

Reply to Referee #1 comment made on bg-2020-300, “Cross-basin differences in the nutrient assimilation characteristics of induced phytoplankton blooms in the subtropical Pacific waters” by Hashihama et al.

Review: The manuscript by Hashihama et al. reports on the results of a set of shipboard incubation experiments in which surface seawater collected at seven stations across the subtropical North and South Pacific ocean was amended with deep water. The additions induced phytoplankton blooms and concomitant nutrient drawdowns in all seven experiments. The novelty of the study resides mainly on the use of nutrient nanomolar measurements to reveal regional patterns in nutrient drawdown ratios. The authors use also the obtained drawdown ratios in an attempt to contribute to current knowledge on the factors that control phosphate distribution in the subtropical Pacific Ocean.

Overall, the manuscript is very well organized and written. The introduction is fairly complete and presents nicely the context of the study. The methodology is well developed and it did not raise any major concerns from my side. The outcome is very interesting and will certainly contribute to the field of nutrient biogeochemistry in the oligotrophic ocean. The study constitutes a nice illustration on how nanomolar measurements of nutrients can give exciting insights on nutrient cycling. I do have a few comments that may contribute to clarify some aspects of the manuscript.

Our reply (in italics hereafter): We appreciate your constructive comments on our manuscript. We have addressed your comments as seen below.

General comments:

1. A recent paper published by the authors (Hashihama et al. 2020, GBC) show data from the same cruises presented in this study. If I am not wrong, this paper is referenced as Hashihama et al (submitted) at some points in the discussion section. Now that this paper has been published, and given the complementarity with this one, I strongly recommend the authors to refer to it to better put into context their outcomes. For instance, it would be interesting to check the influence of the experimental bloom-derived drawdown ratios on regional patterns of P cell quotas, addressed approximately through chlorophyll or phytoplankton biomass normalized POP data.

Yes, we have referred Hashihama et al. (2020, GBC) in the revised manuscript. For the suggested analysis using POP data, we have looked at the ambient data for Chl a and POP (TPP) in the study area of Hashihama et al. (2020, GBC). However, data on Chl a-normalized TPP in the mixed layer did not show a clear regional pattern (no distinctly high TPP/Chl a values in the western North Pacific). Since cell-specific content of Chl a varies depending on seasonal and/or regional light intensity, the use of Chl a-normalized TPP data is unsuitable for the suggested analysis. We consider

that direct observation of particulate C-N-P variability response to deep water addition will be necessary soon.

2. The only methodological issue that raised some concern to me was the differences in incubation times among the experiments since they can affect the drawdown ratios. This is particularly true when DIN and PO₄ decreases are not linear (i.e. station 15, figure 3). I suggest the authors to conduct their calculations using only data obtained during the first 48h to see if the outcomes still stand. Otherwise, please add a statement in the discussion on the fact that different incubation times might have affected the obtained differences in nutrient drawdown ratios.

We have compared slopes of the nutrient decreases during the first 36-48 h, and for the full incubation periods (52-96 h), and both the slopes were not significantly different for DIN or PO₄ (paired t-test, p>0.05). We have added the related statement in the revised manuscript (L223-224).

3. The differences among experiments in nutrient drawdowns and their ratios are sometimes small. In order to add robustness to the interpretation of these differences, the authors should calculate the errors associated to these observations (errors of the slopes of linear regressions in Figs 3a and 3b and errors of calculated Δ DIN and Δ PO₄). More generally, I missed standard errors and/or estimated uncertainties of all data and/or calculations throughout the manuscript.

We have added the errors (95% confidence interval) for the slopes and intercepts of regressions and the Δ values and its ratios in the revised manuscript. The errors of the drawdown ratios were calculated based on the error propagation rule (Miller and Miller, 1993 Statistics for Analytical Chemistry, 2nd edn) as stated in L121-122. In this calculation, the mean drawdown ratios changed slightly from those in the previous manuscript due to the use of different significant digits, but it did not affect the conclusion.

4. I would like to share with the authors some thoughts that came to my mind when reading their manuscript related to the influence of their results on P* regional distribution. I am not sure of the pertinence and accuracy of these thoughts, so I share them just in case they can feed the interpretation of their outcomes. In 4.4 section and in Figure 6, the authors make a parallelism between ambient P* and “bloom P*”. This exercise is interesting and confusing at the same time. The authors state that the observed negative values of bloom P* (not observed in in situ P*) imply the presence of alternative P sources. But isn't this statement only true under the assumption of 16:1 drawdown ratios used to calculate ambient P*? Wouldn't the observed differences in ambient and bloom P* values question the use of 16:1 to estimate ambient P*? This would also be illustrated by the much higher variability in ambient P* compared to bloom P*, wouldn't it?

Apologies, we had stated a slightly confusing discussion in 4.4 section. In the revised manuscript, we

have stated a simple discussion as “The difference between the bloom P and ambient P* largely depends on the different N:P consumption ratios of ≤ 13.3 and 16, respectively. If the low N:P consumption ratios (≤ 13.3) are consistently dominant in the PO₄-depleted WNP, alternative P sources other than PO₄ would be required to fully exhaust DIN. Since lower DOP concentrations and higher alkaline phosphatase activity were observed in the WNP, compared to other subtropical Pacific regions (Hashihama et al., 2019; Hashihama et al., 2020), active DOP utilization in the WNP likely contributes to the DIN exhaustion. These perspectives suggest that, in the studies on subtropical nutrient biogeochemistry using N:P stoichiometry, the bioavailable fraction of DOP could be an important factor as well as DIN and PO₄.” (L413-420)*

Specific comments:

- Page 5, lines 125-126. When available, it would be useful to give the standard deviation of the triplicate analytical data.

The standard deviations have indicated in Table 2.

- Page 6, lines 168-169. The observed trend would be due to the low variability in DIN concentration rather than to low DIN concentration itself.

We have added “the consistently low concentrations of DIN” in the revised manuscript (L172).

- Page 11, line 219. The term ‘net assimilation rates’ is normally associated to tracer incorporation measurements which is not the case. Please, stick to the term drawdown rates.

We have used “drawdown rates” throughout the text.

- Page 11, line 220. Please, add estimated errors of these slope values (cf general comment 3 above)

Yes, we have added as mentioned above.

- Page 11, lines 225-226 and page 13, line 253. I do not understand what these R² values mean, please clarify.

These values were derived from the linear regression analysis. We have added “r²=xxx in linear regression” in these parts (L230; L259).

Reply to Referee #2 comment made on bg-2020-300, “Cross-basin differences in the nutrient assimilation characteristics of induced phytoplankton blooms in the subtropical Pacific waters” by Hashihama et al.

Comments to Author(s): This manuscript by Hashihama et al. describes the variation in macronutrient drawdown among Pacific Ocean surface microbial communities using deep water additions to bottle incubations. The authors present data from seven subtropical gyre sites with distinct nutrient uptake ratios, where the nutrient limiting net biological production is unknown. These observations are linked to pigment proxies for phytoplankton taxa and diatom densities to examine the role of community in nutrient drawdown ratios. The experiments yielded increased phytoplankton biomass at all sites, but varying stoichiometric ratios of uptake for DIN:PO₄:Si(OH)₄. I feel this manuscript strongly expands upon existing studies on the question of nitrogen, phosphorus and iron limitation in the North and South Pacific Ocean.

The data is presented in a straightforward manner and explained well. All sections are well written and figures are easily digested. I have a few concerns on assumptions made regarding phytoplankton pigments proxies and deep DOM composition, but otherwise recommend the manuscript should be accepted with minor revisions.

Our reply (in italics hereafter): Thank you for your constructive comments. We have addressed your comments as seen below.

General comments

1. Regarding the methods, I request that the deep water collection be clarified slightly. It is not stated if the water was filtered to remove living cells. This is of particular concern at Stations A and 2. If unfiltered seawater was used, both grazers and microbial cells could impact the conclusions at these stations. At other stations, freezing the seawater would remove this concern, but introduce additional nutrients from burst cells. The nutrient composition of cellular detritus is likely different and more bioavailable (urea, NH₄, labile DOM) than deep nutrients (NO₃, recalcitrant DOM).

We used the unfiltered deep water from 1500 m depth to avoid ammonium contamination from filtration process. This has been stated in the revised manuscript (L105-106). As you have considered, there might be a potential influence of heterotrophs in the deep water on the results of incubation experiments. However, concentration of particulate organic carbon in the deep layer below 1000 m is generally less than 10% of that in the surface layer (Hebel and Karl, 2001 DSR-II 48, 1669-1695; Yamada et al., 2017 MEPS 583, 81-93) and prokaryotic abundance and production exponentially decrease with depth (Yokokawa et al., 2013 LO 58, 61-73). Furthermore, the proportion of deep water

to the total incubated volume (surface water + deep water) was only 2.1% as stated in the section 2.2. The large dilution was also confirmed from T0 data of DON and DOP in the treated bottles which were not significantly different from those in the control bottles (paired t-test, $p > 0.05$, Fig. A2), indicating that the influence of labile DON and DOP additions were negligible. Thus, we conclude that the influences of heterotrophs and labile DOM supply by freezing were at negligible levels compared to the large enrichments of inorganic nutrients.

2. My largest concern is directly assuming that divinyl chlorophyll A concentrations are representative of Prochlorococcus abundances. While a useful indicator, the concentration of divinyl chlorophyll a could change between sites, season, light/depth level, etc. It is very possible Prochlorococcus cells in the Eastern South Pacific have a lower density of photosynthetic pigments, especially in the summertime at the surface where these cells were collected. In addition, since Nitrogen is limited, the cells may have adapted by lowering the concentration of N-rich photopigments further. This combined effect of photo-acclimation and adaptation to low N could explain the low divinyl chlorophyll A concentrations in the South Pacific subtropical gyre. This caveat should be acknowledged in the Discussion.

We agree that seasonal/regional PAR level influences the cellular pigment quotas. In contrast, N limitation occurred not only in the South Pacific but also in the North Pacific, because ambient DIN:PO₄ ratios (≤ 8) were much lower than the Redfield ratio (16) and subtropical particulate N:P ratio (28) (Martiny et al., 2013 Nat. Geosci. 6, 279-283). Therefore, N limitation might not be a robust reason for any seasonal/regional variations in the pigment concentrations, including DVchl a. Furthermore, we consider that pigment ratios (Tfuco:Zea (no N in either of the pigments) and DVchl a:Tchl a (both pigments contain N)) are useful for comparing the regional variations in phytoplankton composition even if seasonal/regional differences of photo-acclimation/adaptation and N-limitation occur. In the perspective of phytoplankton physiology, Tfuco and Zea play roles in light-harvesting and photoprotection, respectively (Falkowski, 2013 Aquatic Photosynthesis 2nd edn). Based on these roles, Zea content in cyanobacteria should be higher in the high-PAR South Pacific than the low-PAR North Pacific, while Tfuco content in eukaryotes should be higher in the low-PAR North Pacific than the high-PAR South Pacific. However, ambient Zea (Tfuco) concentrations were higher (lower) in the North Pacific than the South Pacific, indicating that the biomass proportion of cyanobacteria to eukaryotes was higher in the North Pacific than the South Pacific. Thus, we did not revise the statements for pigment ratios and seasonal/regional variations in phytoplankton composition.

3. Regarding DOM, I had two points to consider. The authors mention more bioavailable forms of DOM that may not be present in water at 1500m. Perhaps the DOM added then would not be consumed, leading to no net changes over the incubation period. Alternatively, the balance of net uptake and

release could yield no change. This possibility should be acknowledged for silicic acid as well considering the longer incubation time and high diatom abundances at station 15.

We also interpret that DON and DOP in the deep water did not have an influence on various parameters in the incubation, because these additions were negligible due to the large dilution as mentioned above. Since the resident phytoplankton in the DIN and PO₄-depleted subtropical ocean might have a high affinity to DON and DOP, rather than DIN and PO₄, we considered the possibility that phytoplankton in the incubated bottles consume labile DON and DOP in the surface water (not in the deep water) rather than DIN and PO₄. However, such a phenomenon did not occur in the present study and the uptake and release of DON and DOP were balanced. As you suggest, the uptake and release of silicic acid likely occurred. We have added the related statement in the revised manuscript (L380-381).

Specific comments

Line 47 - Changing 'alleviates temporarily' to 'temporarily alleviates' reads a bit better.

We have revised as you have suggested (L47).

Line 74 - 'observation' should be plural

Yes, it has been plural (L74).

Line 74-75 - I suggest changing ', to understand them, experimental validations are required.' to ', and to understand them experimental validations are required'.

We have revised based on your suggestion (L74-75).

Line 95 - See comment on methods

See our answer above.

Line 120 - To what extent does the water volume in the bottle change between T0 and the last time point? Also are collection intervals than shorter for the shorter incubation times?

At the end point, water volumes in the bottles just prior to final sampling were approximately 1.8 L, because the subsamples for nanomolar nutrients and DON/P were collected from the initial volume of 2.35 L (surface water + deep water). Also, the collection intervals were shorter for the shorter incubation times. We subsampled 5-6 times for nanomolar nutrients and 3 times for DON and DOP during the incubation periods for 48-96 h. We have added these statements in the revised manuscript (L114-117).

Line 199 - 'Dominance of Prochlorococcus' not actually quantified. See comment above.

We deleted this statement.

Line 202 - A brief description for how the *Nitzschia longissimi* was identified should be included (i.e. by sight, microscope identification?).

We described the microscopic analysis of diatoms in the section 2.6. of Materials and methods (L147-150).

Lines 312-315 (347) - See comment on phytoplankton proxies above.

See our answer above.

Lines 333-340 - See comment on DOM above.

See our answer above.

Line 347 - Based on uncertainty of phytoplankton composition estimates, I don't believe this phytoplankton uptake theory can be thrown out.

See our answer above.

Line 362 - Alternatively, DOP uptake and release is balanced.

We have revised based on your suggestion (L369-370).

Line 396-7 - This point is very interesting and I point the authors to this short compilation reference of nitrogen fixation estimates by Bonnet et al. 2017 in (<https://doi.org/10.1073/pnas.1619514114>). It is possible that iron (or a trace metal) is limiting, but the microbial population does not have standing stocks of nitrogen fixers.

We have referred Bonnet et al. (2017 PNAS) in the revised manuscript (L408).

Cross-basin differences in the nutrient assimilation characteristics of induced phytoplankton blooms in the subtropical Pacific waters

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15 **Abstract.** To better understand the nutrient assimilation characteristics of subtropical phytoplankton, deep water addition incubation experiments were carried out on surface waters collected at seven stations across the subtropical North and South Pacific Ocean. These deep water additions induced phytoplankton blooms with nutrient drawdown at all stations. The drawdown ratios of dissolved inorganic nitrogen (DIN) to phosphate (PO₄) varied from 14.1 to 30.7 at the PO₄-replete stations in the central North Pacific (CNP) and eastern South Pacific (ESP). These ratios were similar to the range
20 represented by the canonical Redfield ratio (16) through to typical particulate N:P ratios in the surface subtropical ocean (28). In contrast, lower DIN:PO₄ drawdown ratios (7.7-13.3) were observed in induced blooms at the PO₄-depleted stations in the western North Pacific (WNP). The DIN:PO₄ drawdown ratios in the PO₄-replete ESP were associated with eukaryote-dominated blooms, while those in PO₄-depleted WNP were associated with eukaryotic and cyanobacterial blooms. The surplus PO₄ assimilation, relative to DIN, by phytoplankton in the WNP was not expected based on their typical cellular N:P
25 ratio, and was likely due to the high PO₄ uptake capability as induced by low PO₄-adapted phytoplankton. The low and high P* (=PO₄-DIN/16) regimes geographically corresponded to the low and high DIN:PO₄ drawdown ratios in the WNP and the CNP or ESP, respectively. The basin-wide P* distribution in the oligotrophic Pacific surface waters showed a clear regional trend from low in the WNP (<50 nM) to high in the ESP (>100 nM). These results suggest that the subtropical phytoplankton blooms as observed in our experiments could be an important factor controlling P* as well as the commonly
30 recognized dinitrogen fixation and denitrification characteristics.

1 Introduction

The surface waters of the subtropical oceans are characterized by strong stratification, low nutrients, and low phytoplankton biomass (Karl, 2002). In this regime, primary production is largely sustained by regenerated production (f -ratio: ~ 0.1 ,
35 Dugdale and Goering, 1967; Eppley and Peterson, 1979), and driven by small phytoplankton such as the picocyanobacteria *Prochlorococcus* and *Synechococcus* (Waterbury et al., 1979; Chisholm et al., 1988). Despite the persistent oligotrophic regime, phytoplankton blooms with large diatoms and cyanobacteria occur occasionally in the subtropical oceans and have large impacts on new production and export production (Benitez-Nelson et al., 2007; McGillicuddy et al., 2007; Dore et al.,
40 2008; Wilson and Qiu, 2008; Karl et al., 2012; Villareal et al., 2012; Hashihama et al., 2014). The mechanisms that bring about the development of these blooms are not simple, but fundamentally they involve nutrient supply with physical forcing (Wilson et al., 2013; Toyoda and Okamoto, 2017).

The nutrient supply to surface subtropical oceans is important for many aspects of biogeochemical cycling and food-web dynamics as it drives new production and net community production (Sarmiento and Gruber, 2006; Saito, 2019). Seasonal variations in dissolved inorganic carbon and dissolved oxygen in the subtropical oceans highlight the net
45 productive systems, which are potentially sustained by intermittent nutrient supply from deep water (Michaels et al., 1994; Dore et al., 2003; Johnson et al., 2010). Deep water contains high amounts of nutrients such as nitrate (NO_3), phosphate (PO_4), and silicic acid ($\text{Si}(\text{OH})_4$), and their supply into the surface ocean temporarily alleviates phytoplankton nutrient stress. Several ship-based experimental studies indicate that deep water additions to subtropical surface waters have induced phytoplankton blooms (Mahaffey et al., 2012; Lampe et al., 2019; Robidart et al., 2019). These studies highlight the shifts in
50 phytoplankton community structure, growth characteristics, and gene expression during the bloom development. However, although nutrient assimilation characteristics are important mechanisms driving the net production, they were not fully described, for example the drawdown ratios (e.g., $\Delta\text{NO}_3:\Delta\text{PO}_4$ and $\Delta\text{Si}(\text{OH})_4:\Delta\text{NO}_3$), in these studies.

Phytoplankton N:P stoichiometry is generally based on the canonical ratio of 16 (Redfield, 1958). However, subtropical phytoplankton have higher N:P cellular ratios than Redfield and its mean value for subtropical waters is 28 (Martiny et al.,
55 2013). This higher ratio suggests that subtropical phytoplankton assimilate nutrients with higher N:P ratios than 16. If subtropical phytoplankton assimilate the upwelled deep water nutrients which have nearly Redfield $\text{NO}_3:\text{PO}_4$ ratios (~ 16 , Fanning, 1992), PO_4 -excess waters would remain at the surface. The PO_4 anomaly ($P^* = \text{PO}_4 - \text{NO}_3/16$) in the upper 120 m of the water column has indeed positive values throughout the subtropical oceans (Deutsch et al., 2007). As with N^* (Gruber and Sarmiento, 1997; Deutsch et al., 2001), P^* is recognized to be controlled by dinitrogen (N_2) fixation and denitrification
60 (Deutsch et al., 2007), but it may also be influenced by phytoplankton uptake of the upwelled deep water nutrients.

Subtropical phytoplankton utilize not only NO_3 and PO_4 but also nitrite (NO_2), ammonium (NH_4), dissolved organic N (DON), and dissolved organic P (DOP). Amongst them, the concentrations of DON and DOP are one to three orders of magnitude higher than those of dissolved inorganic N (DIN: the sum of NO_3 , NO_2 , and NH_4) and PO_4 in subtropical surface waters (Karl and Björkman, 2015; Sipler and Bronk, 2015). The majority of DON and DOP is likely refractory, but the

65 bioavailable forms such as urea, amino acid, and ATP play important roles in sustaining primary production in the inorganic nutrient-depleted subtropical waters (Kanda et al., 1985; Zubkov et al., 2004; Casey et al., 2009; Hill et al., 2011; Shilova et al., 2017; Björkman et al., 2018). Thus, the dynamics of alternative nutrients other than NO_3 and PO_4 should be considered when examining the nutrient assimilation characteristics of subtropical phytoplankton blooms.

Along with N and P assimilation, Si is also assimilated during diatom blooms. Several field studies reported anomalous
70 Si(OH)_4 removal relative to NO_3 and PO_4 at the sites of diatom blooms in the subtropical oceans (Benitez-Nelson et al., 2007; Hashihama et al., 2014). These Si(OH)_4 removals were not accompanied by stoichiometrically equivalent N and P removals as with a typical Si:N:P ratio of 16:16:1 (Redfield, 1958; Brzezinski, 1985). Given the linkages between Si and other elemental cycles, it is important to understand Si dynamics in the subtropical oceans. However, the Si dynamics cannot be fully explored from snapshot observations in the field (Hashihama et al., 2014), and to understand them experimental
75 validations are required.

In this study, our aim was to reveal N, P, and Si assimilation characteristics of subtropical phytoplankton blooms as induced by deep water additions. The onboard bottle incubation experiments were conducted across the subtropical Pacific Ocean, which has a large geographical variation in surface PO_4 from very low (<10 nM) in the western North Pacific (WNP) to high (>100 nM) in the eastern South Pacific (ESP) (Hashihama et al., 2019; Martiny et al., 2019; Hashihama et al., 2020).
80 Since subtropical phytoplankton respond to nanomolar increases in nutrient concentrations (Garside, 1985; Eppley and Renger, 1988; Eppley et al., 1990), we used sensitive liquid waveguide spectrophotometry for measuring nanomolar NO_3 , NO_2 , NH_4 , PO_4 , and Si(OH)_4 . The nanomolar nutrient data enabled us to calculate accurate stoichiometric ratios for N, P, and Si. Along with the inorganic nutrients, we also examined DON and DOP variations during the incubation experiments. These deep-water addition experiments successfully induced phytoplankton blooms, while the nutrient drawdown ratios
85 showed geographical patterns concomitant with the surface PO_4 distributions. Here we conclude by discussing the mechanism of the regionally different drawdown ratios and its possible influences on P^* distribution in the subtropical Pacific Ocean.

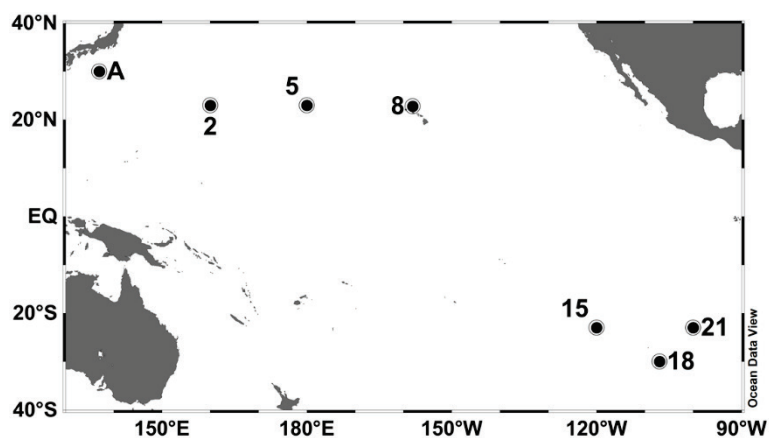
2 Materials and methods

2.1 Study areas and water sampling

90 Observations were conducted at seven stations in the subtropical North and South Pacific Ocean (Table 1 and Fig. 1). Station A in the WNP was occupied in July 2010 during the R/V Tansei Maru KT-10-13 cruise. Stations 2-21 were occupied along the transect from the WNP (2 and 5) to the ESP (15, 18, and 21) through the central North Pacific (CNP, 8) for the period from December 2011 to January 2012 during the R/V Hakuho Maru KH-11-10 cruise. Water sampling was performed using a conductivity-temperature-depth (CTD) system (Sea-Bird Electronics) equipped with HCl-cleaned Teflon-coated Niskin-X
95 bottles (General Oceanics). Water samples for incubation experiments were collected from 10 m depth (hereafter referred to as 'surface') at all stations. At Stations A and 2 in the WNP, deep water from 1500 m depth was also collected.

Table 1. Details of the incubation experiments.

Region	Sampling station	Latitude	Longitude	Sampling date (GMT)	Sampling depth (m)	Incubation period (h)	Mean PAR during incubation ($\mu\text{mol photons m}^{-2} \text{s}^{-1}$)
WNP	A	30.00°N	137.01°E	2010/7/11	10	52	582
WNP	2	23.00°N	160.00°E	2011/12/6	10	96	271
WNP	5	23.00°N	180.00°E/W	2011/12/12	10	96	284
CNP	8	22.77°N	158.09°W	2011/12/18	10	48	240
ESP	15	23.00°S	120.00°W	2012/1/7	10	96	542
ESP	18	30.00°S	107.00°W	2012/1/13	10	96	627
ESP	21	23.00°S	100.00°W	2012/1/17	10	96	549



100 **Figure 1.** Study areas and sampling stations in the subtropical Pacific Ocean. Station A was occupied in July 2010 during the R/V Tansei Maru KT-10-13 cruise. Stations 2-21 were occupied for the period from December 2011 to January 2012 during the R/V Hakuho Maru KH-11-10 cruise.

2.2 Deep-water addition incubation experiments

105 Surface waters collected from 10 m depth were poured into 2.3 L HCl-cleaned polycarbonate bottles and then 50 mL of deep water was immediately added to the triplicate bottles (2.1% v/v deep water addition). **Filtration of the deep water was not conducted to avoid NH_4 contamination.** The deep waters collected at Stations A and 2 were added to the surface water samples at Stations A and 2-21, respectively. The deep waters for the experiments at Stations 5-21 were kept frozen (-20°C) until used at each station. Nutrient concentrations in the deep waters at Stations A and 2 were 37.1 and $39.0 \mu\text{M DIN}$, 2.1 and $2.9 \mu\text{M PO}_4$, and 134.5 and $140.5 \mu\text{M Si(OH)}_4$, respectively. For all experiments, triplicate control bottles were prepared.

110 Both the treated and control bottles were incubated for 48-96 h (Table 1) in an on-deck incubator with flowing surface seawater, which was shaded with appropriate sheeting to give 30% of full sunlight so as to mimic the ambient photosynthetically active radiation (PAR) condition at 10 m depth. The ambient PAR on deck was continuously monitored

by a LI-COR quantum sensor (LI-190R) with a data logger (LI-1400), and its mean values during the incubation periods including day and night times are presented in Table 1. During the incubation periods (48-96 h), the bottles were subsampled 5-6 times for nanomolar nutrients, 3 times for DON and DOP, and 1 time (at the end point) for phytoplankton. Initial phytoplankton samples at time-zero were collected in single or duplicate directly from the Niskin-X bottles. At the end point, water volumes in the bottles just prior to final sampling were approximately 1.8 L. The DON and DOP samples were not collected at Station A during the KT-10-13 cruise. To assess any significant decrease or increase in the concentrations of nanomolar nutrients, DON, and DOP during the incubation periods, linear regression analyses were performed. In addition, Student *t*-test and paired *t*-test were performed to determine significant differences between the measured parameter values. In this paper, the significance is reported where $p < 0.05$. In the calculation of nutrient drawdown ratio, its error was estimated based on the error propagation rule (Miller and Miller, 1993).

2.3 Determinations of nanomolar nutrients

Water samples for nanomolar nutrients were collected in 30 mL HCL-cleaned polypropylene tubes and were frozen at -20 °C until analysis. The concentrations of NO₃, NO₂, NH₄, PO₄, and Si(OH)₄ were determined using an automated liquid waveguide spectrophotometric system equipped with 50-100 cm liquid waveguide capillary cells (LWCC, World Precision Instruments) (Hashihama et al., 2009; Hashihama and Kanda, 2010; Hashihama et al., 2014; Hashihama et al., 2015). The detection limits for NO₃, NO₂, NH₄, PO₄, and Si(OH)₄ were 3, 2, 6, 3, and 11 nM, respectively. Although we collected triplicate water samples from the incubated bottles, triplicate analytical data were not available for several samples.

2.4 Determinations of DON and DOP

Water samples for DON and DOP were collected in HCL-cleaned polypropylene tubes after removing particulate matter by filtering through pre-combusted Whatman GF/F filters. The samples were frozen at -20 °C until analysis. Total dissolved N (TDN) and P (TDP) were quantified by a persulfate oxidation method (Hansen and Koroleff, 1999) with a QuAAtro TN-TP analyser (SEAL Analytical) (Yasui et al. 2016; Yasui-Tamura et al., 2020). Concentration of DON was derived from the difference between TDN and DIN concentrations, and that of DOP was derived from the difference between TDP and PO₄ concentrations. As with the nutrients (2.3), triplicate analytical data on DON and DOP were not obtained for some samples.

2.5 Phytoplankton pigment analysis

Phytoplankton pigment analysis was performed using high-performance liquid chromatography (HPLC). Water volumes of 440-3000 mL were filtered onto GF/F filters, and the filter samples were immediately frozen in liquid nitrogen and stored in a deep freezer (-80 °C) until analysis. Pigment analysis was conducted using the method of Zapata et al. (2000) with a HPLC system (Hashihama et al., 2008). Six phytoplankton pigments - chlorophyll *a* (Chl *a*), divinyl chlorophyll *a* (DVchl *a*), 19'-butanoyloxyfucoxanthin (But-fuco), fucoxanthin (Fuco), 19'-hexanoyloxyfucoxanthin (Hex-fuco), and zeaxanthin (Zea) - were quantified from the peak area calibrated against that of standard pigments (DHI Water and Environment). Total

chlorophyll *a* (Tchl *a*: the sum of Chl *a* and DVchl *a*) was used as an index of total phytoplankton biomass. Total
145 fucoxanthin (Tfuco: the sum of But-fuco, Fuco, and Hex-fuco) was used as a representative marker of eukaryotic
phytoplankton. *Zea* and DVchl *a* were markers of cyanobacteria and *Prochlorococcus*, respectively.

2.6 Microscopic analysis of phytoplankton

Water volumes of 100-1000 mL seawater samples were fixed with neutralized formalin at a final concentration of 1% (v/v).
The fixed samples were concentrated through sedimentation in a land-based laboratory. Diatom **species** were identified and
150 enumerated under an inverted microscope (Utermöhl, 1958).

2.7 P* determination using nanomolar nutrient data

Surface P* at the experimental stations was calculated using the measured nanomolar PO₄ and DIN data through an equation
P*=PO₄-DIN/16. In addition, to reveal basin-wide distribution of surface (≤10 m) P* over the oligotrophic Pacific area (40°
N-40° S), we assembled nanomolar (<1000 nM) data sets of PO₄ and NO₃ plus NO₂ (N+N), most of which were previously
155 published by the authors in this study (Hashihama et al., 2009; Kitajima et al., 2009; Shiozaki et al., 2009; Hashihama et al.,
2010; Sato et al., 2010; Shiozaki et al., 2010; Girault et al., 2013; Sato et al., 2013; Hashihama et al., 2014; Shiozaki et al.,
2014; Girault et al., 2015; Sato et al., 2015; Sato et al., 2016; Shiozaki et al., 2016; Shiozaki et al., 2017; Ellwood et al.,
2018; Horii et al., 2018; Shiozaki et al., 2018; Hashihama et al., 2019; Martiny et al., 2019; Sato and Hashihama, 2019;
Yamaguchi et al., 2019; Hashihama et al. 2020; Yamaguchi et al. in press; Jiang et al. in press). We also included several
160 unpublished data sets collected by F. Hashihama and T. Kodama. These data sets were obtained by using the liquid
waveguide spectrophotometry for PO₄ and N+N (Woodward, 2002; Hashihama et al., 2009). Since surface NH₄
concentrations were typically low at the sub-nanomolar level and the NH₄ data were relatively limited compared to the PO₄
and N+N data, we did not use the NH₄ data to show the basin-wide distribution of P*. Thus, the P* in this case was
calculated through an equation P*=PO₄-N+N/16.

165 3 Results

3.1 Initial conditions

High temperature (21.76-26.91 °C) and high salinity (34.07-36.50) in the surface waters (10 m) of the seven experimental
stations indicated that typical subtropical oceanic waters prevailed in the study regions (Table 2). DIN concentrations at the
surface were consistently lower than 50 nM, while PO₄ concentrations varied geographically and were extremely low in the
170 WNP (<10 nM; Stations A, 2, and 5), intermediate in the CNP (53 nM; Station 8), and high in the ESP (>100 nM; Stations
15, 18, and 21). Surface P* at these stations showed a geographical variation similar to PO₄ concentrations, and this trend
was due to **the consistently low concentrations of DIN** found at those stations. Si(OH)₄ concentrations were higher in the

WNP and CNP (767-1276 nM; Stations A, 2, 5, and 8) than the ESP (427-541 nM; Stations 15, 18, and 21), and DON and DOP concentrations ranged from 3.47 to 4.45 μM and 0.10 to 0.21 μM , respectively.

175 Tchl *a* concentrations at the surface were less than 129 ng L^{-1} with low values at Stations 18 and 21 in the ESP (18 and 3 ng L^{-1} , Table 2). Tfuco, Zea, and DVchl *a* all showed geographical variations similar to Tchl *a*. Tfuco:Zea ratios were lower than 1.0 except for Station 18 in the ESP (2.6), where the biomass proportion of eukaryotes to cyanobacteria was relatively high. DVchl *a*:Tchl *a* ratios (indices of the contribution of *Prochlorococcus* to total phytoplankton) were mostly 0.4-0.5, but the lower ratios were observed at Stations 18 and 21 in the ESP (0.1 and 0.2). Cell densities of surface diatoms
180 were consistently low (3-88 cells L^{-1}) at all stations.

Table 2. Initial conditions for incubation samples collected from 10 m depth. Errors represent standard deviations ($n=3$). No error means no triplicate data set. nd: no data.

Region	Station	Temperature (°C)	Salinity	DIN (nM)	PO ₄ (nM)	P* (nM)	Si(OH) ₄ (nM)	DON (μM)	DOP (μM)	Tchl <i>a</i> (ng L ⁻¹)	Tfuco (ng L ⁻¹)	Zea (ng L ⁻¹)	DVchl <i>a</i> (ng L ⁻¹)	Tfuco :Zea (g:g)	DVchl <i>a</i> :Tchl <i>a</i> (g:g)	Diatoms (cells L ⁻¹)
WNP	A	26.91	34.07	9±2	7±2	6±2	1276±15	nd	nd	45	17	32	18	0.5	0.4	88
WNP	2	26.72	34.92	16±14	2±1	1±1	870±25	3.47±0.16	0.17±0.03	26	13	27	13	0.5	0.5	20
WNP	5	26.37	35.31	11±2	8±1	7±1	767±36	3.65±0.24	0.19±0.05	52	21	32	24	0.6	0.5	18
CNP	8	24.24	35.29	22	53	52	989	4.25	0.21±0.05	129	44	47	65	0.9	0.5	4
ESP	15	25.28	36.50	16±12	228±8	227±8	541±19	4.45±0.37	0.10	49	17	46	25	0.4	0.5	20
ESP	18	21.76	35.57	47±24	124±11	121±11	427±24	3.51±0.05	0.15±0.05	18	11	4	3	2.6	0.1	3
ESP	21	24.09	35.87	40	272	270	439±53	3.62	nd	3	2	5	1	0.4	0.2	19

3.2 Phytoplankton response to deep water additions

185 Mean Tchl *a* concentrations were three to ten times higher in the treated bottles than observed for the control bottles in all experiments, although no significant difference was observed at Station A due to a highly variable results in the treated bottles (Fig. 2a). These trends in increasing Tchl *a* indicate that the deep water additions positively induced phytoplankton blooms. The Tchl *a* differences between the treated and control bottles were greatest at Station A in the WNP (178 ng L⁻¹) and Station 15 in the ESP (306 ng L⁻¹), and net growth rates of the blooms (as Tchl *a*, relative to the control) were higher at
190 these two stations (0.70 and 0.58 d⁻¹) than other stations (0.27-0.50 d⁻¹). Mean Tfuco concentrations were significantly higher in the treated than in the control bottles except for Station A (Fig. 2b). Mean Zea concentrations were significantly higher in the treated bottles in the WNP (Stations A, 2, and 5) and at two sites of the ESP (Stations 15 and 21) (Fig. 2c). Mean DVchl *a* concentrations were significantly higher in the treated bottles at two sites in the WNP (Stations 2 and 5) and Station 21 in the ESP (Fig. 2d). The pigment concentrations in the control bottles were similar to those in the initial conditions, indicating
195 that changes in pigment concentrations due to photoacclimation during the incubation periods are small.

Mean Tfuco:Zea ratios were higher in the treated than in the control bottles, although no significant differences were observed at Stations A and 18 (Fig. 2e). The higher Tfuco:Zea ratios in the treated bottles imply that biomass increases in eukaryotes were relatively large compared to those of cyanobacteria. The ratios of Tfuco:Zea in the treated bottles were higher in the ESP (Stations 15, 18, and 21) than the WNP and CNP (Stations A, 2, 5, and 8), indicating that the proportions
200 of eukaryotes (cyanobacteria) were higher (lower) in the ESP than the WNP and CNP. Mean DVchl *a*:Tchl *a* ratios were significantly lower in the treated bottles than the control, except for Stations 18 and 21 in the ESP, where the ratios in the control were quite low (<0.1) as observed in the initial conditions (Fig. 2f).

Cell densities of diatoms were significantly higher in the treated bottles at two sites of the WNP (Stations A and 2) and Station 18 in the ESP (Fig. 2g). An exceptionally high mean density of diatoms (907 cells L⁻¹), mostly consisting of
205 *Nitzschia longissima*, was observed in the treated bottles at Station 15 in the ESP, although no significant difference between the densities in the control and treated bottles was seen.

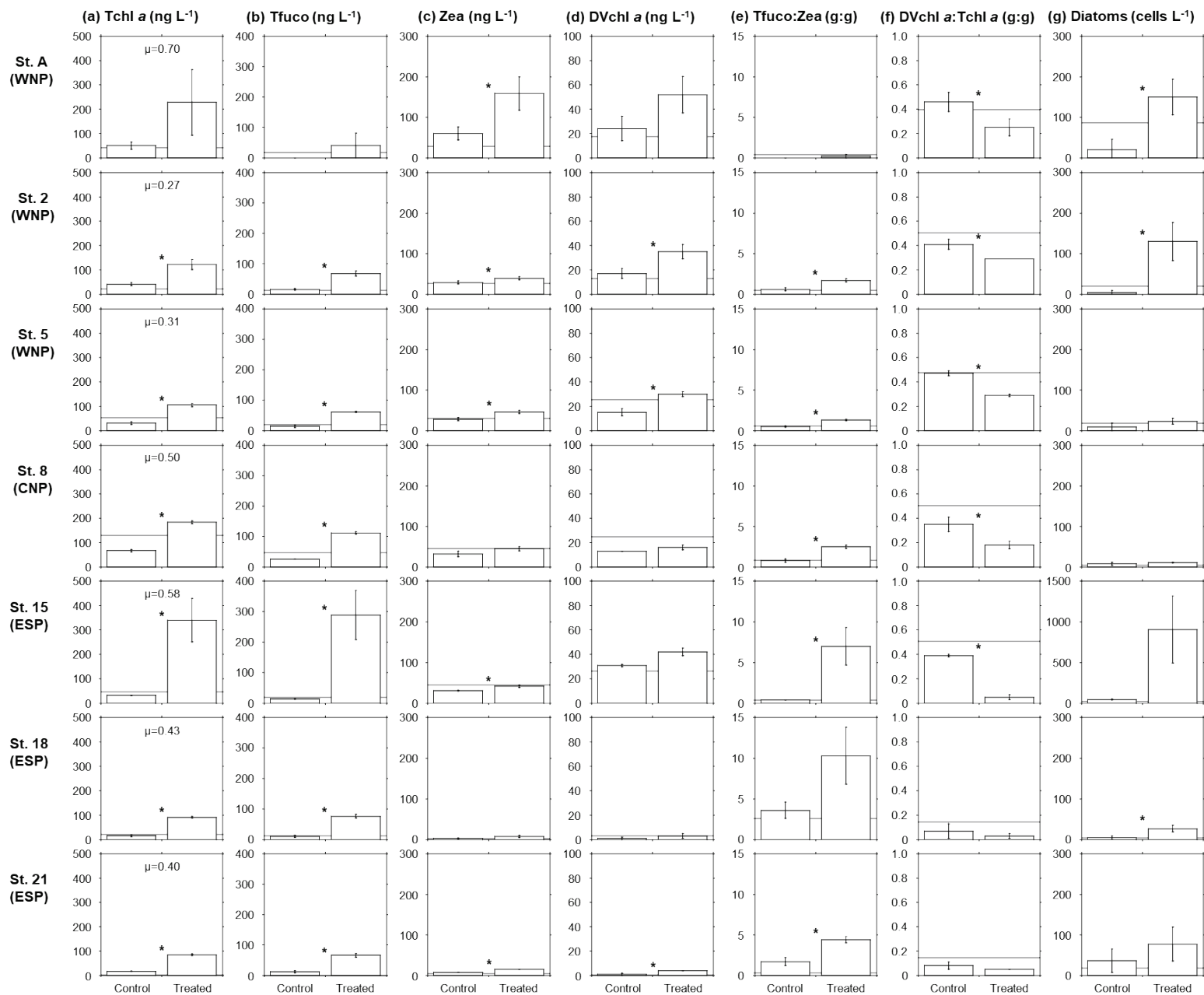


Figure 2. (a) Tchl *a*, (b) Tfuco, (c) Zea, (d) DVchl *a*, (e) Tfuco:Zea ratios, (f) DVchl *a*:Tchl *a* ratios, and (g) diatom cell densities in the control and treated bottles after 48-96 h incubations at seven stations. A grey horizontal line in each panel indicates a mean or single concentration of each pigment at the initial point. Error bars denote standard deviations ($n=3$). Significant differences (t -test, $p<0.05$) between the values in the control and treated bottles are depicted by asterisks. Net growth rates, μ (d^{-1}) of the phytoplankton blooms (as Tchl *a*, relative to the control) are denoted in (a) Tchl *a*. In (g) diatom cell densities, a scale of y-axis at Station 15 (~ 1500 cells L^{-1}) is different from those at other stations (~ 300 cells L^{-1}).

3.3 Nutrient drawdown

Following the phytoplankton blooms, DIN and PO_4 concentrations in the treated bottles at all stations showed significant linear decreases as a function of the incubation times ($r^2>0.82$, Figs. 3a and 3b). In contrast, the concentrations in the control bottles at all stations showed no significant trends. The DIN decreases were largely ascribed to NO_3 decreases (Fig. A1a), and interestingly, NO_2 concentrations in the treated bottles at all stations significantly increased with time (Fig. A1b). There were also significant linear increases in the control bottles for NO_3 at Stations 8 and 15, and NO_2 at Station 18 but these changes were quite small (<13 nM). NH_4 concentrations in the treated and control bottles showed no significant trends with time, but occasionally high standard deviations (>100 nM) were observed (Fig. A1c).

Drawdown rates (slopes of linear regression lines in Figs. 3a and 3b) of DIN and PO_4 in the treated bottles varied from 1.38 to **7.37** $nmol\ N\ L^{-1}\ h^{-1}$ and from 0.13 to 0.71 $nmol\ P\ L^{-1}\ h^{-1}$, respectively. **These rates were not significantly different from the rates derived from the first 36-48 h incubation data (paired t -test, $p>0.05$).** The **drawdown** rates were relatively low (≤ 1.49 $nmol\ N\ L^{-1}\ h^{-1}$ and ≤ 0.15 $nmol\ P\ L^{-1}\ h^{-1}$, respectively) in the WNP during winter (December, Stations 2 and 5) where relatively low mean PAR was observed during the incubation periods (≤ 284 $\mu mol\ photons\ m^{-2}\ s^{-1}$, Table 1). Differences between the control-corrected mean concentrations of DIN and PO_4 at the start and end points of the incubation (Δ DIN and Δ PO_4) varied from 123 to **749** nM and from 9 to 53 nM, respectively (Table 3). The values of Δ DIN and Δ PO_4 normalized by the incubation times (h) were almost identical to the **drawdown** rates of DIN and PO_4 in the treated bottles ($r^2=0.99$ and $r^2=0.91$ **in linear regressions**, respectively).

Unlike the DIN and PO_4 concentrations, $Si(OH)_4$ concentrations in the treated bottles at all stations did not show any significant linear decreases with the occasionally high standard deviations (>500 nM) (Fig. 3c). The insignificant trends in $Si(OH)_4$ concentrations were also observed in the control bottles at all stations. Difference between the control-corrected mean $Si(OH)_4$ concentrations at the start and end points (Δ $Si(OH)_4$) showed net drawdown values ranging from 7 to **465** nM, although mean $Si(OH)_4$ concentrations in the treated bottles at the start and end points were not significantly different except at Station A (Table 3).

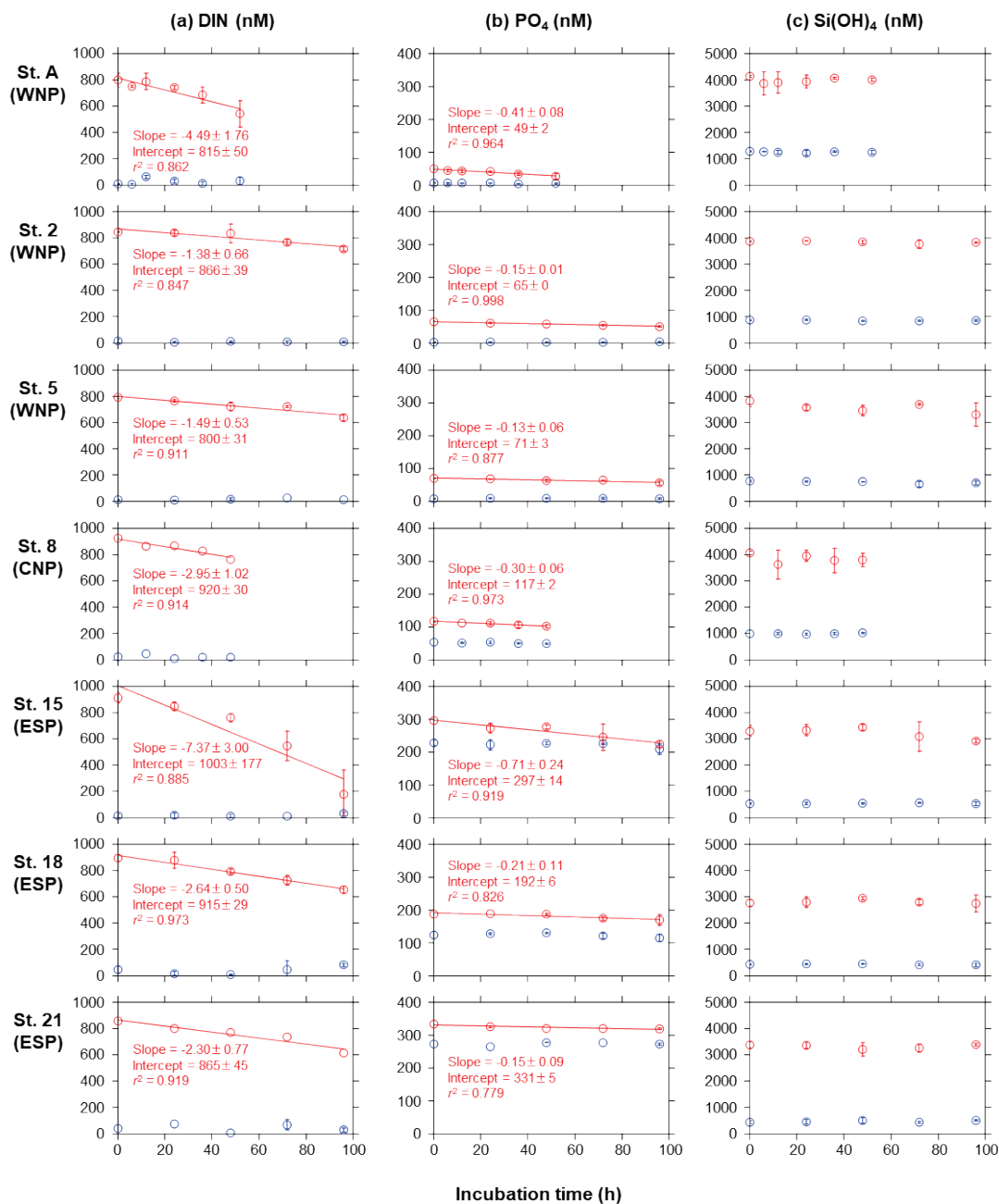


Figure 3. Temporal changes in concentrations of (a) DIN, (b) PO₄, and (c) Si(OH)₄ in the control (blue) and treated (red) bottles during the incubation periods at seven stations. Error bars denote standard deviations ($n=3$). Duplicate or single data are denoted as mean or single values without error bars. Linear regression lines are depicted when significant decreases ($p < 0.05$) in the mean concentrations against time were observed. **Errors of slope and intercept represent 95% confidence intervals.**

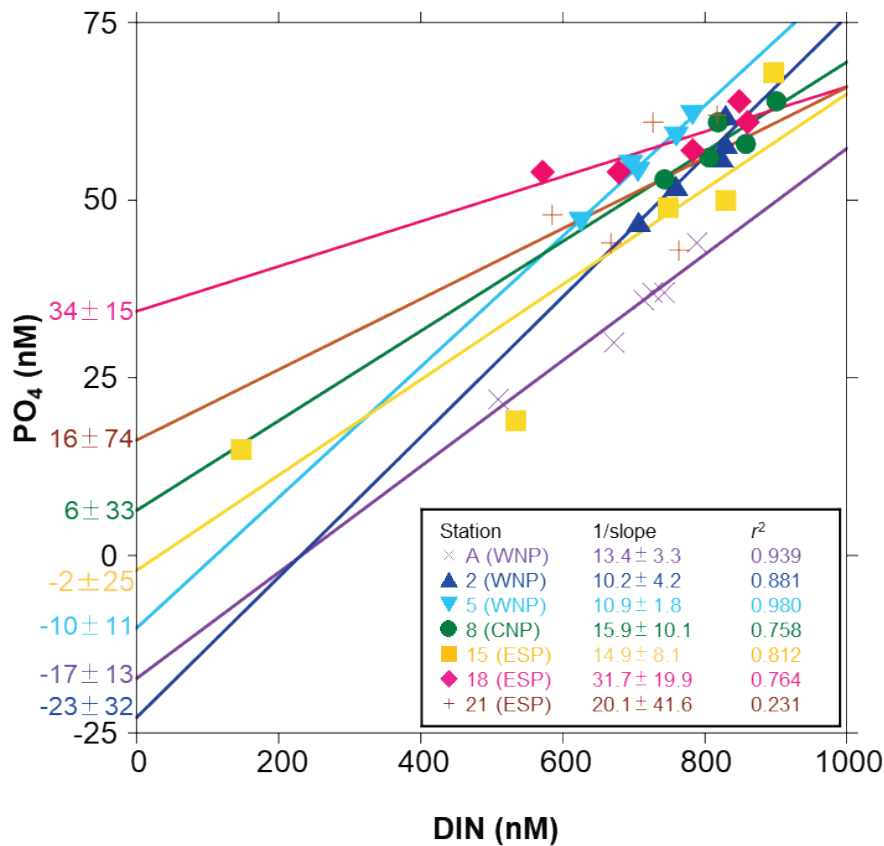
Table 3. Nutrient drawdowns and their ratios throughout the incubation periods. **Errors represent 95% confidence intervals.**
 245 **No error means no triplicate data set.** ^a Difference between the control-corrected mean concentrations at start and end points of incubations. ^b No significant difference between the mean Si(OH)₄ concentrations in the treated bottles at start and end points of incubations (*t*-test, *p*>0.05).

Region	Station	Incubation period (h)	ΔDIN ^a (nM)	ΔPO ₄ ^a (nM)	ΔSi(OH) ₄ ^a (nM)	ΔDIN:ΔPO ₄ (mol:mol)	ΔSi(OH) ₄ :ΔDIN (mol:mol)
WNP	A	52	279±131	21±10	101±111	13.3±82.8	0.36±0.22
WNP	2	96	123±26	16±4	21±55 ^b	7.7±12.6	0.17±0.08
WNP	5	96	157±32	14±7	465±561 ^b	11.2±26.0	2.96±10.59
CNP	8	48	158	11±7	304 ^b	14.4	1.92
ESP	15	96	749±215	53±24	374±270 ^b	14.1±57.7	0.50±0.23
ESP	18	96	276±52	9	7±392 ^b	30.7	0.03±0.04
ESP	21	96	232	15	41±163 ^b	15.5	0.18

3.4 Nutrient drawdown ratio

In all experiments, the enriched nutrients in the treated bottles were not fully taken up by phytoplankton during the
 250 incubation periods (Fig. 3). Therefore, we assessed the nutrient drawdown ratios using ΔDIN, ΔPO₄, and ΔSi(OH)₄ (Table 3). ΔDIN:ΔPO₄ ratios varied from 7.7 to 30.7. **Although 95% confidence intervals of these ratios were large (12.6-82.8), the relatively lower ratios (≤13.3) were convergent in the WNP (Stations A, 2, and 5).** ΔSi(OH)₄:ΔDIN ratios varied from 0.03 to 2.96 with most stations less than 1 except for Stations 5 and 8. However, these ratios, except at Station A (0.36), involved uncertainties due to no significant decreases in Si(OH)₄ concentrations in the treated bottles during the incubations.

255 We also evaluated the drawdown characteristics using the control-corrected mean concentrations of DIN and PO₄ at sampling points during the incubation periods. A plot of PO₄ against DIN showed strong negative linear relationships (*r*²>0.75), except for Station 21 (*r*²=0.23) (Fig. 4). Here, the drawdown ratio of DIN to PO₄ was expressed as 1/slope of the linear regression, and ranged from 10.2 to 31.7 **with 95% confidence intervals between 1.8 and 41.6.** These drawdown ratios were almost identical to the ΔDIN:ΔPO₄ ratios in Table 3 (*r*²=0.95 in linear regression). In addition, we observed unique
 260 variations in PO₄-intercepts of the linear regression lines. **The PO₄-intercepts varied from -23 to 34 nM with 95% confidence intervals between 11 and 74 nM, and the relatively lower values (<10 nM) being convergent in the WNP (Stations A, 2, and 5).**



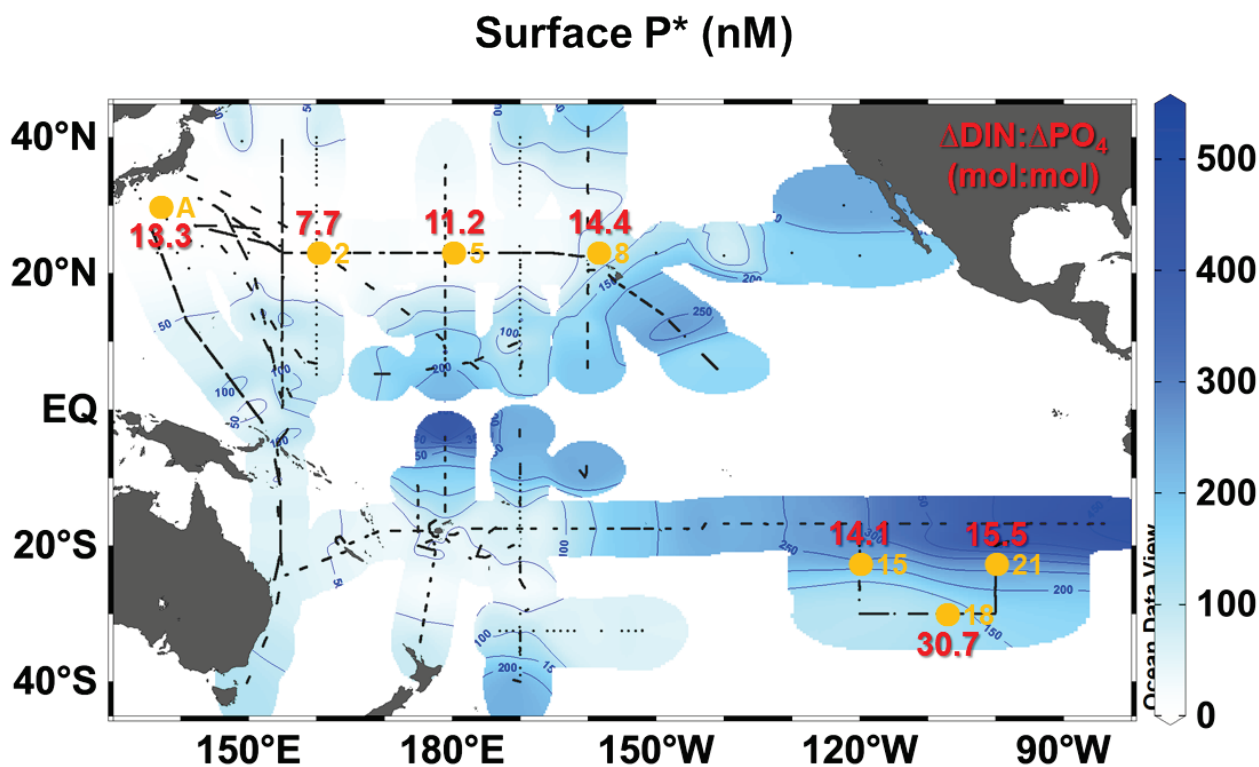
265 **Figure 4.** Scatter plots of the control-corrected mean concentrations of PO_4 against DIN in the incubation experiments at seven stations. Linear regression lines with their parameters (1/slope, PO_4 -intercept, and r^2) at seven stations were denoted by the different colours. Errors of 1/slope and PO_4 -intercept represent 95% confidence intervals.

3.5 DON and DOP

270 DON and DOP concentrations in both the treated and control bottles did not show any significant increase or decrease as a function of the incubation time, except for DOP in the treated bottles at Station 8 in the CNP (Figs. A2a and A2b). At this station, the DOP concentrations in the treated bottles significantly increased with time, although the change over 48 h ($0.03 \mu\text{M}$) was smaller than that from 0 to 24 h ($-0.07 \mu\text{M}$) in the control bottles. Overall, the DON and DOP concentrations in the treated bottles were similar to those in the control bottles, indicating that the deep water additions did not alter DON and DOP regimes during the incubation periods.

275 **3.6 Basin-wide P* distribution in the oligotrophic Pacific**

The assembled surface N+N and PO₄ data (Figs. A3a and A3b) revealed a detailed surface P* distribution over the oligotrophic Pacific Ocean (Fig. 5). The distributional pattern of P* was similar to that of PO₄ (Fig. A3b), mainly due to a broad area with low N+N (<100 nM) (Fig. A3a). The P* showed a clear west-east gradient from <50 nM in the western basin to ~500 nM in the eastern basin. In the western basin, the extremely low P* (<10 nM) was found in the WNP. Stations A, 2, and 5 were located within this low P* area. Station 8 was in the intermediate P* area (50-100 nM) in the CNP, while Stations 15, 18, and 21 were within the high P* area (>100 nM) in the ESP.



285 **Figure 5.** Surface distributions of P* in the oligotrophic Pacific Ocean (40° N-40° S). The P* data were derived from the assembled data on nanomolar concentrations (<1000 nM) of PO₄ and N+N. Small black dots denote sampling stations for the nanomolar PO₄ and N+N. Large orange circles denote the stations where the incubation experiments were conducted. Red values indicate the experimentally-determined $\Delta\text{DIN}:\Delta\text{PO}_4$ ratios (Table 3).

4 Discussion

4.1 Phytoplankton blooms following deep water additions

290 Our study confirms that deep water additions induced phytoplankton blooms in various regions of the subtropical Pacific Ocean. Such induced blooms have also been reported at Station ALOHA in the subtropical North Pacific Ocean (Mahaffey et al., 2012). The ALOHA experiments revealed that Tchl *a*-based growth rates following deep water additions were higher in boreal summer than boreal winter. A similar seasonal trend was observed in the North Pacific stations reported here; net growth rates were higher at the westernmost site in July (Station A, 0.70 d⁻¹) than other sites in December (Stations 2, 5, and 295 8, 0.27-0.50 d⁻¹) (Fig. 2a). The low growth rates in winter could be explained by low assimilation rates of nutrients (Fig. 3). These results suggest that the magnitude of a subtropical phytoplankton bloom is regulated by additional seasonal factors such as PAR (Table 1).

The net phytoplankton growth rates were relatively low at Stations 18 and 21 in the ESP (0.43 and 0.40 d⁻¹, respectively, Fig. 2a), even for austral summer conditions with high PAR (Table 1). Since the ESP is known as a low dust deposition area, growth by the resident phytoplankton is considered to be limited by iron (Fe) (Jickells et al., 2005; Blain et al., 2008; 300 Wagener et al., 2008; Moore et al., 2013). However, a Fe-enrichment incubation experiment at Station 18 demonstrated that there was no significant difference between the Fe-enriched and control bottles for phytoplankton, nutrients, DON, and DOP during a 96 h incubation, and also these parameter values were little changed from their initial values as the initial values lay within standard deviations of the mean values in the Fe-enriched and control bottles (Appendix methods, Table A1). Similar 305 experimental results were reported by Bonnet et al. (2008) and they suggested that phytoplankton growth in the ESP is limited by N rather than Fe. Furthermore, the deep waters used in this study were from a depth (1500 m) within North Pacific intermediate water that typically contains high dissolved Fe (>0.6 nM, i.e., not iron-limiting) (Nishioka et al., 2013; Nishioka et al., 2020). These lines of evidence imply that surface phytoplankton in the ESP and their bloom formation were not primarily limited by Fe. Although grazing by zooplankton is a possible factor controlling phytoplankton net growth, it 310 was suggested to be not strong in the case of the subtropical phytoplankton blooms following deep water additions (Mahaffey et al., 2012). There is further research required to understand the factors controlling the bloom development in the ESP.

The induced phytoplankton blooms in this study were accompanied by changes in community structure. Several nutrient enrichment experiments conducted in the subtropical oceans have demonstrated significant increases in eukaryotic 315 phytoplankton following nutrient enrichments, particularly of N (Bonnet et al., 2008; Moore et al., 2008; Mahaffey et al., 2012; Shilova et al., 2017; Rii et al., 2018; Lampe et al., 2019; Robidart et al., 2019). Similar blooms dominated by eukaryotic phytoplankton were also observed at most stations in this study as evidenced by the Tfuco increases (Fig. 2b). In addition, significant increases in cyanobacteria (Zea and DVchl *a*) following deep water additions were also observed particularly in the WNP and CNP (Figs. 2c and d). The relative proportions of cyanobacteria to eukaryotes in the treated 320 bottles were lower in the ESP than the WNP and CNP (Fig. 2e). At Stations 18 and 21 in the ESP, the low proportions of

Prochlorococcus at time zero (Table 2) might influence the low proportions of cyanobacteria in the induced phytoplankton blooms. On the other hand, the relative increases of cyanobacterial abundances in the North Pacific experiments are likely driven by seasonal phytoplankton response to nutrient enrichment. Mahaffey et al. (2012) reported that there were less increases in eukaryote abundances in winter than in summer at Station ALOHA following deep water additions. Because of this, the relative increases in cyanobacteria in the winter-time North Pacific (Stations 2, 5, and 8) might be significant. However, although Station A in the WNP was occupied in summer, we did not observe any significant eukaryotic bloom following deep water addition (Fig. 2e). This opposing trend at Station A could be due to regional differences of seasonal phytoplankton responses to nutrient enrichments, as summer eukaryotic blooms frequently occur in the eastern basin compared to the western basin in the North Pacific (Wilson, 2011; Villareal et al., 2012; Hashihama et al., 2014; Jiang et al. in press).

4.2 N and P drawdown characteristics

Our incubation experiments have revealed consistent linear decreases in DIN and PO₄ concentrations, at nanomolar levels, along with the development of phytoplankton blooms (Figs. 3a and 3b). Additionally, accurate measurements of nanomolar inorganic N species detected the consistent increases in NO₂ concentrations in the treated bottles in which there were large decreases in NO₃ concentrations. This trend was particularly prominent at Station A (Figs. A1a and A1b). These NO₂ increases could be due to in vitro nitrification of NH₄ and/or incomplete assimilation of NO₃ by phytoplankton (Lomas and Lipschultz, 2006). While the factors controlling regional or seasonal differences in these NO₂ increases remains unknown, we demonstrate that sensitive measurements for multiple nutrients enable us to detect trace, but important, biogeochemical dynamics in response to the addition of nutrient-rich deep water.

Unlike the DIN and PO₄, DON and DOP concentrations did not show any consistent changes over time despite the occurrence of the phytoplankton blooms (Fig. A2). Although natural phytoplankton in the nutrient-depleted oligotrophic oceans show high affinity to DON and DOP (Karl and Björkman, 2015; Sipler and Bronk, 2015), the DON and DOP concentrations here did not indicate net drawdown in either the treated or control bottles. For the treated bottles, phytoplankton N and P demands were largely met by the enriched inorganic nutrients - which were not exhausted during the incubation periods (Figs. 3a and 3b). The DON and DOP dynamics during the development of phytoplankton blooms appear to be in an equilibrium between uptake and release, as seen under ambient conditions like those in the control bottles (Fig. A2).

Although the induced phytoplankton blooms in this study were solely dependent on DIN and PO₄, regionally unique Δ DIN: Δ PO₄ ratios were unveiled (Table 3). The Δ DIN: Δ PO₄ ratios showed a geographical trend with relatively lower ratios in the PO₄-depleted WNP (7.7-13.3) than in the other PO₄-replete regions (14.1-30.7) although there were large variation in 95% confidence intervals (12.6-82.8). While the ratios in the PO₄-replete regions were similar to the range (16-28) from the Redfield ratio to the subtropical particulate N:P ratios (Redfield, 1958; Martiny et al., 2013), those in the WNP were convergently lower than 16. Cellular N:P ratios in the subtropical phytoplankton are higher in cyanobacteria (25-35) than for

eukaryotes (16) (Martiny et al., 2013). Because the phytoplankton blooms in the WNP were composed of communities with
355 a relatively high proportion of cyanobacteria (Figs. 2e and 2f), the lower $\Delta\text{DIN}:\Delta\text{PO}_4$ ratios (7.7-13.3) could not be explained
by phytoplankton cellular N:P ratios. Since macro-scale (>2000 km) exhaustion of PO_4 in the WNP is coupled with N_2
fixation (Hashihama et al., 2009; Martiny et al., 2019), N_2 fixation by diazotrophic cyanobacteria might potentially meet
some of phytoplankton N demand. However, assuming a relatively high N_2 fixation rate in the WNP ($5 \text{ nmol N L}^{-1} \text{ d}^{-1}$,
Hashihama et al., 2020), the contributions of N_2 fixation to $\Delta(\text{DIN}+\text{N}_2)$ were small (4-16%), leading to still lower
360 $\Delta(\text{DIN}+\text{N}_2):\Delta\text{PO}_4$ ratios (8.9-13.8) than 16.

Other explanations for the regionally unique $\Delta\text{DIN}:\Delta\text{PO}_4$ ratios we observed (Table 3) are based on resident
communities being 'primed' for rapid P acquisition. Lomas et al. (2014) reported that phytoplankton in the western
subtropical North Atlantic have active PO_4 transporters which are rapidly induced under severely PO_4 -depleted condition. In
addition, several studies have reported that microbial genes for high-affinity PO_4 transporter (*pstSCAB*) are enriched in the
365 PO_4 -depleted regions in the western North Atlantic and WNP compared to the PO_4 -replete regions in the CNP and ESP
(Coleman and Chisholm, 2010; Hashihama et al., 2019). These studies indicated that the phytoplankton in the PO_4 -depleted
regions possess a high PO_4 uptake capability. Since N perturbation to phytoplankton little alters their cellular N quota
(Moreno and Martiny, 2018), a high cellular P accumulation by the high uptake capability could induce the surplus
drawdowns of PO_4 relative to DIN in the incubation experiments of the PO_4 -depleted WNP. Since **DOP uptake and release**
370 **in the treated bottles in the WNP were balanced** (Fig. A2), the assimilated PO_4 would be sustained in the cellular P
components such as polyphosphate, that typically accumulates in particulate P in the PO_4 -depleted regions such as the
western North Atlantic and WNP (Martin et al., 2014; Hashihama et al., 2020).

4.3 Si and N drawdown characteristics

Previous field studies have reported that natural diatom blooms in the subtropical North Pacific were accompanied by high
375 $\text{Si}(\text{OH})_4:\text{DIN}$ drawdown ratios (>1) (Benitez-Nelson et al., 2007; Hashihama et al., 2014). However, in our incubation
experiments, most of the $\Delta\text{Si}(\text{OH})_4:\Delta\text{DIN}$ ratios were less than 1 and involved no significant values of $\Delta\text{Si}(\text{OH})_4$ (Table 3).
Furthermore, there was no significant $\Delta\text{Si}(\text{OH})_4$ even with a large increase (806 cells L^{-1} relative to the control) in diatom
stocks at Station 15 (Table 3 and Figs. 2g and 3c). This increased diatom density is comparable to that in the natural bloom
in the WNP reported by Hashihama et al. (2014). This mismatch between increased stocks and little change in $\Delta\text{Si}(\text{OH})_4$
380 implies that $\Delta\text{Si}(\text{OH})_4:\Delta\text{DIN}$ ratio is not so high in an early stage of diatom blooms. **$\text{Si}(\text{OH})_4$ uptake and release would be**
balanced in the early bloom phase. In addition, low $\Delta\text{Si}(\text{OH})_4:\Delta\text{DIN}$ ratio in the early bloom phase was also observed in a
mesoscale Fe enrichment experiment in the northeastern part of the subarctic Pacific, suggesting no Fe limitation of diatoms
in the early bloom phase (Boyd et al., 2005). Probably, in a Fe- or DIN-depleted late stage of the bloom, selective $\text{Si}(\text{OH})_4$
removal by diatoms (>1 of $\Delta\text{Si}(\text{OH})_4:\Delta\text{DIN}$) occurs through putative biogeochemical processes such as selective Si export
385 (Si pump), anomalous Si uptake associated diatom physiology, and/or Si uptake supported by N_2 fixation (Dugdale and

Wilkerson, 1998; Takeda, 1998; Boyd et al., 2005; Benitez-Nelson et al., 2007; Brzezinski et al., 2011; Krause et al., 2013; Hashihama et al., 2014).

4.4 Possible influences of the subtropical phytoplankton blooms on P* distribution

The present study is the first to reveal basin-wide distributions of surface P* in the oligotrophic Pacific Ocean using
390 nanomolar N+N and PO₄ data (Fig. 5). The distributional pattern of surface P* coincided with that obtained in the upper 120
m using micromolar NO₃ and PO₄ data (Deutsch et al., 2007), indicating that a relatively homogenous P* condition prevails
throughout the water column from the surface to 120 m depth. Based on the conventional concept, the upper layer P* is
likely controlled by N₂ fixation and denitrification (Deutsch et al., 2007). However, by comparing surface P* distribution
with experimentally determined $\Delta\text{DIN}:\Delta\text{PO}_4$ ratios across the subtropical Pacific, we see insights into additional controls on
395 P*. The low (≤ 13.3) and high (≥ 14.1) $\Delta\text{DIN}:\Delta\text{PO}_4$ ratios geographically corresponded to the low and high P* in the WNP
(<50 nM, Stations A, 2, and 5) and the CNP and ESP (>50 nM, Stations 8, 15, 18, and 21), respectively (Fig. 5).

A comparison of the PO₄-intercepts that were determined from the deep-water addition experiments with surface P*
(from ambient concentrations) was conducted (Fig. 6). This cross-comparison was valuable since both represent metrics for
excess PO₄ that remains after DIN exhaustion. The PO₄-intercepts and P* are hereafter referred to as ‘bloom P*’ and
400 ‘ambient P*’, respectively. The bloom and ambient P* showed a similar geographical trend both being relatively low in the
PO₄-depleted WNP (Stations A, 2, and 5) when compared with the PO₄-replete regions (Stations 8, 15, 18, and 21). The one
to two orders of magnitude higher ambient P* than bloom P* in the PO₄-replete regions suggests that, rather than
phytoplankton uptake, denitrification (and also anammox) has a more pronounced influence on setting excess PO₄ in those
regions. This trend may be particularly important in the ESP which is the vicinity of an oxygen minimum zone with active
405 denitrification and anammox conditions (Paulmier and Ruiz-Pino, 2009). Alternately, N₂ fixation may exert an influence on
ambient P* in the PO₄-depleted WNP. However, several studies reported that directly measured N₂ fixation rates are not
consistently high in the WNP compared to other subtropical Pacific regions (Shiozaki et al., 2009; Shiozaki et al., 2010;
Bonnet et al., 2017; Hashihama et al., 2020). Given that natural phytoplankton blooms in the subtropical oceans have a large
impact on nutrient dynamics through new production (Benitez-Nelson et al., 2007; McGillicuddy et al., 2007; Dore et al.,
410 2008), the surplus PO₄ removal by phytoplankton bloom as observed in our experiments might play a significant role in
maintaining low ambient P* in the WNP.

Furthermore, we found the unique result that bloom P* showed negative values in the PO₄-depleted WNP (Stations A,
2, and 5), while the ambient P* did not exhibit negative values (Fig. 6). **The difference between the bloom P* and ambient
P* largely depends on the different N:P consumption ratios of ≤ 13.3 and 16, respectively. If the low N:P consumption ratios
415 (≤ 13.3) are consistently dominant in the PO₄-depleted WNP, alternative P sources other than PO₄ would be required to fully
exhaust DIN. Since lower DOP concentrations and higher alkaline phosphatase activity were observed in the WNP,
compared to other subtropical Pacific regions (Hashihama et al., 2019; Hashihama et al., 2020), active DOP utilization in the
WNP likely contributes to the DIN exhaustion. These perspectives suggest that, in the studies on subtropical nutrient**

biogeochemistry using N:P stoichiometry, the bioavailable fraction of DOP could be an important factor as well as DIN and
420 PO_4 .

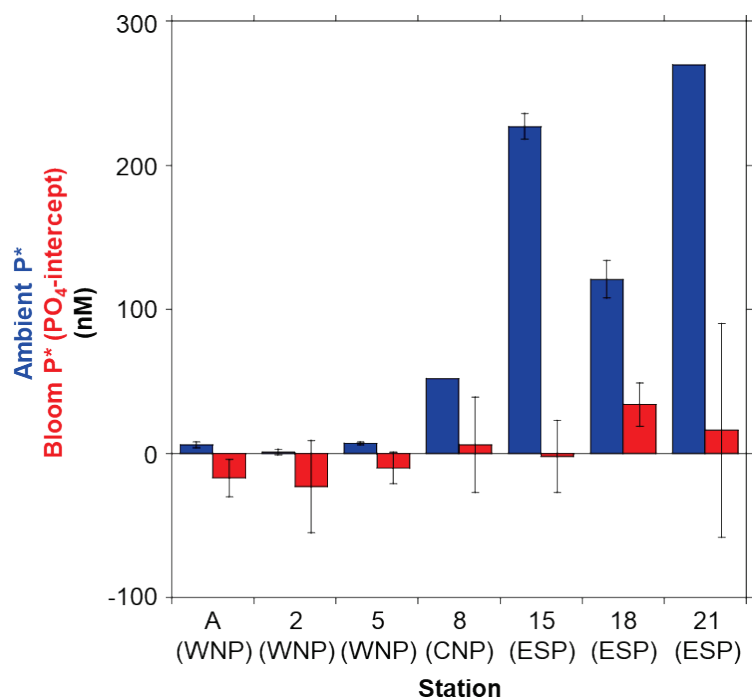


Figure 6. Comparison of ambient P* (blue) and bloom P* (red) at seven stations. The bloom P* is identical to the experimentally determined PO_4 -intercept in Fig. 4. Error bars denote 95% confidence intervals. The ambient P* at Stations 8
425 and 21 have no error bar due to no triplicate data set.

5 Conclusions

By applying highly sensitive analytical methodology, we have revealed nutrient drawdowns and their ratios during the developments of phytoplankton blooms as induced by deep water additions to the surface water of the oligotrophic Pacific Ocean. The $\Delta\text{DIN}:\Delta\text{PO}_4$ ratios showed a geographical variation from low in the PO_4 -depleted WNP to high in the PO_4 -replete ESP. While the $\Delta\text{DIN}:\Delta\text{PO}_4$ ratios in the PO_4 -replete regions were similar to the range from the Redfield ratio to
430 typical subtropical particulate N:P ratio (16-28), those in the PO_4 -depleted regions (7.7-13.3) could not be expected from the conventional phytoplankton N:P ratios. The lower $\Delta\text{DIN}:\Delta\text{PO}_4$ ratios were likely due to the high PO_4 uptake capability of low PO_4 -adapted subtropical phytoplankton. The regional trend in $\Delta\text{DIN}:\Delta\text{PO}_4$ ratios was aligned with that of ambient P* in the oligotrophic Pacific. Although it remains necessary to examine nutrient assimilation characteristics in natural
435 phytoplankton blooms, the regional variation in $\Delta\text{DIN}:\Delta\text{PO}_4$ ratios as observed in our experiments appears to at least control

basin-scale ambient P* distribution in addition to conventional N₂ fixation and denitrification (also anammox). We have also demonstrated that accurate measurements of nanomolar nutrients are powerful tools in investigating trace nutrient dynamics. Further application of these tools to the field and experimental studies would be beneficial for understanding of nutrient biogeochemistry in the oligotrophic ocean.

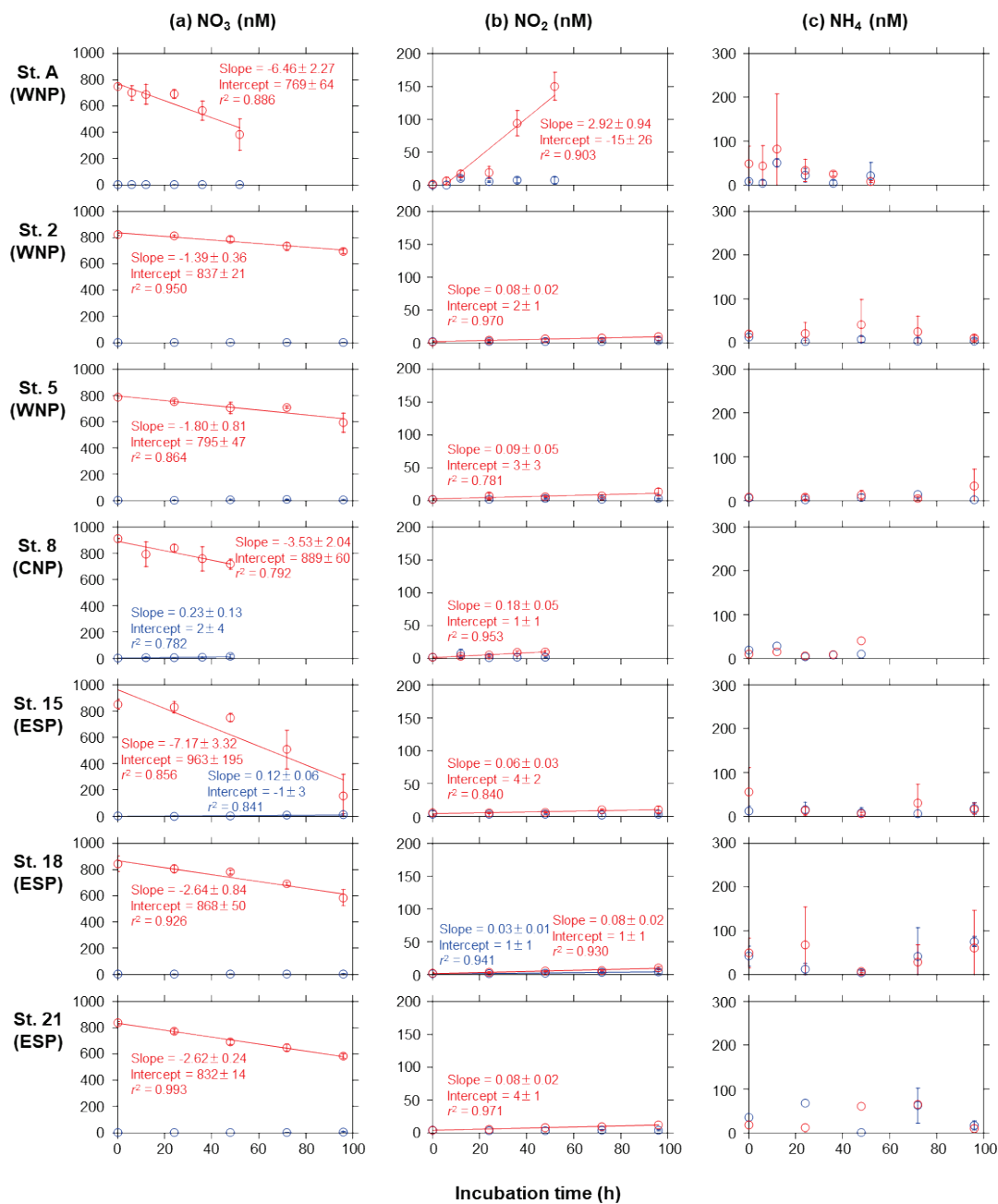
440 Appendices

Appendix methods

The Fe-enrichment incubation experiment was conducted using the surface water (10 m depth) collected at Station 18 in the ESP (Table 1 and Fig. 1). Water sampling was performed using HCl-cleaned Teflon-coated Niskin-X bottles (General Oceanics) on a CTD system (Sea-Bird Electronics) attached to a titanium-armored cable. This sampling procedure succeeded
 445 in avoiding Fe contamination as reported previously (Shiozaki et al., 2018). The surface water was poured into 1.19 L polycarbonate bottles and then Fe was enriched as iron chloride (FeCl₃, Iron Standard Solution Fe 1000, Wako) at the final concentration of 1.8 nM. The triplicate bottles for either Fe-enrichment or control were prepared. These bottles were pre-cleaned sequentially with neutral detergent, 1 M HCl, and 0.3 M hot HCl (for Analysis of Poisonous Metals, Wako), and filled with pure water for a day (Takeda and Obata, 1995). Both the Fe-enriched and control bottles were incubated for 96 h
 450 in the on-deck incubator as described in 2.2. After 96 h, the incubated bottles were sampled for nanomolar nutrients, DON, DOP, and Tchl *a*. Initial samples for nanomolar nutrients, DON, DOP, and Tchl *a* were collected in duplicate directly from the Niskin-X bottles. The samples for nanomolar nutrients, DON, and DOP were processed and analysed as described in 2.3 and 2.4. For Tchl *a* here, a water volume of 100 mL was filtered onto GF/F filters, and the filter samples extracted with *N,N*-dimethylformamide (DMF, Wako) were analysed using a Turner Design fluorometer (Suzuki and Ishimaru, 1990). Student *t*-
 455 test was performed to determine significant differences ($p < 0.05$) between the measured parameter values in the Fe-enriched and control bottles.

Table A1. Results of a Fe-enrichment incubation experiment at Station 18 in the ESP. **Errors represent standard deviations ($n=3$).** Differences between mean values in control and Fe-treatment were insignificant for all parameters (*t*-test, $p > 0.05$).

Parameter	Initial (0 h)	Control after 96 h	Fe-treatment after 96 h
Tchl <i>a</i> (ng L ⁻¹)	31	29±2	30±3
DIN (nM)	7	13±6	9±3
PO ₄ (nM)	133	131±6	133±3
Si(OH) ₄ (nM)	444	438±37	467±26
DON (µM)	3.52	3.57±0.12	3.63±0.14
DOP (µM)	0.12	0.09±0.03	0.09±0.07



460

Figure A1. Temporal changes in concentrations of (a) NO_3 , (b) NO_2 , and (c) NH_4 in the control (blue) and treated (red) bottles during the incubation periods at seven stations. Error bars denote standard deviations ($n=3$). Duplicate or single data are denoted as mean or single values without error bars. Linear regression lines are depicted when significant decreases or increases ($p < 0.05$) in the mean concentrations against time were observed. Errors of slope and intercept represent 95% confidence intervals.

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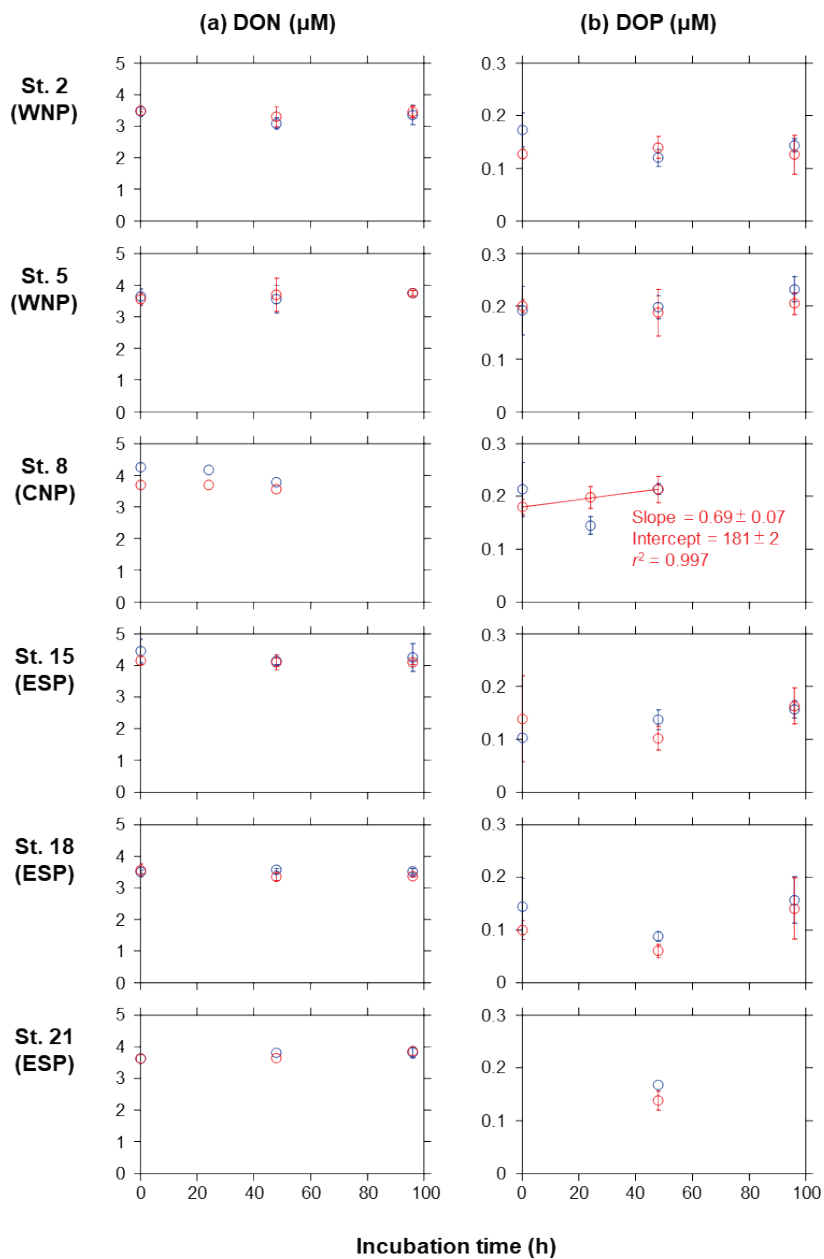
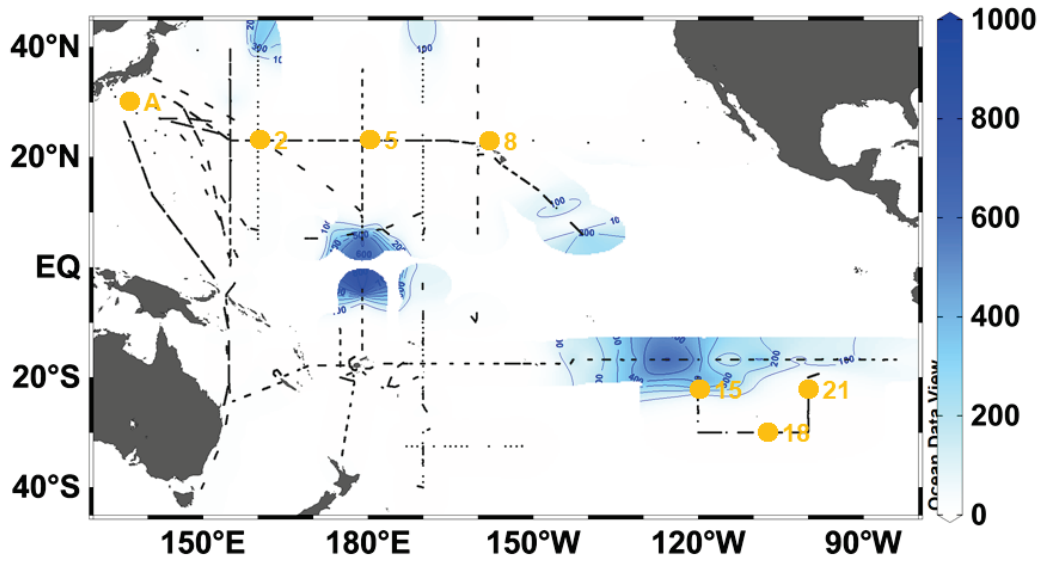


Figure A2. Temporal changes in concentrations of (a) DON and (b) DOP in the control (blue) and treated (red) bottles during the incubation periods at six stations during the KH-11-10 cruise. Error bars denote standard deviations ($n=3$). Duplicate or single data are denoted as mean or single values without error bars. A linear regression line is depicted in the DOP concentrations in the treated bottle at Station 8 as a significant increase ($p<0.05$) in the mean concentration against time was observed. Errors of slope and intercept represent 95% confidence intervals.

(a) Surface N+N (nM)



(b) Surface PO₄ (nM)

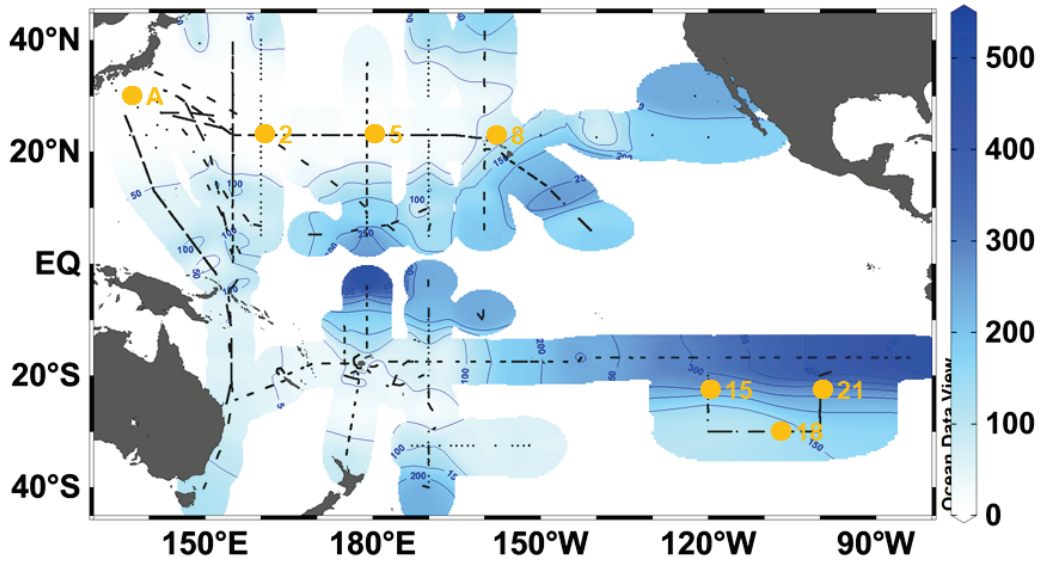


Figure A3. Surface distributions of nanomolar concentrations (<1000 nM) of (a) N+N and (b) PO₄ in the oligotrophic Pacific Ocean (40° N-40° S). Small black dots denote sampling stations for the nanomolar PO₄ and N+N. Large orange circles denote the stations where the incubation experiments were conducted.

Data availability. The data are available upon request to the corresponding author (Fuminori Hashihama).

480 **Author contribution.** FH designed the incubation experiments. KF, HS, HO, and PWB designed the sampling schemes across the subtropical North and South Pacific. FH, TK, JK, and EMSW collected nanomolar nutrient data. SY-T, FH, and JK collected DON and DOP data. FH collected phytoplankton data. FH and IT performed the Fe-enrichment incubation experiment. FH wrote the manuscript. All authors reviewed and approved the manuscript.

485 **Competing interests.** The authors declare that they have no conflict of interest.

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