Impact of ocean acidification on phytoplankton assemblage, growth and
DMS production following Fe-Dust additions in the NE Pacific HNLC waters

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Abstract

Ocean acidification (OA) is likely to have an effect on the fertilizing potential of desert dust in high-nutrient, low-chlorophyll oceanic regions, either by modifying Fe speciation and bioavailability, or by altering phytoplankton Fe requirements and acquisition. To address this issue, short incubations (4 days) of northeast subarctic Pacific waters enriched with either FeSO$_4$ or dust, and set at pH 8.0 (in situ) and 7.8 were conducted in August 2010. We assessed the impact of a decrease in pH on dissolved Fe concentration, phytoplankton biomass, taxonomy and productivity, and the production of dimethylsulfide (DMS) and its algal precursor dimethylsulfoniopropionate (DMSP).

Chlorophyll a (chl a) remained unchanged in the controls and doubled in both the FeSO$_4$-enriched and dust-enriched incubations, confirming the Fe-limited status of the plankton assemblage during the experiment. In the acidified treatments, a significant reduction (by 16-38\%) of the final concentration of chl a was measured compared to their non-acidified counterparts, and a 15\% reduction in particulate organic carbon (POC) concentration was measured in the dust-enriched acidified treatment compared to the dust-enriched non-acidified treatment. FeSO$_4$ and dust additions had a fertilizing effect mainly on diatoms and cyanobacteria. Lowering the pH affected mostly the haptophytes, but pelagophyte concentrations were also reduced in some acidified treatments. Acidification did not significantly alter DMSP and DMS concentrations. These results show that dust deposition events in a low-pH iron-limited Northeast subarctic Pacific are likely to stimulate phytoplankton growth to a lesser extent than in today’s ocean during the few days following fertilization and point to a low initial sensitivity of the DMSP and DMS dynamics to OA.
1 Introduction

The northeast subarctic Pacific is a high-nutrient, low-chlorophyll (HNLC) region characterized by a phytoplankton assemblage dominated in summer by calcifying coccolithophores and extremely high concentrations of the biogenic climate-active gas dimethylsulfide (DMS) (Levasseur et al., 2006; Wong et al., 2006). Several studies have demonstrated that iron (Fe) addition in these Fe-poor waters stimulated phytoplankton growth, in which diatoms often outcompeted other phytoplankton groups, including coccolithophores, and resulted in a decrease in DMS concentrations (Boyd et al., 2005; Hamme et al., 2010; Mélançon et al., 2014). Sources of iron to the northeast subarctic Pacific include vertical mixing, eddies, tidal currents and convection (Cullen et al., 2009; Royer et al., 2010), volcanic ash (Mélançon et al., 2014; Olgun et al., 2011), and desert dust (Boyd et al., 1998; Jickells et al., 2005). Dust, which is considered one of the most important sources, is deposited sporadically mostly in the spring during occasional dust storms originating from the deserts of northern Asia (Duce and Tindale, 1991). A natural strong dust deposition event has been shown to nearly double particulate organic carbon (POC) concentration in the northeast subarctic Pacific in 2001 (Bishop et al., 2002). The importance of eddies and vertical diffusion in the Gulf of Alaska was recently reviewed and found to be greater than previously thought (Crawford et al., 2007; Johnson et al., 2005; Lam and Bishop, 2008) Ongoing ocean acidification (OA) is however likely to compromise our current understanding of the ecosystem’s response to Fe addition by potentially altering Fe bioavailability (Breitbarth et al., 2010; Shi et al., 2010) and phytoplankton physiology and community composition (Tortell et al., 2002).

OA is currently in progress, is measurable and is caused by CO₂ emissions to the atmosphere that end up in the ocean (Gattuso et al., 2013). The ocean has taken up one third of the CO₂ emissions since the beginning of the industrial era. The dissolution of CO₂ in seawater increases the concentration of bicarbonate (HCO₃⁻), protons (H⁺) (thereby decreasing pH) and decreases the concentration of carbonate (CO₃²⁻), leading to calcite and aragonite undersaturation. Studies have shown that calcifiers growing in acidified conditions generally present lower net calcification, although their growth, synthesis and abundance are not generally affected (Kroeker et al., 2013 and references
therein). On the other hand, the increase in dissolved CO$_2$ in seawater could favor the
growth of phytoplankton groups with low surface area to volume ratios (S/V) that are
limited by the diffusion of CO$_2$ across their surface or with low-efficiency carbon
concentrating mechanisms (CCMs) by reducing the energetic cost of carbon (C)
assimilation. Coccolithophores and diatoms generally exhibit low- and high-efficiency
CCMs, respectively (Reinfelder, 2011). Alternatively, fertilisation with FeSO$_4$ usually
favors the growth of diatoms in HNLC waters (Boyd et al., 2007 and references therein).
It is not known whether Fe-Dust deposition will favor diatoms or coccolithophores in the
context of OA. OA could profoundly modify the structure and functioning of a
phytoplankton community typically dominated by calcifying haptophytes.

Studies examining the effects of acidification on the bioavailability of Fe in HNLC
regions have provided contrasting results. Breitbarth et al. (2010) observed a significant
increase in Fe(II) half-life and concentration in response to CO$_2$ enrichment, suggesting
that a lower pH could increase Fe bioavailability. However, Fe bioavailability could also
decrease during acidification due to changes in dissolved Fe speciation. Shi et al. (2010)
observed that complexation of Fe(III) by organic ligands containing acidic, unprotonated
functional groups (e.g. carboxylic acid) is strengthened in response to small decreases in
surface water pH, resulting in decreased inorganic Fe concentrations - the more
bioavailable form of Fe. Furthermore, Fe uptake rates decrease when acquiring
organically complexed Fe – such as Fe(III) bound to desferrioxamine B (DFB) – because
the enzymatic reduction of Fe(III) at the cell surface may release protons (Shi et al.,
2010). Experimental studies with natural communities also yield inconsistent results. A
study combining CO$_2$ and Fe manipulations of a natural northwest subarctic Pacific
community showed a decrease in coccolithophore abundance at higher CO$_2$ levels (750
and 1000 µatm) regardless of the Fe status, but no effect of the CO$_2$ level on diatoms nor
on total chlorophyll $a$ (chl $a$) concentrations (Endo et al., 2013). A similar study
conducted in HNLC waters of the Weddell Sea, Antarctica, showed an increased C-specific primary productivity with increasing CO$_2$ concentrations in Fe-enriched
treatments but not in Fe-depleted treatments (Hoppe et al., 2013).
By altering algal physiology and community composition, OA is likely to influence dimethylsulfoniopropionate (DMSP) and DMS production. DMS originates mostly from the enzymatic cleavage of DMSP, an osmolyte produced by several groups of phytoplankton. DMSP quotas in phytoplankton vary by three orders of magnitude, with coccolithophores and diatoms known as strong and poor producers, respectively (Keller et al., 1989). Results from the few studies which have looked at the impact of OA on DMS production are inconsistent. Several of them have reported a decrease in DMS production in acidified waters (Archer et al., 2013; Arnold et al., 2013; Avgoustidi et al., 2012; Hopkins et al., 2010b). However, an increase in the concentration of DMS at high CO₂ was measured in five bioassays conducted in northwest European waters (Hopkins and Archer, 2014), during the first ten days of the Third Pelagic Ecosystem CO₂ Enrichment Study (PeECE III) (Vogt et al., 2008; Wingenter et al., 2007), and during a mesocosm study conducted in the coastal waters of Korea (Kim et al., 2010). The effect of acidification on DMSP concentration is usually smaller than on DMS, and a greater variability in responses is generally observed: particular or total DMSP increases in some studies (Archer et al., 2013; Arnold et al., 2013), decreases in others (Avgoustidi et al., 2012; Hopkins and Archer, 2014; Hopkins et al., 2010b) or shows no response to increased pCO₂ (Lee et al., 2009; Vogt et al., 2008). The causes for this variability are not well known.

The objective of this study was to determine how a decrease in pH by 0.2 units could influence the impact of Fe delivered as FeSO₄ or Asian dust on the growth and taxonomic composition of the phytoplankton assemblage of the Fe-limited northeast subarctic Pacific in summer, and to explore how these pH-induced changes could affect the production of the climate-active gas DMS.
2 Material and Methods

2.1 Experimental setting and location

On deck incubations were conducted during a cruise along the Line-P transect aboard the Canadian Coast Guard Ship John P. Tully. Water was collected at Ocean Station Papa (OSP) (50°N, 145°W) from 10 m depth on 27 August 2010 using a Teflon® diaphragm compressed air activated pump with Teflon® tubing and filtered through a 200 µm nylon mesh to remove large zooplankton. A CTD profile conducted at the same station 2h after all bags were filled showed a temperature of 13.5°C and a salinity of 32.6 at 10 m depth. Water was incubated in 5 L collapsible bags (Hyclone® Labtainers™). A flow of surface water was continuously pumped through the incubators to keep the temperature at in situ levels. Measured transmittance shows that the incubation bags filtered 55 % of ultraviolet A (UVA) radiation, 70 % of ultraviolet B (UVB) radiation, and 33 % of photosynthetically active radiation (PAR), irradiance corresponding roughly to a depth of 10 m for the 400-600 nm wavelengths (Sasaki et al., 2001). The incubations lasted 4 days and subsampling took place at T0 (0-20 min after acidification and enrichment), T2 (after 2 days), and T4 (at the end of the incubation). All materials in contact with seawater were cleaned to prevent trace-metal contamination according to protocols established by the international GEOTRACES program and available in GEOTRACES’ Methods Manual (Cutter et al., 2010).

2.2 Treatments and acidification protocol

Incubation bags were submitted to six treatments (in triplicate) representing the following combination of dust or Fe addition and acidification: Control, Control+Acid, Fe, Fe+Acid, Dust, Dust+Acid (Table 1). The carbonate system parameters and methods used for acidification were based on the recommendations of Riebesell et al. (2010). The technique chosen was the addition of a strong acid (HCl) and bicarbonate (NaHCO₃). The target value of 750 µatm CO₂ was chosen to reproduce the concentration of CO₂ expected in 2100 following the “business as usual” scenario IS92a by the Intergovernmental Panel on Climate Change (IPCC) (Meehl et al., 2007). Target values of the carbonate system parameters (DIC, pCO₂, alkalinity and pH) were calculated using the MS Excel macro
CO2SYS (Pierrot et al., 2006) (sets of constants: K1, K2 from Mehrbach et al. (1973) refitted by Dickson and Millero (1987), KH2SO4: Dickson (1990), pH scale: seawater scale (mol kg\(^{-1}\) – SW)) and are presented in Table 2. To reach these target values, a final concentration of 122.4 µmol kg\(^{-1}\)\(_{sw}\) of trace-metal clean 6 mol L\(^{-1}\) Seastar\textsuperscript{TM} Baseline HCl solution and a final concentration of 115.1 µmol kg\(^{-1}\)\(_{sw}\) of a trace-metal clean solution of NaHCO\(_3\) were added to each acidified treatment bag through the Luer lock port using a syringe. The piston was activated several times to ensure proper mixing.

2.3 Fe and dust addition

Two sources of Fe were used for the fertilization: FeSO\(_4\) and standardized Asian dust CJ-2. Briefly, CJ-2 dust was collected from the Tengger desert surface soil, roughly sieved and blown through a wind tunnel designed to collect fine particles. Median diameter of CJ-2 dust is 24.1 µm. CJ-2 dust is characterized by a Fe content of 3.02 ± 0.12% and Fe solubility of 0.33% (Ooki et al., 2009). For more information on CJ-2 dust, see Nishikawa et al. (2000) and Hwang and Ro (2006). FeSO\(_4\) was added at the final concentration of 0.6 nmol L\(^{-1}\) and CJ-2 dust was added at the final concentration of 2 mg L\(^{-1}\). Twenty hours prior to their addition in the bags, dust and Fe solutions were prepared by adding CJ-2 dust samples or FeSO\(_4\) to MilliQ water. Proper quantities of Fe-enriched solutions were added to each bag with a syringe through the Luer lock port. The piston was pushed and pulled several times to ensure proper mixing. Then, each bag was gently shaken for 5 minutes to homogenize its content prior to sampling.

2.4 Chemical and biological variables

2.4.1 Carbonate system

Dissolved inorganic carbon (DIC) and alkalinity were measured at T0 and T4. DIC was measured by a coulometric method using a Single operator multi-parameters metabolic analyzer (SOMMA) (Johnson et al., 1993) coupled to a UIC 5011 coulo meter, according to standard protocols (Dickson et al., 2007). Samples were calibrated against Andrew Dickson (Scripps) CRM water batch 101. Alkalinity was measured using an open cell method consisting of a Brinkmann Dosimat 665 an Alpha PHE-4841 glass body
combination electrode according to standard protocols outlined in the Guide to best practices for ocean CO₂ measurements (Dickson et al., 2007). Samples were calibrated against Andrew Dickson (Scripps) CRM water batch 101. Partial pressure of CO₂ (pCO₂) and pH were calculated using the MS Excel macro CO2SYS as described above.

2.4.2 Macronutrients and dissolved Fe

For macronutrients, at each subsampling time, ~10 mL of each bag was filtered in an in-line syringe filter to remove particles (Polycarbonate 0.8 Micron, 25 mm filters) and placed in polystyrene test tubes. The test tubes were immediately frozen at -20°C in an aluminium freezer block. Concentrations of nitrate, silicic acid and phosphate were measured ashore using a Technicon AA autoanalyzer II following the methods described in Barwell-Clarke and Whitney (1996).

Dissolved Fe (DFe) samples were collected at each subsampling time by filtering ~125 mL (0.22 µm) into acid-cleaned low density polyethylene bottles under a clean laminar flow hood or in the trace-metal clean positive pressure plastic tent (the “bubble”) constructed in the main lab as described in Johnson et al. (2005). DFe samples were placed in trace-metal clean, low-density polyethylene bottles and acidified to pH 1.7 for 20-30 h. Samples were then buffered to pH 3.2 using a formic acid-ammonium formate buffer. DFe was quantified according to GEOTRACES protocols available in the Methods Manual (Cutter et al., 2010). All DFe samples were analyzed on board in the “bubble” using a flow injection analysis (FIA) method where Fe is concentrated from the seawater matrix onto a chelating resin and detected by chemiluminescence as first described by Obata et al. (1993) with modifications presented in Obata et al. (1997). Samples were pre-concentrated on a resin column of 8-hydroxyquinoline immobilized on silica gel. The eluent was then combined with ammonia, hydrogen peroxide and luminol. A Hamamatsu photomultiplier tube quantified the light emitted by the reaction of Fe and luminol as it passed through the detection cell. Fe concentration was determined using an external standard curve. Accuracy of the system was checked by regular measurements of the standard reference seawaters SAFe D1 and D2 (Johnson et al., 2007). Our average values of the SAFe D1 and two SAFe D2 standards were 0.63 ± 0.05 nmol L⁻¹ Fe (n = 7),
0.91 ± 0.05 nmol L\(^{-1}\) Fe (n = 6) and 0.90 ± 0.03 nmol L\(^{-1}\) (n = 4), consistent with the community consensus value of 0.67 ± 0.04 nmol L\(^{-1}\) (D1) and 0.93 ± 0.02 nmol L\(^{-1}\) (D2).

### 2.4.3 Pigments and particulate organic carbon

For chl \(a\) determination, 305 mL of water was withdrawn from each bag at each subsampling time. The water was filtered through 25 mm GF/F filters at ≤9.33 kPa vacuum and filters were frozen at -20\(^{\circ}\)C until analysis. Acetone (90 %) was added to extract the pigments 24 h prior to analysis. Pigments were quantified with a Turner 10 AU fluorometer as described in Strickland and Parsons (1972). To characterize the initial pigment composition by High Performance Liquid Chromatography (HPLC), two samples of ca. 2 L were taken directly from the pump when treatment bags were filled at T0. At T4, ca. 1 L was sampled from two incubation bags. Samples were filtered at ≤9.33 kPa vacuum on 47 mm GF/F filters and filters were frozen at -80\(^{\circ}\)C until analysis onshore. Pigments were extracted by placing the filters in 95 % methanol at -20\(^{\circ}\)C in the dark for 24 h prior to analysis. The extracts were filtered through 25 mm diameter polytetrafluoroethylene (PTFE) syringe filters (0.2 µm pore size) and analyzed using a Waters Alliance 2695 (HPLC) system equipped with a 2996 Photodiode Array Detector (PDA) and a reverse phase C8 column (Waters Symmetry), with a pyridine-containing mobile phase (Zapata et al., 2000). Analysis was performed on a 200 µl injection of sample mixed with water in the autosampler at a ratio of 5:1 immediately prior to injection. Pigment concentrations were quantified using commercially available standards (Danish Hydraulic Institute). The initial pigment ratio matrix loaded into the CHEMTAX program (Table 3) was obtained by averaging the minimum and maximum values of pigment : Chl a ratios given in Table 1 of Mackey et al. (1996) and is similar to that used by Suzuki et al. (2002) and Royer et al. (2010) for samples collected in the subarctic North Pacific. Eight algal groups were quantified using the chemotaxonomy program CHEMTAX (Mackey et al., 1996): cyanobacteria, pelagophytes, haptophytes (including coccolithophores), diatoms, dinoflagellates, prasinophytes, cryptophytes and chlorophytes. For a description of the pigment types, see Zapata et al. (2004). Particulate organic carbon (POC) concentrations were measured at T0 and at T4. At T0, three samples of 500 mL of seawater were pumped from the sampling station. At T4, 500 mL
were sampled from each one of the incubation bags. Samples were filtered on pre-combusted 25 mm GF/F and the filters were placed in open cryovials and allowed to dry in an oven at 60°C for 48 h. The cryovials were then capped and kept in drierite until onshore analysis. POC and particulate nitrogen were measured using a mass spectrometer (Delta Plus, Thermo Finnigan Mat) coupled with an elemental analyzer (CE Instrument model 1110).

2.4.4 C and Fe uptake rates

The siderophore desferrioxamine B (DFB) has been used as a model ligand for studying the bioavailability of strongly organically complexed Fe in seawater (e.g. Maldonado and Price 1999; Hutchins et al. 1999; Shi et al. 2010). Iron uptake from DFB occurs through a high affinity Fe transport system, and can reflect the Fe nutritional status of laboratory phytoplankton strains (Maldonado and Price 1999; Maldonado et al. 2006) and natural phytoplankton assemblages (Semeniuk et al. 2009; Taylor et al. 2013; Semeniuk et al. in press). Thus, we examined whether uptake rates of Fe complexed to DFB varied among our treatments with varying Fe bioavailability.

Uptake of 55Fe bound to DFB was performed as previously described (Maldonado and Price 1999; Semeniuk et al. 2009). The 0.5 nM 55Fe (Perkin Elmer) was complexed with 5% excess DFB in pH 3.5 Milli Q for 30 minutes (Maldonado and Price, 1999). The resulting 55FeDFB complex was subsequently equilibrated in 0.22 µm filtered seawater for 2 h. Approximately 250 mL of seawater was subsampled from each incubation bag into acid-cleaned polycarbonate bottles. Just before dawn, the equilibrated 55FeDFB complex and 10 µCi of H14CO3- (Perkin Elmer) were added to each 250 mL bottle. From each assay bottle, 1 mL of sample was taken in order to determine the initial total added activities of 55Fe and 14C. To prevent inorganic 14C from off-gassing in the initial activity sample vial, 500 µL of 5 M NaOH was added.

After 24 h, the content of each bottle was gently filtered onto a 47 mm diameter, 1 µm porosity polycarbonate filter (AMD) under low vacuum pressure (≤70 mm Hg). Just before going dry, the filters were immersed in 5 mL of Titanium-ethylenediaminetetraacetic acid (Ti-EDTA) wash to remove extracellular Fe (Hudson and Morel, 1989). The filters were then rinsed with 10 mL of filtered seawater to remove any
loosely associated tracer. Filters were placed into 7 mL borosilicate scintillation vials, immersed in 5 mL Scintisafe 50% scintillation cocktail, and conserved in the dark until analysis on a Beckman LS65005514 scintillation counter.

Volumetric Fe uptake and C-fixation rates were calculated as described elsewhere (Maldonado and Price 1999; Semeniuk et al. 2009). Previous work with phytoplankton assemblages along Line P has demonstrated that uptake of Fe from DFB by natural phytoplankton communities is linear over 24 h (Maldonado and Price 1999). Thus, Fe uptake rates were calculated assuming the accumulation of 55Fe by cells was linear during the assay. In order to compare Fe uptake rates among treatments, volumetric rates were normalized to the amount of C fixed during the assay. The 14C uptake rates normalised to chl a is used here as an indicator of the growth status of the autotrophic assemblage.

DMS and DMSP concentrations were measured following the techniques described in Royer et al. (2010). Briefly, for DMS, water samples were withdrawn from the bags at every subsampling time in 50-mL serum bottles and analyzed on board using a purge and trap system coupled to a gas chromatograph following methods described in Scarratt et al. (2000). Total DMSP (DMSP\textsubscript{t}) was measured in an unfiltered water sample of 3.5 mL. The samples were acidified with 50 µL of 50% H\textsubscript{2}SO\textsubscript{4} and conserved at 4ºC in the dark until analysis.

2.5 Statistical Analysis

All statistical analyses were run on Statistical Analysis System (SAS) software. The threshold value for determining statistical significance was p < 0.05. Repeated-measures ANOVA were used to test the difference between treatments and the changes in time for the means of biological and chemical variables. Normality of the data was determined using the Shapiro-Wilk test. ANOVA on ranks was used when normality of the data could not be assumed. Differences between the mean concentrations of phytoplankton groups, as measured by HPLC, were assessed using one factor ANOVA. Two-way ANOVAs were used to isolate the effect of one factor (acid, Fe addition).
3 Results

3.1 State of the carbonate system, macronutrients and Fe concentrations

Table 2 presents the average and standard deviation of the four parameters of the carbonate system measured (DIC and alkalinity) and calculated (pCO$_2$ and pH) for each treatment at T0 and T4. Target values of pH and pCO$_2$ in the acidified treatments were reached with averages of 7.80 ± 0.01 and 740 ± 23 µatm CO$_2$ respectively, at T0. DIC values in acidified treatments reached an average of 2139 ± 4 µmol kg$^{-1}$sw, a value 1.5% higher than the target value but consistent among acidified treatments. Alkalinity values in the acidified treatment were 2243 ± 7 µmol kg$^{-1}$sw, a value 2.5% higher than the in situ (and target) value of 2187 µmol kg$^{-1}$sw. DIC, alkalinity and pH values all remained stable during the 4-d incubations. From T0 to T4, pCO$_2$ values varied by 3-12% in the acidified treatments and by 2-15% in the non-acidified treatments, but values of the acidified versus non-acidified treatments remained different from each other.

Initial concentrations of nitrate, silicate and phosphate were 8.0 ± 0.2 µmol L$^{-1}$, 14.2 ± 0.2 µmol L$^{-1}$ and 0.88 ± 0.02 µmol L$^{-1}$. Nutrient concentrations remained high during the course of the experiments, with nitrate, silicate and phosphate decreasing by less than 6%, 4% and 13%, respectively (data not shown).

DFe, operationally defined as the fraction that passes through a 0.22 µm filter, includes soluble and colloidal Fe (de Baar et al., 2005). DFe concentrations started at 0.41 ± 0.09 nmol L$^{-1}$ and 0.21 ± 0.02 nmol L$^{-1}$ in the Control and Control+Acid treatments, respectively, and decreased to 0.07 ± 0.01 nmol L$^{-1}$ and 0.04 ± 0.01 nmol L$^{-1}$ over the time course of the experiment (Fig. 1a). In the Fe and Fe+Acid treatments, DFe started at 0.65 ± 0.32 nmol L$^{-1}$ and 0.47 ± 0.23 nmol L$^{-1}$, respectively. The DFe concentration decreased to ca. 0.11 nmol L$^{-1}$ on day 2 and to ca. 0.06 nmol L$^{-1}$ on day 4 in both treatments (Fig. 1b). In the Dust treatment, DFe started at 0.28 ± 0.10 nmol L$^{-1}$, decreased to 0.12 ± 0.01 nmol L$^{-1}$ at T2 and remained at this level at T4. In the Dust+Acid treatment, DFe started at 0.18 ± 0.05 nmol L$^{-1}$, increased to 0.28 ± 0.01 nmol L$^{-1}$ at T2 and decreased slightly to 0.21 ± 0.07 at T4 (Fig. 1c).
3.2 Plankton biomass

Average initial chl $a$ concentration in all treatments was $0.39 \pm 0.03 \text{µg L}^{-1}$ (Fig. 2). In the Control and Control+Acid treatments, chl $a$ concentration remained stable for the length of the experiment (Fig. 2a). In the Fe treatment, chl $a$ concentrations reached $0.76 \pm 0.16 \text{µg L}^{-1}$ after 4 days, a value significantly higher than measured in the control at the same time ($p$-value = 0.0269; Fig. 2b). In the Fe+Acid treatment, chl $a$ concentrations increased to $0.58 \pm 0.15 \text{µg L}^{-1}$ after 4 days, a value not significantly different from the one reached at the end of the Fe treatment. The addition of dust also had a significant stimulating effect on phytoplankton growth compared to the Control ($p$-value: 0.0071) with chl $a$ reaching $0.88 \pm 0.23 \text{µg L}^{-1}$ at T4 (Fig. 2c). The chl $a$ concentration reached at T4 in the Dust treatment was not statistically different than the one reached in the Fe treatment. In the Dust+Acid treatment, chl $a$ concentration reached $0.74 \pm 0.01 \text{µg L}^{-1}$ at T4, a value again not significantly different from the concentrations reached at the end of the Dust treatment (Fig. 2c). Although the difference is not significant, a trend appears in chl $a$ concentrations: chl $a$ concentrations are always slightly lower in the acidified vs non-acidified treatment. In order to detect any interactive effect of Fe or acidification on chl $a$ and to further explore these apparent trends in chl $a$, all treatments were grouped and tested with a two-factor ANOVA. The first factor, enrichment, had three possible states (Fe, Dust, nil) and the second factor, acid, had two possible states (+ acid, control). This analysis showed a significant effect of the Fe enrichment ($p$-value = 0.0060) and a significant effect of the acidification ($p$-value = 0.0385) on chl $a$ concentration. However, no combined effect (synergic or antagonistic) was detected with the two factors.

Initial POC concentration was $75.3 \pm 11.2 \text{µg L}^{-1}$ (not shown) and increased in all treatments including the Control. After 4 days, POC concentrations were similar in the Control ($125.5 \pm 0.3 \text{µg L}^{-1}$) and Control+Acid ($122.5 \pm 17.6 \text{µg L}^{-1}$) treatments (Fig. 2a). The average POC concentration at T4 in the Fe treatment ($169.2 \pm 55.8 \text{µg L}^{-1}$) was not statistically different than in the Control. Final POC concentration in the Fe+Acid treatment ($189.3 \pm 29.2 \text{µg L}^{-1}$) was not significantly different than in the Fe treatment, but significantly higher than in the Control (Fig. 2b). The highest POC concentration was measured in the Dust treatment ($286.7 \pm 39.7 \text{µg L}^{-1}$) and lowering the pH resulted in a
24% decrease in POC concentration at T4 (Dust+Acid treatment: 217.4 ± 2.0 µg L⁻¹) (Fig. 2c).

3.3 Taxonomy

The initial phytoplankton biomass (T0) was dominated by chlorophytes (37% of total chl a), followed by haptophytes (31%), pelagophytes (19%) and dinoflagellates (13%) (from Fig. 3). Prasinophyte and cryptophyte biomarkers were not detectable during our experiments. Dinoflagellates were present in low concentrations at T0 and became undetectable at T4 in the Control and in all treatments (Fig. 3b). In contrast with the dinoflagellates, diatoms and cyanobacteria were below the detection limit at T0 and became detectable at T4 in the Control and in all treatments (Fig. 3a, f). These changes in community composition in the control show that the sampling and/or incubation conditions had a negative effect on the growth of dinoflagellates and a positive effect on the growth of diatoms and cyanobacteria. Figure 3 shows that diatoms were responsible for most of the increases in chl a measured in the Fe, Fe+Acid, Dust and Dust+Acid treatments compared to the control. The only treatment to show a statistically significant difference with the Control is the Dust treatment, which had significantly higher concentrations of chl a attributable to diatoms and cyanobacteria than the Control at T4. No statistical difference between any treatment and its acidified counterpart could be detected.

3.4 Carbon and Fe uptake rates at T4

At T4, C assimilation rate was 92 ± 50 nmol C L⁻¹ h⁻¹ in the Control (Fig. 4a). In the Control+Acid treatment, C assimilation was 195 ± 21 nmol C L⁻¹ h⁻¹, a value significantly higher than in the Control. C assimilation rates in the Fe and Fe+Acid were significantly higher than the Control, but not different from each other with values of 189 ± 23 nmol C L⁻¹ h⁻¹ and 243 ± 66 nmol C L⁻¹ h⁻¹, respectively. C assimilation rate in the Dust treatment was similar to the Control with 59 ± 24 nmol C L⁻¹ h⁻¹. Lowering the pH significantly increased C assimilation rate in the Dust+Acid treatment (145 ± 61 nmol C L⁻¹ h⁻¹) compared to the Dust treatment.
The chl *a*-specific C assimilation rate (size fraction > 1 µm) in the Control at T4 was 0.23 ± 0.01 µmol C µg chl *a*⁻¹ h⁻¹ (Fig. 4b). Lowering the pH increased significantly the chl *a*-specific C assimilation rate to 1.03 ± 0.13 µmol C µg chl *a*⁻¹ h⁻¹ in the Control+Acid treatment. A similar albeit less pronounced pH-induced increase in chl *a*-specific C assimilation was observed between the Fe (0.25 ± 0.03 µmol C µg chl *a*⁻¹ h⁻¹) and the Fe+Acid treatments (0.43 ± 0.15 µmol C µg chl *a*⁻¹ h⁻¹). The Dust and Dust+Acid treatments had chl *a*-specific C assimilation rates of 0.07 ± 0.04 and 0.17 ± 0.10 µmol C µg chl *a*⁻¹ h⁻¹, respectively. It is noteworthy that although comparable biomasses were achieved after 4 days in the FeSO₄ or Dust treatments, the chl *a*-specific C assimilation rate was significantly lower in the Dust treatment as compared to the Fe treatment and in the Dust+Acid compared to the Fe+Acid treatment.

At T4, FeDFB uptake rates normalized to C assimilation were 0.94 ± 0.55 µmol Fe mol C⁻¹ in the Control, 0.34 ± 0.07 in the Control+Acid treatment, 0.33 ± 0.16 in the Fe treatment and 0.39 ± 0.20 in the Fe+Acid treatment (Fig. 4c). In the dust treatment, FeDFB uptake rate normalized to C assimilation was high (1.44 ± 0.66 µmol Fe mol C⁻¹ h⁻¹), also suggesting a rapid return to Fe-limiting conditions in this treatment. Lowering the pH decreased the FeDFB uptake rates normalized to C in the Dust+Acid treatment (0.55 ± 0.39 µmol Fe mol C⁻¹), but had no effect on the Fe+Acid treatment (0.39 ± 0.20 µmol Fe mol C⁻¹).

### 3.5 DMSPₜ and DMS

Initial average DMSPₜ concentration was 39.8 ± 3.6 nmol L⁻¹ and decreased between 10 and 21 nmol L⁻¹ at T4 in all treatments (Fig. 5a, b, c). Lowering the pH resulted in no significant change in DMSPₜ concentrations in the control and in the Fe and Dust treatments. Initial average DMS concentration in all treatments was 10.0 ± 1.1 nmol L⁻¹ and decreased in all treatments to reach concentrations varying between 3.4 – 6.6 nmol L⁻¹ at T4 (Fig. 5d, e, f). Neither the addition of Fe/dust nor the decrease in pH had an effect on DMS concentrations.
4 Discussion

4.1 Considerations on the experimental protocol

The experimental approach used in this study has limitations, some of which deserve to be addressed forefront in order to avoid misinterpretations of the results. First, the sampling and incubation procedures negatively affected the growth of dinoflagellates. For this reason, no conclusion could be drawn on the effect of the treatments on dinoflagellates. For all other taxa, the influence of the treatments could only be addressed by comparing the samples collected at T4. Second, the abrupt acidification rate imposed to the plankton assemblage during our study is not representative of the slow process that is currently taking place in the ocean. Hence, acclimation and adaptation to acidification which will most probably take place in the natural system cannot take place during our transient and short experiment. Transient experiments, in the manner conducted here, are nevertheless useful to characterize the direct impact of OA on Fe bioavailability and to observe short-term resistance/sensitivity of organisms to OA. It is likely that organisms capable of withstanding rapid decreases in pH will also display tolerance to a more gradual decrease in pH.

4.2 Initial in situ conditions and impact of acidification

Oceanic conditions encountered during the experiments were typical of this part of the northeast Pacific and time of year. Macronutrients and chl a concentrations were high and low, respectively, indicative of the HNLC conditions characterizing the Gulf of Alaska in summer (Harrison et al., 1999; Hopkinson et al., 2010). DMS concentrations were high, but usual for this region in the summer (Wong et al., 2005). The DFe concentration of 0.4 nmol L\(^{-1}\) measured in the Control at T0 was higher than expected, but the Fe-limited status of the plankton community was confirmed by the absence of chl a accumulation in the Control and the increase in chl a induced by the addition of FeSO\(_4\) (Fig. 2a, b). Also as expected for this time of the year, the greatest contributors to total chl a included chlorophytes, haptophytes and pelagophytes (Fig. 3), while diatoms represented minor contributors. Hence, the combined influences of dust and pH on phytoplankton growth, taxonomy, and DMS production reported in this study can be extrapolated to the
northeast subarctic Pacific summer conditions. However, for the reason mentioned above, our protocol does not allow us to draw conclusions on how dinoflagellates, which represented ca. 13 % of the autotrophic biomass in situ, respond to OA.

The abrupt decrease in pH (by 0.2 units) and increase in pCO$_2$ (by 335 µatm) had no detectable effect on the Fe-limited phytoplankton biomass and community structure (Figs. 2a and 3c). During a comparable experiment conducted in the same region and under similar oceanographic conditions (HNLC waters, phytoplankton dominated by haptophytes and chlorophytes), Hopkinson et al. (2010) showed that increasing CO$_2$ to 760-1204 µatm had little effect on chl a, nutrient drawdown, or phytoplankton growth rates after 5 days in Fe-limited conditions. In their experiment, they did not observe a decrease in the biomass of haptophytes as during our study, but the absence of present-day CO$_2$ control (observations restricted to low and high CO$_2$ treatments) limits the comparison between the two studies. Another similar experimental study conducted in the northwest subarctic Pacific revealed a small decrease in haptophyte relative biomass at high CO$_2$ levels (1000 µatm) compared to low CO$_2$ level (180 µatm) after 6 days and an increase in diatoms biomass in all treatments but no pCO$_2$-related statistical change in the abundance of diatoms and total chl a after 5 days (Endo et al., 2013). Another experiment conducted with water from the diatom-dominated Fe-limited Bering Sea has shown a negative effect of elevated CO$_2$ on diatoms (Sugie et al., 2013, Endo et al., 2015), which was not observed in the Fe-enriched treatments. Thus, albeit the differences mentioned above, results from these three experiments suggest that pH expected toward the end of this century will only have a small negative impact, if any, on total autotrophic biomass in the HNLC waters of the subarctic North Pacific.

Acidification resulted in the up-regulation of C assimilation in the control (Fig. 4a,b). Such pH-induced increases in C assimilation have previously been reported in pH manipulation experiments (Riebesell et al., 2007; Tortell et al., 2008). During their study in the same oceanic region, Hopkinson et al. (2010) reported an increase in photosynthetic efficiency in their low pH treatment, an increase they attributed to energy savings from down-regulation of the CCMs. Increasing CO$_2$ concentrations (lowering
pH) could have resulted in a down-regulation of this costly mechanism, freeing energy for other metabolic pathways such as C assimilation.

Unexpectedly, the low pH-induced up-regulation of C assimilation measured at day 4 did not directly translate into an increase in POC in the non-Fe-enriched waters. In point of fact, acidifying non-Fe-enriched waters (Control+Acid treatment) had no effect on chl a or POC concentrations but resulted in a 2-fold and 4-fold increase in absolute and chl a-normalized C assimilation rates, respectively, after 4 days (Fig. 4). The absence of a higher POC concentration in the Control+Acid treatment during our study suggests that the newly assimilated C was not converted into biomass or that a loss mechanism would impede the buildup of POC. Such mechanisms could include increased grazing by micrograzers or DOC exudation and subsequent transparent exopolymer particle (TEP) formation. An increase in C uptake without biomass accumulation under acidified conditions has previously been observed during a similar 12-day experiment (Riebesell et al., 2007). This unexpected result was attributed to the release of DOC and subsequent formation of TEPs which are known to accelerate particle aggregation and sinking. Similar significant stimulation of DOC production at elevated CO₂ was reported by Engel et al. (2014) in coastal waters but only a weak and inconsistent CO₂ induced decrease in DOC production was observed by Yoshimura et al. (2014). In the absence of sinking as in our experiment, the aggregation of DOC into TEP may only explain the absence of increase in POC if a large proportion of the TEP produced adsorbed on the walls of the incubation bags. Even though our measurements do not allow identifying the fate of the increased assimilated C in the high pCO₂ treatment, they point toward a perturbation of the C cycling, either by an increase in DOC exudation or grazing. If confirmed, such pH-induced modification of C cycling and pools in HNLC waters could have important impacts on microbial dynamics and C export.

Overall, our results show that OA in the HNLC waters of the northeast subarctic Pacific may initially negatively impact the growth of haptophytes but stimulate phytoplankton C assimilation by the Fe-limited cells. In spite of these effects at the cellular and taxonomic levels, lowering the pH had little effect on the net accumulation of biomass (chl a and POC) after 4 days.
4.3 Dust fertilization in a high CO\textsubscript{2} northeast subarctic Pacific

During the 4 days of the experiment, dust fertilization had the same stimulating effect on net chl \textit{a} production as the addition of FeSO\textsubscript{4}. This similarity confirms that the phytoplankton assemblage was Fe limited when the study was conducted, and that Fe was responsible for the stimulating effect of dust on phytoplankton growth (Fig. 2). However, the average final POC concentration in the Dust treatment was 69% higher than in the FeSO\textsubscript{4} treatment, suggesting a more efficient Fe stimulation of phytoplankton C assimilation in the former. These results suggest that 2 mg L\textsuperscript{-1} of CJ-2 dust releases at least as much bioavailable Fe during the first 4 days of the experiment as the addition of 0.6 nmol L\textsuperscript{-1} FeSO\textsubscript{4}. We can only speculate as to why Fe contained in dust stimulated more growth than FeSO\textsubscript{4}. Part of the answer may be related to the speed at which the Fe is released from dust, a relatively low release rate allowing a more efficient utilization of Fe. In comparison, and as can be seen in Fig. 1, FeSO\textsubscript{4} is rapidly utilized and probably scavenged by particles and ligands following fertilization. Diatoms, cyanobacteria and, to a lesser extent pelagophytes, benefited the most from the dust enrichment, reaching higher group-specific chl \textit{a} concentrations than the Control incubation at T4. Except dinoflagellates, which did not thrive in the Control, all other groups maintained their biomass in the Dust treatment. This response to Fe addition is comparable to the one reported for previous small and large-scale Fe fertilization experiments conducted in the Gulf of Alaska showing an initial increase in the abundance of major taxa and a dominance of diatoms (Boyd et al., 1996; Levasseur et al., 2006; Marchetti et al., 2006; Martin and Fitzwater, 1988; Mélançon et al., 2014). We saw no clear difference in the structure of the phytoplankton assemblage whether dust or FeSO\textsubscript{4} was used as fertilizer. In that regard, FeSO\textsubscript{4} seems to be a good substitute for dust in studies of the early response (< 4 days) of plankton communities to dust deposition in the northeast subarctic Pacific.

DFe measurements were poor indicators of Fe bioavailability following dust deposition in our study. In contrast with the FeSO\textsubscript{4} treatment where almost all the added Fe was still present and measured in the dissolvable pool at T0 (ca. 20 min. following the addition), DFe concentrations remained low and near \textit{in-situ} levels during the entire length of the
dust-addition experiments (Fig. 1b, c). Considering that the addition of Dust did stimulate algal growth, the low and constant concentration of DFe in the Dust treatment suggests that the release of bioavailable Fe from dust was matched by bacterial and phytoplankton Fe acquisition. Re-adsorption of the released Fe by the dust particles may also be responsible for the low levels of DFe measured during the experiment (Ye et al., 2011). These results also show that DFe may not be a good indicator of dust deposition events in the oceanic environment where the release of DFe from the low concentrations of dust deposited is unlikely to exceed bio-uptake and re-adsorption on particles. Similar conclusions were reached in a previous study (Mélançon et al., 2014), where total dissolvable Fe (TDFe) was also measured and shown to be a better indicator of Fe bioavailability than DFe.

Our results suggest that after a period of active growth, phytoplankton in the Dust treatment became Fe-limited 4 days into the experiment. This conclusion is supported by the very low absolute and chl a normalized C fixation rates in the Dust treatment as compared to the Control and FeSO₄ treatments at T4 (Fig. 4a), as well as by the FeDFB uptake rates normalized to C uptake rates which were higher in the Dust treatment than in the Control (Fig. 4b). Dust particles are known to efficiently adsorb Fe (Ye et al., 2011). The rapid return to Fe deficiency in the Dust treatment may thus result from a combination of increased Fe demand and re-adsorption of Fe onto dust particles. This explanation implies however that the re-adsorbed Fe becomes less prone to desorption, which needs to be demonstrated. These results suggest that the influence of the Fe released from dust lasted less than 4 days during our experiments. In natural environments, this period of influence may be even shorter due to fast sinking of larger dust particles. Based on Stoke’s Law and assuming our dust particles were spherical, CJ-2 dust particles may sink at an average speed of 32 m/day, which would take particles out of a 60 m-deep euphotic zone in ~2 days. Due to the combined effect of both Fe re-adsorption on particles and fast sinking, the impact of natural dust deposition may thus be of relatively short duration in the environment, similar to the time-frame of our in vitro study.
Decreasing the pH resulted in a slightly lower biomass (chl *a* by 16% and POC by 15%) in the Dust+Acid treatment than in the Dust treatment (Fig. 2c). The decrease in biomass corresponded to decreasing trends in the contribution of haptophytes and to a lesser extent, of pelagophytes and cyanobacteria to total chl *a* concentrations. As discussed above, a likely explanation for the lower biomass reached after 4 days is the negative effect of acidification on the growth of the coccolithophores (Engel et al., 2005; Harvey et al., 2013; Kroeker et al., 2010). C assimilation rates (absolute and chl *a*-normalized) were ca. 2-fold higher in the Dust+Acid treatment than in the Dust treatment at T4, probably reflecting the stimulating effect of higher CO₂ concentrations on diatom C assimilation. These results reinforce the aforementioned hypothesis that the up-regulation of C assimilation was paired with an increased particulate C loss via enhanced C exudation from the cells or increased grazing.

It is obvious from Figure 2 that the natural variability among the bags of a same treatment as well as the short length of the incubations limited our capacity to statistically distinguish trends resulting from the two treatments (i.e. Fe/dust addition and acidification). The two-factor ANOVA allowed us to increase the statistical power and detect a negative effect of acidification on chl *a* concentration after 4 days. This suggests that OA will not increase Fe bioavailability to natural HNLC phytoplankton communities. However, it does not preclude the possibility that acidification may have exacerbated Fe limitation in our experiment. If the effect of acidification on the growth of calcifying haptophytes was expected, it is not expected in the case of diatoms, pelagophytes and cyanobacteria. Since these non-calcifying organisms also presented a lower biomass in acidified treatments, it is likely that Fe bioavailability has been reduced via interactions between pH, ligands and Fe speciation.

### 4.4 Impact of acidification and dust deposition on DMSP₁ and DMS

The general decrease in DMSP₁ and DMS concentrations measured in all treatments likely reflects the loss of DMSP-rich dinoflagellates due to sampling and/or bottle effect and their replacement by diatoms with low DMSP quotas. In spite of the increase in chl *a* and POC measured in the FeSO₄ and Dust treatments compared to the Control, the alleviation of Fe limitation had no impact on the concentrations of DMSP₁ or DMS. This
can be explained by the lack of significant difference in the abundance of DMSP-rich haptophytes in the Control, FeSO$_4$ and Dust treatments at T4. Indeed, the addition of FeSO$_4$ or dust mostly stimulated the growth of DMSP-poor diatoms, which would have little effect on DMSP$_t$ concentrations.

Our results show no statistical differences between DMSP$_t$ concentrations between the acidified and non-acidified treatments after 4 days. Although the short duration of our incubations may explain this lack of response, other studies conducted over longer period of time have reported similar results. For example, lowering the pH had no effect on DMSP$_t$ concentrations during the nutrient-stimulated bloom of a community from a Norway fjord (Vogt et al., 2008). In their 24-day experiment the absence of effect was attributed to the resistance of the planktonic community considering that similar increases in chl $a$ concentration and species succession were observed in all CO$_2$ treatments. Studies reporting pH-induced changes in DMSP$_t$ concentrations are usually associated with alterations of the structure of the phytoplankton assemblage after several days. For example, a low pH-induced increase in dinoflagellates after 13 days resulted in higher DMSP$_t$ concentrations during a mesocosm study with Arctic water (Archer et al., 2013).

Lowering the pH had no measurable effect on DMS concentrations, a result probably also related to the short duration of the experiment. Previous studies showed either an increase or decrease in DMS concentrations with acidification, but these differences become measurable late in the experiment at or after the peak of the blooms. For example, during the experiment PeECE III in Raunefjorden, Norway, Wingenter et al. (2007) observed an increase in time-integrated average amount of DMS at high-CO$_2$, but the pH-related differences could only be observed after 6 days. In this case, the authors attributed the pH effect to a difference in viral attack and phytoplankton lysis at the chl $a$ peak, a situation that is not likely to have occurred in our short incubations. Contrastingly, a decrease in DMS concentrations with high CO$_2$ was observed during other mesocosm studies conducted in the same fjord (Hopkins et al., 2010; Avgoustidi et al. 2012). These authors suggested that the dominance of flagellates and picoeucaryotes during their study as compared to coccolithophores during PeECE III could explain the divergent responses observed in regards of DMS production.
Archer et al. (2013) also measured a decrease in DMS concentration under high CO$_2$ conditions in Arctic waters, a decrease they attributed to an increase in bacterial production and decrease in DMS yield. It should be noted that none of the experiments described above reports an effect of CO$_2$ on DMS in the first 4 days. The sole study so far showing a rapid effect of a decrease pH is by Hopkins and Archer (2014) who measured a decrease in DMSP and an increase in DMS after 4 days during shipboard bioassays experiments with NW European waters. In that case, the changes in DMSP and DMS were associated with a rapid decline in the abundance of small cells in the acidified treatments and a possible cellular release and cleavage of DMSP to DMS. Their results show nonetheless a regional variability of the responses. The lack of response of DMS concentration to pCO$_2$ in our incubations might reflect this natural variability and a particular resistance of our initial community to acidification.
5 Conclusion

This study demonstrates that the degree of OA expected to occur by the end of the century is likely to have a detectable but minor impact on the short-term response of Fe-limited planktonic communities to sporadic atmospheric Fe-dust depositions. The addition of FeSO$_4$ and Asian dust stimulated the growth of all major phytoplankton groups during our 96-h incubations, thereby confirming the Fe-depleted status of the community and the potential of natural dust deposition for stimulating phytoplankton growth in this HNLC region. In the acidified treatments, Fe in the form of FeSO$_4$ or Asian dust still had a fertilizing effect on the algal assemblage, but to a lesser extent than in the non-acidified treatment. The trends reported here suggest that OA could moderate the growth response of pelagophytes and haptophytes to dust deposition. The fact that non-calcifying taxa were also affected by acidification suggests that the lower pH possibly interferes with ligands, Fe speciation and transporters to reduce Fe bioavailability in these HNLC waters. Finally, our results suggest a low sensitivity of the DMS dynamics to acidification, both under Fe-limited and Fe-replete conditions, in the northeast subarctic Pacific. In order to understand the mechanisms behind this apparent resistance, studies on DMSP and DMS phytoplankton and bacterioplankton metabolisms during longer incubation periods are advisable.
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References


Dickson, A. G.: Standard potential of the reaction: \( \text{AgCl(s)} + \text{H}_2(\text{g}) = \text{Ag(s)} + \text{HCl(aq)} \), and the standard acidity constant of the ion \( \text{HSO}_4^- \) in synthetic sea water from 273.15 to 318.15 K, J. of Chem. Thermodyn., 22, 113-127, 1990.


Hopkins, F. E. and Archer, S. D.: Consistent increase in dimethyl sulfide (DMS) in response to high CO$_2$ in five shipboard bioassays from contrasting NW European waters, Biogeosciences, 11, 4925-4940, 2014.


Royer, S. J., Levasseur, M., Lizotte, M., Arychuk, M., Scarratt, M. G., Wong, C. S.,
Lovejoy, C., Robert, M., Johnson, K., Pena, A., Michaud, S., and Kiene, R. P.: Microbial
dimethylsulfoniopropionate (DMSP) dynamics along a natural iron gradient in the

western subarctic gyre and alaskan gyre in the subarctic North Pacific and the southern

Scarratt, M., Cantin, G., Levasseur, M., and Michaud, S.: Particle size-fractionated
kinetics of DMS production: where does DMSP cleavage occur at the microscale?, J. Sea

Semeniuk, D. M., Cullen, J. T., Johnson, W. K., Gagnon, K., Ruth, T. J., and Maldonado,
M. T.: Plankton copper requirements and uptake in the subarctic Northeast Pacific Ocean,

Semeniuk, D.M., Taylor, R.L., Bundy, R.M., Johnson, W.K., Cullen, J.T., Robert, M.,
Barbeau, K.A., and M.T. Maldonado.: Iron–copper interactions in iron-limited
phytoplankton in the northeast subarctic Pacific Ocean, Limnol. Oceanogr. doi:
10.1002/lno.10210, in press.

Shaked, Y., Kustka, A. B., and Morel, F. M. M.: A general kinetic model for iron


Strickland, J. D. H. and Parsons, T. R.: A practical handbook of sea-water analysis (2nd

Maldonado, M. T.: Colimitation by light, nitrate, and iron in the Beaufort Sea in late


Table 1: Description of Fe/dust additions and acidification status of each treatment. All treatments were conducted in triplicate.

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<th>Treatment</th>
<th>Addition of Fe</th>
<th>Acidification</th>
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<tbody>
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<td>Control</td>
<td>No</td>
<td>No</td>
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<tr>
<td>Control+Acid</td>
<td>No</td>
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</tr>
<tr>
<td>Fe</td>
<td>FeSO$_4$ (+ 0.6 nmol L$^{-1}$)</td>
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</tr>
<tr>
<td>Fe+Acid</td>
<td>FeSO$_4$ (+ 0.6 nmol L$^{-1}$)</td>
<td>Yes</td>
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<tr>
<td>Dust</td>
<td>CJ2 dust (+ 2.0 mg L$^{-1}$)</td>
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<td>CJ2 dust (+ 2.0 mg L$^{-1}$)</td>
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Table 2: Values of DIC, alkalinity, pH and pCO$_2$ in each treatment at T0 and T4.

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<tr>
<th></th>
<th>DIC ($\mu$mol kg$^{-1}_{sw}$)</th>
<th>Alkalinity ($\mu$mol kg$^{-1}_{sw}$)</th>
<th>pH</th>
<th>pCO$_2$ (µatm)</th>
</tr>
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<td><strong>Target values</strong> (acidified)</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>2107</td>
<td>2187 (in situ)</td>
<td>7.78</td>
<td>750</td>
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<tr>
<td><strong>T0</strong></td>
<td>T4</td>
<td>T0</td>
<td>T4</td>
<td>T0</td>
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<tr>
<td>Control</td>
<td>Avg 1998</td>
<td>1998</td>
<td>2184</td>
<td>2180</td>
</tr>
<tr>
<td></td>
<td>SD 1</td>
<td>2</td>
<td>8</td>
<td>16</td>
</tr>
<tr>
<td>Control+Acid</td>
<td>Avg 2141</td>
<td>2147</td>
<td>2244</td>
<td>2248</td>
</tr>
<tr>
<td></td>
<td>SD 5</td>
<td>8</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>Fe</td>
<td>Avg 1992</td>
<td>1989</td>
<td>2186</td>
<td>2187</td>
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<tr>
<td></td>
<td>SD 2</td>
<td>3</td>
<td>2</td>
<td>5</td>
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<tr>
<td>Fe+Acid</td>
<td>Avg 2137</td>
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<td>2240</td>
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<td></td>
<td>SD 5</td>
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<td></td>
<td>SD 3</td>
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Table 3: Biomarker pigment initial ratio matrix for CHEMTAX: Chl a ratios for eight algal groups.

<table>
<thead>
<tr>
<th>Class / Pigment</th>
<th>Chl c3</th>
<th>Chl c2</th>
<th>Peri</th>
<th>19'-but</th>
<th>Fucox</th>
<th>Prasinox</th>
<th>Violax</th>
<th>19'-hex</th>
<th>Diadinox</th>
<th>Allox</th>
<th>Zeax</th>
<th>Chl_b</th>
<th>Chl_a</th>
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</thead>
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<td>Prasinophytes</td>
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<td>0</td>
<td>0</td>
<td>0</td>
<td>0.360</td>
<td>0.114</td>
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<td>0</td>
<td>0</td>
<td>0.142</td>
<td>0.888</td>
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<td>0</td>
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<td>0</td>
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<td>0</td>
<td>0</td>
<td>0</td>
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<td>1</td>
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<td>0</td>
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<tr>
<td>Dinoflagellates</td>
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<td>0.532</td>
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<td>0</td>
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<td>0.192</td>
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**Abbreviations:** Chl c3, chlorophyll c3; Chl c2, chlorophyll c2; Peri, peridinin; 19'-but, 19'-butanoyloxyfucoxanthin; Fucox, fucoxanthin; Prasinox, prasinoxanthin; Violax, violaxanthin; 19'-hex, 19'-hexanoyloxyfucoxanthin; Diadinox, diadinoxanthin; Allox, alloxanthin; Zeax, zeaxanthin; Chl b, chlorophyll b; Chl a, chlorophyll a.
Figure 1: Average concentration of DFe in each treatment during the incubations measured at T0, T2 and T4. (a) Control and Control+Acid. (b) Fe and Fe+Acid. (c) Dust and Dust+Acid. Error bars indicate standard deviations. n = 3 except for Acid, T0, T2, Dust+Acid, T0, and Control (all times) where n = 2.

Figure 2: Average concentration of chl $a$ (left axis) during the incubations and POC at T4 (right axis) in each treatment. (a) Control and Control+Acid. (b) Fe and Fe+Acid. (c) Dust and Dust+Acid. Error bars indicate standard deviations. Dashed line indicates POC concentration at T0. Chl $a$: n = 3 except for Acid, T0, Dust+Acid, T4, Fe+Acid, T0 and Control (all times) where n=2 because of missing/unreliable data or contamination (Control 1). POC: n=3 except Control where n=2.

Figure 3: Average chl $a$ concentration ($\mu$g L$^{-1}$) attributable to each of the measured groups of phytoplankton initially (T0: white bar) and for each of the treatments after 4 d (T4) of incubation (Control, Control+Acid, Fe, Fe+Acid, Dust, Dust+Acid; gray bars). (a) Diatoms. (b) Dinoflagellates. (c) Haptophytes. (d) Pelagophytes. (e) Chlorophytes. (f) Cyanobacteria. Error bars indicate standard deviations. n = 2 except for T0 (all groups but dinoflagellates) where n = 3 and Acid, diatoms, Control, haptophytes and pelagophytes, Dust+Acid, cyanobacteria and Fe+Acid, haptophytes and cyanobacteria where n = 1.

Figure 4: Average (a) Absolute C assimilation rates. (b) C assimilation rates normalized to chl $a$ concentration at T4 and (c) Fe uptake rates normalized to chl $a$ concentration at T4. Error bars indicate standard deviations. Absolute C assimilation rates: n = 3 except Control and Acid where n = 2, C assim norm to chl $a$: n = 3 except Control, Acid and Dust+Acid where n=2, Fe uptake rates normalized to chl $a$: n = 3 except Control and Dust+Acid where n = 2.

Figure 5: DMSP$_t$ (a, b and c) and DMS (d, e and f) concentrations (nmol L$^{-1}$) in the Control and Control+Acid treatments (a and d), the Fe and Fe+Acid treatments (b and e), and the Dust and Dust+Acid treatments (c and f). Error bars indicate standard deviations. n = 3 except Control where n = 2.