitrophenyl) (DNP-PE; Avanti Polar Lipids) as an internal recovery standard (Bligh and Dyer, 1959; Popendorf et al., 2011).

2.4 IP-DAG quantification

Total lipid extracts were analyzed using high performance liquid chromatography/mass spectrometry (HPLC-MS) with an Agilent 1200 HPLC and Thermo Scientific TSQ Vantage triple quadrupole MS with a heated electrospray ionization (ESI) interface following Popendorf et al. (2011). HPLC methods are given in Sturt et al. (2004). Nine IP-DAGs were quantified: three phospholipids, phosphatidylglycerol (PG), phosphatidylethanolamine (PE), phosphatidylcholine (PC); three glycolipids, sulfoquinovosyldiacylglycerol (SQDG), monoglycosyldiacylglycerol (MGDG), diglycosyldiacylglycerol (DGDG); and three betaine lipids, diacylglyceryl hydroxymethyl-trimethyl- β -alanine (DGTA), diacylglyceryl trimethylhomoserine (DGTS), and diacylglyceryl carboxyhydroxymethylcholine (DGCC). MS detection and quantification of phospholipids and glycolipids followed the methods given in Popendorf et al. (2011). Additionally, be-

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taine lipids were analyzed. Detection of the betaine lipids was by positive ion mode product ion scans, for DGTS and DGTA product ion m/z 236; for DGCC product ion m/z 104. Quantification of DGCC used an external standard curve prepared with natural DGCC extracted from Thalassiosira pseudonana grown in the Van Mooy lab, purified by preparative HPLC and then quantified by gas chromatography flame-ionization detector (GC-FID) analysis of transesterified fatty acid methyl esters. The response factor of DGTA on the TSQ MS was established previously using standard curves of DGCC and DGTA prepared in a similar fashion. MS response of DGTS and DGTA were compared on a Thermo-Finnigan LCQ Deca XP ion trap MS and the TSQ MS to calculate a response factor for DGTS.

Nutrients 2.5

Dissolved nitrate (NO_3^-) , nitrite (NO_2^-) and phosphate (PO_4^{3-}) were measured on board using the automated colorimetric technique on a segmented flow Bran Luebbe autoanalyser II; ammonium (NH₄) determinations were performed on board by fluorometry on a Jasco FP-2020 fluorometer. For full method description see Pujo-Pay et al. (2011).

Microbial cell abundance 2.6

Abundances of the following microbial populations were assessed using flow cytometry: heterotrophic bacteria, Synechococcus, Prochlorococcus, picoeukaryotic phyotoplankton, and nanoeukaryotic phytoplankton. Total cells are presented as a sum of these groups. Full methods for flow cytometry enumeration of both autotrophic and heterotrophic cells are provided in Talarmin et al. (2011) and Tanaka et al. (2011).

Chlorophyll 2.7

Depth profiles across the transect were sampled for pigments by filtering 2 L of seawater onto 25 mm GF/F filters, which were frozen until extraction. HPLC analysis was used to quantify pigments, see Ras et al. (2008) for full analytical methods. Total **BGD**

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Statistical treatment of data 2.8

Averages are reported ± one standard deviation. All reported correlations are based on linear regressions. Significant differences in total IP-DAGs and IP-DAG ratios across different nutrient amendments in the microcosm incubations (Fig. 6) were identified by ANOVA and post-hoc Tukey-Kramer HSD tests using MatLab software (version R2009b), at a confidence level of 95 %. Principal component analysis (PCA) was conducted using MatLab software (version R2009b) following procedures and code outlined in Glover et al. (2011). The purpose of the PCA was to determine if the variability of individual IP-DAGs and total IP-DAGs was similar to the variability of nutrients (PO₄, NO_3 , NH_4), pigments (chl a), or microbial groups (*Prochlorococcus*, *Synechococcus*, heterotrophic bacteria, picoeukaryotic phytoplankton, nanoeukaryotic phytoplankton, total cells as the sum of these groups). Accordingly a covariance matrix was constructed as the minor product of twenty standardized parameters; 76 samples were included where each parameter was measured at the same depth from the same CTD cast. Two principal component factors were selected which captured the greatest variability in the data, and the loading of each parameter on the varimax rotation of these two factors is presented. Loadings were considered notable when their absolute values were greater than 0.6.

Results

Environmental conditions across the Mediterranean

The surface waters of the Mediterranean basins were distinguished by their variation of salinity (Figs. 1 and 2a): relatively low salinity in the Algero-Provencal basin in the

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west (stations A, 21), and across the Sicilian channel (stations 19 and 15) and the Sicilian strait (station 17), increasing salinity through the Ionian basin (stations 13, B, 1, 3), and highest salinity in the Levantine basin in the east (stations 5, 7, 9, 11, C). Intense summer stratification was apparent in the potential density (σ_{θ} , Fig. 2b), which also displayed downwelling at anticyclonic eddies near stations A, B, 9 and C. There was a west-east gradient in phosphate concentration with high phosphate in the west. The phosphocline was shallowest through the Sicilian channel (Fig. 2c) and deepened to the east (Pujo-Pay et al., 2011). Further discussion of the oceanographic conditions during the BOUM campaign were described by Moutin et al. (2011).

3.2 Distribution of IP-DAGs across the Mediterranean

The IP-DAG membrane lipid composition of particulate material was analyzed at 14 stations across the Mediterranean. In each sample the nine most abundant classes of marine IP-DAGs (Van Mooy and Fredricks, 2010) were quantified: three phospholipids, PG, PE, PC; three glycolipids, SQDG, MGDG, DGDG; and three betaine lipids, DGTA, DGTS, DGCC (Fig. 3). Total IP-DAG concentration, as the sum of these nine classes, averaged $2.4\pm1.5\,\mathrm{nM}$ in the upper 250 m of the Mediterranean. The general distribution of total IP-DAGs paralleled that of total cells (Fig. 2), with a significant positive linear correlation ($R^2=0.64$, n=159, p<0.01). The average IP-DAG content per cell was found to be 6.6 ± 2.4 amol/cell, or approximately $3.7\pm1.3\,\mathrm{fg}$ lipid C/cell. Total lipid concentration was highest in the western Mediterranean with a maximum of $6.3\,\mathrm{nM}$ at approximately 50 m in the Sicilian channel at station 15. The depth of the total IP-DAG maximum deepened from west to east, reaching 100 m in parts of the Levantine basin and a maximum concentration of only $4.3\,\mathrm{nM}$ in the east.

The distribution of the total IP-DAGs by depth showed low concentration in the surface (1.0–5.0 nM), a maximum around 50–100 m (2.5–6.0 nM), and decreasing concentration down to 150 m (1.0–3.0 nM), then relatively low and constant down to 250 m (0.3–0.7 nM; Fig. 2). Near the surface, total IP-DAGs were dominated by the glycolipids (Fig. 3), closely followed by phospholipids. Averaging across the whole transect,

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from the surface to 150 m depth the most abundant IP-DAG was the glycolipid SQDG, roughly 30% of the total IP-DAGs at many depths, with a maximum of 2.5 nM at 50 m in the Sicilian strait. Below 150 m the phospholipid PE was most abundant.

The distribution of each lipid class varied both longitudinally and with depth, however the classes within each type of IP-DAG (phospholipids, glycolipids, betaine lipids) tended to have similar characteristics. This was particularly clear in the trends with depth: at depths below approximately 100 m betaine lipid concentrations decreased sharply; at depths below 150 m glycolipid concentrations decreased steeply, thus at depths greater than 150 m phospholipids dominated the reservoir of IP-DAGs (Fig. 3).

3.3 Variation within types of lipids

3.3.1 Phospholipids

Phospholipids demonstrated a strong longitudinal gradient, with a clear maximum in the west and decreasing concentrations to the east (Fig. 3a,b,c). The phosphate concentration was also higher in the west and decreased to the east (Fig. 2c). As a fraction of total IP-DAGs, total phospholipid concentration (the sum of PG, PE and PC) was significantly positively correlated with phosphate concentration ($R^2 = 0.30$, n = 198, p < 0.01). The longitudinal and depth distribution of the three phospholipids, PG, PE, and PC, was quite similar (Fig. 3a, b, c). By depth, each phospholipid class was approximately 200-600 pM in the surface, increasing to 200-800 pM at 50-75 m, then decreasing to 100-200 pM at 250 m. Though the basin-wide distribution of the three phospholipids was similar, their relative abundance varied with depth. From 5 m to 100 m, PC was the most abundant phospholipid, and in many places the second most abundant IP-DAG after SQDG. At depths below 100 m PE was the most abundant phospholipid; and below 150 m PE was the most abundant IP-DAG. Maxima occurred at station 17 for PG and PE; station 21 for PC; at station 9 high concentrations of all three phospholipids persisted much deeper than the surrounding areas consistent with downwelling seen in isolines (Fig. 2). At several locations samples were taken from 250 m down to 2500

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In contrast to the phospholipids, the glycolipids did not display a strong longitudinal gradient, nor was their distribution as similar across the three classes of IP-DAGs. By far the most abundant glycolipid was SQDG (Fig. 3). Its vertical distribution showed a distinct minimum in concentration near the surface, an increase at 50-100 m, and a sharp decrease at 100 m in the Algero-Provencal basin in the west, a slightly deeper decline at 150 m to the east. The distributions of MGDG and DGDG were more similar to each other than to SQDG, being less abundant overall (~200 pM near the surface), and having less pronounced minimums near the surface. DGDG was a little less abundant than MGDG, though MGDG and DGDG both had maxima at 75 m at station A, roughly 600 and 400 pM, respectively.

m (data not shown); below approximately 175 m all of the detectable IP-DAGS were phospholipids, which persisted in measurable quantities down to 2700 m (the deepest

depth sampled). At 500 m and deeper, each phospholipid was roughly 50-100 pM, with

PE being most abundant (40 % of total IP-DAGs).

3.3.3 **Betaine lipids**

5 3.3.2 Glycolipids

The three betaine lipids had similar patterns of distribution, but their absolute concentrations were quite different with DGTA having the highest concentration by a factor of 2 at most locations, a maximum of almost 900 pM compared to roughly 300 pM for DGTS and DGCC. For all three betaine lipids the highest concentration was in the Sicilian channel (stations 19 and 17), with a second maximum in the eastern Mediterranean at the boundary of the Ionian and Levantine basins (stations 3, 5 and 7). Depth profiles of the betaine lipids were different from the phospholipids and glycolipids in several ways: they did not exhibit a pronounced minimum near the surface; the maximum was slightly shallower, 50–75 m as opposed to 50–100 m; and their concentration decreased more steeply with increasing depth than either glycolipids or phospholipids.

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by 100 m depth their concentration was near undetectable.

3.4 Ratio of IP-DAGs

From west to east across the Mediterranean, the strong gradient of the phospholipids from high to low led to a gradient in the ratio of non-phosphorus lipids to phospholipids (Fig. 4). The ratio of the sulfolipid SQDG to the phospholipid PG (SQDG:PG; Fig. 4a) was low in the western Algero-Provencal basin $(1.6\pm0.3 \text{ in the upper } 100 \text{ m})$, began to increase across the Sicilian channel $(1.9\pm0.7 \text{ in the upper } 100 \text{ m})$ into the Ionian basin $(2.5\pm1.3 \text{ in the upper } 150 \text{ m})$, and was highest in the eastern Levantine basin $(3.1\pm1.6 \text{ in the upper } 150 \text{ m})$. The maximum deepened from west to east, from 75 m in the Algero-Provencal basin to 125 m in the Levantine basin.

The ratio of the betaine lipids to PC also exhibited a longitudinal gradient (betaine lipids:PC; Fig. 4b), being lowest in the west $(1.0\pm0.5 \text{ in the upper } 50 \text{ m}$ in the Algero-Provencal basin) and highest in the east $(2.7\pm0.9 \text{ in the upper } 75 \text{ m}$ in the Levantine basin). The ratio approached zero at a much shallower depth than SQDG:PG, due to the shallower depths of the betaine lipids relative to SQDG (Fig. 3). The maximum also deepened from west to east, more steeply than SQDG:PG, from near 5 m in the Sicilian channel to 50 m in the Levantine basin.

Across the Mediterranean there was a strong positive correlation between SQDG and chlorophyll a ($R^2 = 0.50$, n = 135, p < 0.01). The maxima of both SQDG and chlorophyll a occured at depth (Figs. 2 and 3), between 50 and 100 m (shallower in the western basin, deeper in the eastern basins), however the ratio of SQDG to chlorophyll a was highest near the surface (Fig. 4c). The deep chlorophyll maximum paralleled the abundance of Prochlorococcus (see Mella-Flores et al. (2011) for cyanobacteria abundance), and while the maximum of SQDG also followed the Prochlorococcus abundance, SQDG was also relatively abundant near the surface (Fig. 3d). There was not a longitudinal gradient in SQDG:chl a, with the maxima of the ratio in the Sicilian strait (station 15) and across the boundary of the Ionian and Levantine basins (stations 3 and 5).

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Relationship between IP-DAGs and other biogeochemical parameters

Principal component analysis (PCA) showed that total cells, heterotrophic bacteria, and total IP-DAGs had similar loadings on both of the two most heavily loaded principal component factors (factors 1 and 2, Fig. 5). Almost all of the phospholipids and glycolipids also had similar, heavy loadings on factor 1, except for DGDG which loaded heavily on factor 2. Additionally most of the microbial populations loaded closely to the glycolipids and phospholipids on factor 1, including heterotrophic bacteria, Prochlorococcus, nanoeukaryotic phytoplankton and picoeukaryotic phytoplankton. Synechococcus did not load closely to the other microbial populations on factor 1, though its loadings on both factors 1 and 2 were similar to DGDG. Total chlorophyll a had similar loading to Prochlorococcus. The nutrients NH₄, NO₃, and PO₄ did not have loadings similar to any of the cells, chlorophyll a, glycolipids or phospholipids in this analysis. However, NO₃ and PO₄ had heavy positive loadings on factor 2, while the three betaine lipids had a heavy negative loading on factor 2, and both nutrients and betaine lipids had similarly little loading on factor 1.

Response of IP-DAGs to nutrient amendments in microcosm incubations

Microcosm incubations were conducted at three locations (stations A, B, and C) in the Mediterranean to determine the response of the biological community to changes in nutrient availability, specifically an increase in available ammonium (N condition), phosphate (P condition), or an increase in both ammonium and phosphate (NP condition). Over three or four days an array of biogeochemical parameters were measured to assess community response (see Tanaka 2011 for a full description of results). Comparison of IP-DAGs in the different nutrient amendments on the final day of the incubation showed that within the space of a few days the membrane lipid composition had changed dramatically.

Total IP-DAG concentration increased dramatically in the NP condition in all three experiments (stations A, B and C) relative to the control. With the addition of only **BGD**

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ammonium or only phosphorus there were not significant changes in total IP-DAGs relative to the control (Fig. 6a-c).

In contrast to the change in total IP-DAG concentration, the ratio of SQDG to PG increased in the N condition at all three stations. At station B there was an also a large increase in SQDG:PG in the NP condition. At station C there was a small decrease in SQDG:PG in the P condition as well as the NP condition. Shifts in betaine lipid:PC ratio in the N conditions were similar to those observed in SQDG:PG, with an increase relative to the control at stations B and C (changes were not significant at station A). In contrast to SQDG:PG there also appeared to be an increase in betaine lipids:PC in the NP condition at all stations, however the variability across the three triplicates was slightly larger for the betaine:PC ratios than for SQDG:PG and thus the increase in the NP condition was only significant at station C. In the P condition there were no significant differences in betaine lipids:PC relative to the control.

The ratio of SQDG to chlorophyll *a* also changed with nutrient additions, with a decrease in the ratio in both the N and NP conditions relative to the control at all stations. At station C there was also a decrease in the P condition, and no significant change in the P condition at the other two stations.

4 Discussion

Membrane lipids have a rich history of being applied as a proxy for microbial biomass in the environment (Balkwill et al., 1988, White et al., 1979), and recent work has shown that they may also be applied as indicators of community structure and physiology (Popendorf et al., 2011; Van Mooy et al., 2006, 2009). In this study, across the Mediterranean total cell abundance was highest in the west (Fig. 2) as expected from previous studies (Siokou-Frangou et al., 2010 and references therein), and total IP-DAGs were significantly positively correlated with total cells, reaffirming the application of lipids as a measure of biomass. Among the different IP-DAGs the strongest west to east gradient was seen in the phospholipids. When normalized to total IP-DAGs,

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the gradient of phospholipids was positively correlated with phosphate concentration, with phospholipids making up less of the membrane lipids in the ultraoligotrophic east. This aligns with the hypothesis that microbes will direct less of their cellular phosphate towards phospholipid production when the nutrient is increasingly scarce (Van Mooy et ₅ al., 2006, 2009). With less phospholipids in the membrane, the balance of IP-DAGs is expected to come from glycolipids and betaine lipids. The particular substitutions that have been observed in cultures are SQDG substituting for PG, and betaine lipids substituting for PC (Benning et al., 1993, 1995; Martin et al., 2011; Van Mooy et al., 2006, 2009). Across the Mediterranean, the ratios of SQDG:PG and betaine lipids:PC were both highest in the east. This gradient in membrane composition corresponded with the change in nutrients across the Mediterranean (Pujo-Pay et al., 2011), however there are two potential mechanisms that could contribute to this variation in IP-DAGs: (1) physiological shifts within organisms, leading to increased non-phosphorus lipids per cell, or (2) changes in the composition of the microbial community leading to more abundant low-phospholipid containing organisms (for example Synechoccocus; Popendorf et al., 2011; Van Mooy et al., 2006) when phosphorus is scarce. By combining measurements of IP-DAGs across the Mediterranean with measurements in microcosm incubations we were able to deconvolve the relative contribution of these two mechanistic influences.

4.1 Factors influencing IP-DAG ratios

The relative influence of community composition and physiological response to nutrients was assessed by principal component analysis (Fig. 5), comparing the variability in IP-DAG concentrations to the variability in nutrients and variability in the abundances of different planktonic groups. All of the microbial groups, except for Synechococcus, loaded heavily on factor 1 and only minimally on factor 2, thus factor 1 loads heavily on parameters representing community structure. Additionally, all of the phospholipids and glycolipids, except DGDG, load heavily on factor 1 and not heavily on factor 2, indicating that much of the variability in phospholipids and glycolipids across the transect

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might be associated with the variability in the planktonic community. *Synechococcus* and DGDG did not load heavily on factor 1, but they loaded closely to each other on factor 2. Previous studies have shown that DGDG is a significant component of *Synechococcus* membranes and is not as abundant in other cyanobacterial or heterotrophic bacterial membranes (Popendorf et al., 2011). Furthermore, the distribution across the transect showed that maxima of both DGDG and MGDG corresponded with high abundances of *Synechococcus* and *Prochlorococcus* respectively (see Mella-Flores et al. (2011) for the distribution of cyanobacteria) reinforcing the relevance of the PCA for the observed trends across the Mediterranean. The similar loading of *Procholorococcus* and chlorophyll *a* in the PCA also corresponded with the observed co-occurrence of the deep chlorophyll maximum and *Prochlorococcus* maximum across the Mediterranean. Similar associations between *Prochlorococcus* and chlorophyll *a* were previ-

ously observed in the South Pacific (Van Mooy and Fredricks, 2010).

The other parameters that did not load heavily on factor 1 were nutrient concentrations and betaine lipid concentrations, which instead loaded heavily on factor 2. This may indicate that across the Mediterranean the concentration of betaine lipids was heavily influenced by the concentration of nutrients, implying a stronger signal of physiological response as opposed to variation in community structure. Though principal component analysis is a useful tool for indicating potential connections it cannot determine if community or physiology are causal in the variability of IP-DAGs. The variation of IP-DAGs with different microbial groups has been well studied in cultures (Oliver and Colwell, 1973; Guschina and Harwood, 2006; Kato et al., 1996; Van Mooy et al., 2006), and some work has also been done measuring the diversity of IP-DAGs in microbial groups from environmental samples (Popendorf et al., 2011). The physiological response has also been well studied in monocultures (Benning et al., 1993, 1995; Martin et al., 2010; Van Mooy et al., 2009), however it remains to be shown whether the influence of changes in physiology can be detected in changes in IP-DAGs in a natural, mixed planktonic community.

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4.2 Influence of physiology vs. community on IP-DAGs assessed in microcosm incubations

The microcosm incubations with nutrient amendments conducted on the BOUM cruise provided a unique opportunity for assessing the relative contributions of community composition and physiological response to the changes in IP-DAGs. Within the microcosms, we could account for changes in the community composition, enabling an examination of the changes in IP-DAGs that resulted from solely physiological adaptation. The addition of nitrogen and phosphorus to the oligotrophic, phosphorus-depleted waters of the Mediterranean elicited a diverse array of both community and physiologic shifts in the microbial community.

With the addition of phosphate (P condition) in the microcosms no significant changes were observed in the abundance of various microbial groups (Tanaka et al., 2011). With the lack of change in community composition, the observed change of IP-DAGs in this condition would be expected to be primarily due to physiological response. However, primary production, bacterial production, and alkaline phosphate activity did not change in response to the P amendment (Tanaka et al., 2011; Lagaria et al., 2010), which indicates that this system was not phosphate limited despite the low ambient phosphate concentration and high N:P at all three stations (Pujo-Pay et al., 2011). Relative to the control, phosphate addition elicited a change in the ratio of IP-DAGs at only one station: a small decrease in SQDG:PG at station C, as would be expected of a physiological response to increased phosphate. At all other stations there was no significant change in the ratio of SQDG:PG or betaine lipids:PC. This lack of strong response to phosphorus addition is consistent with the observations of a previous phosphate addition experiment in the Mediterranean (Thingstad et al., 2005), which found that increased phosphate did not elicit a change in the phytoplankton community despite the expectation of phosphate limitation. The lack of observed shifts in IP-DAG ratios simply reflects a community that was not P-limited.

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The addition of ammonium (N condition), on the other hand, elicited a strong physiologic response. In the N condition alone there was a significant increase in alkaline phosphatase activity at all three stations (Tanaka et al., 2011), demonstrating clear enhancement of phosphorus stress (Lomas et al., 2010). The changes in IP-DAG ratios that were observed are what would be expected from a physiologic response to increased phosphorus stress: at all three stations the ratio of SQDG:PG increased, and at two stations the ratio of betaine lipids:PC increased relative to the control (changes were not significant at station A, although the mean was higher). These changes were consistent across the stations, despite inconsistent changes in the community structure which were generally small relative to changes observed in the NP condition (Tanaka et al., 2011). Thus these changes in IP-DAG ratios indicate that the increase in phosphorus stress led to a dynamic, physiologic cellular response resulting in IP-DAG substitution in a natural, mixed planktonic community. This is the first observation of a change in membrane composition as a physiological response in the environment, and it indicates that elevated ratios of these IP-DAGs observed in other phosphorus limited environments could be an indicator of phosphorus stress.

The increase of the betaine lipid:PC ratio in the N condition provided validation of the lipid substitution expected from observations in culture (Benning et al., 1995; Martin et al., 2010). This was particularly notable since betaine lipids in marine environments are not well understood (Van Mooy and Fredricks, 2010; Popendorf et al., 2011) and therefore mechanistic explanations for the observed trends are more limited. In all of the amendment conditions the change in betaine lipids:PC was more variable within the triplicate measurements than the change in SQDG:PG. The PCA indicated that betaine lipids might be more strongly influenced by physiological response than the glycolipids or phospholipids (Fig. 5), and the longitudinal trend in the betaine:PC ratio showed an increase in the ultraoligotrophic eastern Mediterranean Sea (Fig. 4), indicating an increase in response to low phosphate conditions as expected. However the sharp decrease in betaine lipid abundance with depth, relative to glycolipids and phospholipids, indicates that the abundance of these IP-DAGs may be controlled by unique

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factors, related either to physiology or community, which are thus far unidentified. The microcosm incubations support the conclusion that physiological response contributes to the observed trends in betaine lipid concentration, but future studies could further refine our understanding of betaine lipid sources and the factors that influence them.

In the NP condition there was a significant change in both the community composition and metabolic activity at all three stations. These changes included an increase in primary production at all three stations, an increase in bacterial production at stations B and C, and an increase in the abundance of many microbial groups including *Synechococcus* (all stations), autotrophic nanoplankton (stations B and C), and ciliates (stations A and C; Tanaka et al., 2011; Lagaria et al., 2010). The total IP-DAG concentration increased significantly at all three stations, as would be expected from the large increase in microbial abundance. The significant shifts in community structure, however, meant that little insight on physiology could be drawn from the observed changes in IP-DAG ratios.

4.3 Relationship between SQDG and chlorophyll a

Previous studies have suggested that SQDG originates solely in photoautotrophic organisms (Popendorf et al., 2011; Van Mooy et al., 2006, 2009) and is associated with the organelles of photosynthesis (Sakurai et al., 2006; Sato, 2004; Wada and Murata 1998). Thus we would predict that across the Mediterranean there would be a strong correlation between SQDG and chlorophyll *a*, and indeed this was observed. The maxima of both SQDG and chlorophyll *a* occurred at depth, however the ratio of SQDG to chlorophyll *a* was highest near the surface. While the maximum of SQDG paralleled the maximum in *Prochlorococcus* abundance (Mella-Flores et al., 2011), SQDG is known to also occur in other phytoplankton including *Synechoccocus* and pico- and nanoeukaryotic phytoplankton (Popendorf et al., 2011; Sato, 2004; Van Mooy et al., 2006, 2009), which were more abundant near the surface than at the deep chlorophyll maximum (Mella-Flores et al., 2011). The increase in the ratio of SQDG to chlorophyll *a* near the surface therefore indicates that though both molecules correlate with phyto-

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plankton abundance, their ratio varies with depth. This may be due to a combination of both photoacclimation, decreasing cellular chlorophyll content with high surface irradiance (McIntyre et al., 2002), or higher cellular SQDG content in the phytoplankton

Though no longitudinal gradient was observed in SQDG:chl a, the change in the ratio in the microcosm incubations demonstrated a response to changes in nutrients. In the N and NP conditions at all stations the ratio of SQDG to chlorophyll a decreased significantly relative to the control (at station C the ratio decreased relative to the control in all amendment conditions). This change in the N condition is likely representative of a physiological response (as previously discussed the changes in the NP condition are likely due to a combination of community changes and physiological response). Chlorophyll has a high nitrogen content, and is known to increase proportionally more than phytoplankton abundance when nitrogen is added to an N-starved system (Zohary et al., 2005). The addition of ammonium in these incubations led to an increase in chlorophyll a in the N condition relative to the control at stations A and B (Tanaka et al., 2011), and, though the ratio of SQDG:PG was seen to increase in these conditions, the ratio of SQDG:chl a decreased, indicating that this ratio is sensitive to nitrogen abundance. Thus the ratios of SQDG:chl a, as an indicator of nitrogen availibility, and SQDG:PG, as an indicator of phosphorus stress, may be complementary indicators of planktonic nutrient stress in oligotrophic marine environments.

species that are more abundant near the surface.

Conclusions

We observed gradients in IP-DAGs that correlated with the phosphate gradient across the Mediterranean, which is consistent with the hypothesis that nutrient availability controls the relative abundance of IP-DAGs. Gradients in community structure undoubtedly contributed to this gradient, but our microcosm incubations showed that physiological shifts likely also contribute to the observed changes. The phospholipid concentration as a percent of total lipids was positively correlated with phosphate concentra**BGD**

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tion across the Mediterranean, both phospholipids and phosphate concentration being higher in the west and lower in the east. Glycolipids and betaine lipids did not show a strong west to east gradient, however the ratio of non-phosphorus lipids to phospholipids varied across the Mediterranean, with SQDG:PG and betaines:PC being higher in the ultraoligotrophic eastern Mediterranean. Throughout the Mediterranean, total IP-DAGs in the upper 250 m were correlated with cell abundance, reaffirming their direct representation of microbial biomass in the ocean. Comparison of the variation in IP-DAGs to variations in microbial groups and nutrients across the Mediterranean was used to examine the relative influence of microbial community composition changes and cellular physiology changes on the observed IP-DAG changes. Microcosm incubations of whole seawater with nutrient amendments provided further insight on these two mechanisms, and provided direct evidence from a mixed community under in situ conditions that shifts in IP-DAG ratios are part of a physiologic response to changes in nutrients. Additionally, the microcosm incubations demonstrated that an increase in ammonium led to a decrease in the ratio of SQDG to chl a, indicating that this ratio may be an indicator of nitrogen availability. With knowledge of the factors influencing the variation of IP-DAGs in the environment, IP-DAGs can be molecular tools for the study of the microbes in the ocean and their role in biogeochemical cycles.

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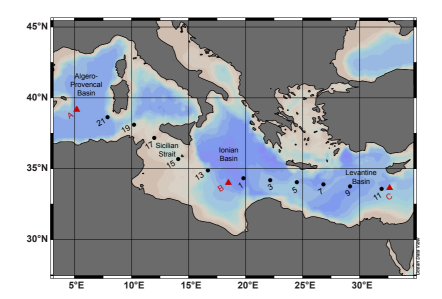


Fig. 1. Map of the Mediterranean Sea, colored to indicate bathymetry, with locations of sampling on the BOUM cruise. Numbered stations (black circles) denote locations where a single depth profile was sampled; lettered stations (red triangles) denote locations where multiple profiles were sampled in near proximity, and where microcosm incubations were conducted.

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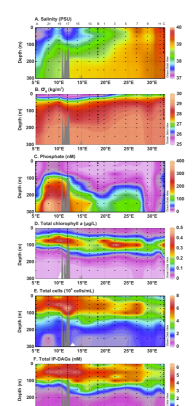


Fig. 2. Distribution of several parameters measured in the upper water column (300 m) from west to east across the Mediterranean Sea. Salinity (A) and potential density (σ_{Θ}) (B) were measured with the CTD sensors; phosphate (C) is soluble reactive phosphorus as measured by an autoanalyzer instrument; total chlorophyll a (D) is the sum of chlorophyll a, divinyl chlorophyll a and chlorophyllide a; total cells (E) is the sum of *Prochlorococcus*, *Synechococcus*, heterotrophic bacteria, picoeukaryotic phytoplankton, and nanoeukaryotic phytoplankton; total IP-DAGs (F) is the sum of the nine most abundant classes of IP-DAGs found in the water column. Numbers across the top panel indicate station labels; black dots indicate individual samples, colored contours indicate interpolated gradients.

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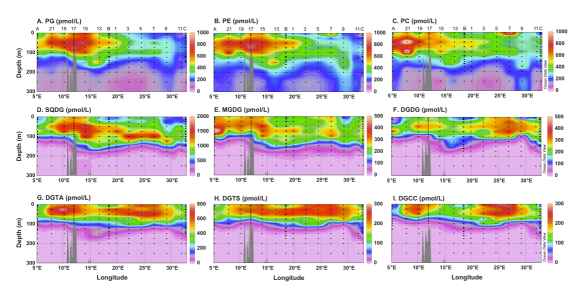


Fig. 3. Distribution of the nine most abundant IP-DAGs in the upper water column (300 m), each panel showing the concentration of a single class of IP-DAG from west to east across the Mediterranean Sea. The first row (A–C) is phospholipids; the second row (D–F) is glycolipids; the third row (G–I) is betaine lipids. Numbers across the top panels indicate station labels; black dots indicate individual samples, colored contours indicate interpolated gradients.

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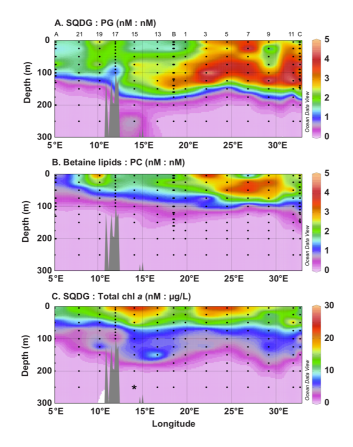


Fig. 4. Distribution of the ratio of the sulfolipid SQDG to the phospholipid PG **(A)**, the ratio of the sum of the betaine lipids to the phospholipid PC **(B)**, and the ratio of SQDG to total chlorophyll *a* **(C)**, each shown in the upper 300 m from west to east across the Mediterranean Sea. Numbers across the top panel indicate station labels; black dots indicate individual samples, colored contours indicate interpolated gradients; * indicates one outlier value not plotted.

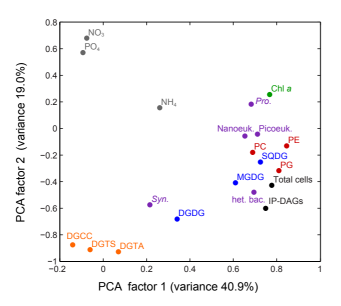


Fig. 5. Principal component analysis was applied to 20 parameters: concentration of three phospholipids (PG, PE, PC; red circles); concentration of three glycolipids (SQDG, MGDG, DGDG; blue circles); concentration of three betaine lipids (DGTA, DGTS, DGCC; orange circles); abundance of five plankton populations (purple circles): *Synechococcus* (*Syn.*), *Prochlorococcus* (*Pro.*), nanoeukaryotic phytoplankton (nanoeuk.), picoeukaryotic phytoplankton (picoeuk.), heterotrophic bacteria (het. bac.); concentration of three nutrients (grey circles); concentration of total chlorophyll *a* (Chl *a*; green circle); and total cell abundance and total IP-DAG concentration (black circles). Each parameter is plotted as its loading on the two principal component factors that captured the greatest amount of variance in the data.

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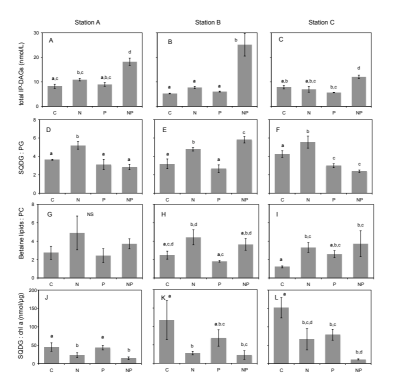


Fig. 6. Total IP-DAG concentration **(A–C)**, SQDG:PG ratio **(D–F)**, betaine lipid: PC ratio **(G–I)**, and SQDG: chlorophyll a ratio **(J–L)** on the final day of incubations from three locations, stations A, B, and C (organized by column) with nutrient amendments: C is control (no amendment), N is addition of ammonium, P is addition of phosphate, NP is addition of ammonium and phosphate. Error bars are the standard deviation of triplicate carboys. In each plot, lowercase letters (a, b, c, d) indicate statistical post-hoc comparisons: bars labeled with the same letter are significantly different from those without the letter (p<0.05); letters in boldface (a) indicate comparison to the control; NS indicates no significant difference.

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