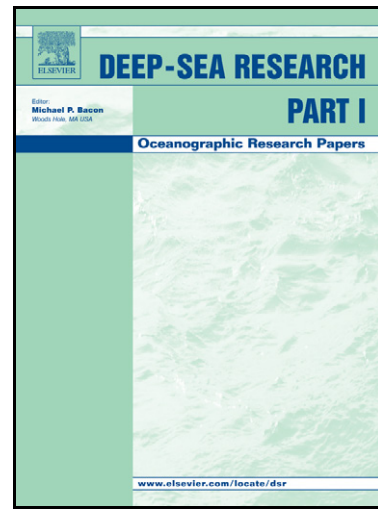


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1 Biogenic silica cycling during summer phytoplankton blooms in the North Pacific subtropical  
2 gyre

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17 **Abstract**

18 Biogenic silica ( $\text{bSiO}_2$ ) cycling, diatom abundance and floristics were examined within  
19 summer-period diatom blooms in the North Pacific Subtropical Gyre (NPSG) in 2008 and 2009.  
20 *Hemiaulus hauckii* was the most abundant diatom observed in an expansive ( $100,000 \text{ km}^2$ )  
21 bloom near the subtropical front in the northeastern NPSG in 2008 and the small pennate diatom  
22 *Mastogloia woodiana* dominated a smaller ( $30,000 \text{ km}^2$ ) bloom sampled in 2009 in the gyre  
23 interior. In both blooms, the  $\text{bSiO}_2$  stock and production rates were up to an order of magnitude  
24 higher relative to non-bloom areas. Remnants of a  $\text{bSiO}_2$  export event was sampled in the *H.*  
25 *hauckii* bloom area where the export rate at 300 m exceeded that at 150 m, and was among the  
26 highest values recorded in the NPSG. The *M. woodiana* bloom was very active with specific  
27  $\text{bSiO}_2$  production rates of  $0.50\text{-}0.75 \text{ d}^{-1}$  and net  $\text{bSiO}_2$  production rates were among the highest  
28 observed in any subtropical-gyre diatom bloom to date. Net silica production rates in the  
29 euphotic zone were strongly positive within blooms and near zero outside of blooms, consistent  
30 with an important role for blooms in  $\text{bSiO}_2$  export. The difference in the areal extent of the *H.*  
31 *hauckii* and *M. woodiana* blooms was consistent with remote-sensing observations that blooms  
32 in the northeastern portion of the NPSG, near the subtropical front, are typically more extensive  
33 than those in the gyre interior near Hawaii Ocean Time-series station ALOHA. Initial estimates  
34 suggest that blooms in the northeast region produced 3 to 25 times more  $\text{bSiO}_2$  in 2008 and 2009,  
35 respectively, than did blooms in the gyre interior; and due to the large areal extent these blooms,  
36 their area-integrated production of  $\text{bSiO}_2$  is similar to intense diatom blooms coastal upwelling  
37 systems (e.g. Monterey Bay, Santa Barbara Channel) despite significantly lower production rates  
38 and standing stock.

39 Keywords: North Pacific subtropical gyre, blooms, diatoms, silica production, net silica  
40 production, export

41 **1. Introduction**

42 Compared to blooms in other subtropical gyres, phytoplankton blooms in the North Pacific  
43 subtropical gyre (NPSG) are different in their timing and in the hydrographic conditions under  
44 which they occur. Winter convective mixing in the NPSG is weak and regular spring blooms do  
45 not ensue; instead, phytoplankton blooms occur during summer when the upper-water column  
46 stratification is at its annual maximum and macronutrient concentrations are at their annual  
47 minimum (Wilson, 2003; Dore *et al.*, 2008; Wilson *et al.*, 2008). This situation sharply contrasts  
48 with that in the North Atlantic Subtropical Gyre (NASG) where strong winter convection  
49 typically erodes the pycnocline and entrains nutrients into the euphotic zone, thereby stimulating  
50 a regular spring bloom after water-column restratification (Sverdrup, 1953). Summer blooms in  
51 the NPSG interior show enhanced abundance of diazotrophs (e.g. *Trichodesmium*) and diatoms  
52 (Dore *et al.*, 2008; Fong *et al.*, 2008; Villareal *et al.*, 2011; 2012); however, *Trichodesmium* is  
53 rarely observed in the northeastern NPSG near the subtropical front (Venrick, 1997; Dore *et al.*,  
54 2008) where satellite ocean color observations reveal blooms that are generally more expansive  
55 and that longer-lived than those in the gyre interior near the Hawaii Ocean Time-series (HOT)  
56 station ALOHA (Wilson, 2003; Wilson *et al.*, 2008). At least some of the blooms occurring in  
57 the northeastern gyre are dominated by diatoms (Brzezinski *et al.*, 1998; Villareal *et al.*, 2011).

58 Why diatom blooms in the NPSG occur during the highly oligotrophic conditions of summer  
59 remains enigmatic. Some blooms contain diatom-diazotrophic associations (DDAs, e.g. the  
60 diatom genera *Hemiaulus* or *Rhizosolenia* and the nitrogen-fixing symbiont, *Richelia*; Villareal  
61 *et al.*, 2011), suggesting that the biological input of fixed N may play a role in bloom initiation  
62 and development. Recently, it was also suggested that summer blooms in the northeastern NPSG  
63 can occur in response to increased vertical mixing due to internal wave breakdown at the 30°N  
64 “critical latitude” (Wilson, 2011). A diatom bloom also requires adequate silicic acid; relative to  
65 the nanomolar levels of inorganic N and P in the surface waters of the NSPG, the silicic-acid  
66 requirement is easily met as surface-water concentrations are typically 0.5 – 1.5 μM. There is  
67 also evidence that the diatom taxa which dominate summer blooms have very efficient Si uptake  
68 kinetics, such that ambient silicic acid concentrations of ~1 μM would support high growth rates  
69 (Brzezinski *et al.*, 1998; Krause *et al.*, 2012).

70 While there currently is no consensus on the factor(s) initiating summer diatom blooms in the  
71 NPSG, there is clear evidence as to their importance in annual biogeochemical budgets. This is

72 somewhat counter intuitive as diatom biomass during non-bloom periods at station ALOHA, as  
73 measured by biogenic silica ( $b\text{SiO}_2$ ) concentration, is the lowest thus far observed in the global  
74 ocean (Brzezinski *et al.*, 2011). However, at the HOT station ALOHA the contribution of  
75 diatoms to new production exceeds both their contribution to autotrophic biomass and their  
76 estimated contribution to primary production (Brzezinski *et al.*, 2011), implying a  
77 disproportionately important role for diatoms in carbon export. This is especially true for  
78 summer blooms which have been estimated to account for 18% of annual new production at  
79 station ALOHA (Dore *et al.*, 2008) and recent estimates suggest that summer blooms account for  
80 29% of the annual production of  $b\text{SiO}_2$  at ALOHA (Brzezinski *et al.*, 2011). Blooms in the  
81 northeastern gyre near the subtropical front are typically longer-lived than those at or near  
82 ALOHA (Wilson *et al.*, 2008), but their role in annual biogeochemical budgets is less clear.  
83 There is also evidence for a significant role for diatoms in the export of carbon to the ocean  
84 interior at ALOHA where observations from both shallow (150 m) and deep ( $\geq 2,800$  m)  
85 sediment traps show an annual summer maximum in carbon export associated with increased  
86 diatom and  $b\text{SiO}_2$  export (Scharek *et al.*, 1999a; b; Karl *et al.*, 2012). To further evaluate the  
87 contribution of summer diatom blooms to silica production and organic matter cycling in the  
88 NPSG, we examined rates of gross and net silica production, diatom abundance and taxonomic  
89 composition within summer phytoplankton blooms in the NPSG.

90

## 91 2. Methods

### 92 2.1 Study area and sampling

93 The Pacific Open-Ocean Bloom (PB) cruises were conducted aboard the R/V *Kilo Moana*  
94 from 4 –19 July 2008 (PB08) and from 29 July – 12 August 2009 (PB09). Using daily-  
95 composite images from the Moderate Resolution Imaging Spectroradiometer (MODIS,  
96 <http://modis.gsfc.nasa.gov>), bloom regions were identified as areas with chlorophyll a (Chl *a*)  
97 concentrations  $\geq 0.12 \mu\text{g L}^{-1}$ . As the feature sampled in 2008 was relatively weak, visualizing  
98 the full extent of the bloom area required the [Chl *a*] threshold value to be slightly lower than the  
99  $0.15 \mu\text{g L}^{-1}$  value used previously (Wilson *et al.*, 2008); for consistency we use the same [Chl *a*]  
100 threshold for both PB cruises. Three regions with bloom-level [Chl *a*] were sampled, one in  
101 2008 and two in 2009. Within each bloom region, stations were classified as representing a  
102 diatom bloom condition by meeting two criteria: 1) the  $>10 \mu\text{m}$  size-fraction [Chl *a*] was  $>30\%$

103 of the total [Chl *a*] collected on a >0.4  $\mu\text{m}$  filter. This threshold value represents a three-fold  
104 increase in >10  $\mu\text{m}$  [Chl *a*] fraction over that present under non-bloom conditions in the gyre  
105 during summer (Villareal *et al.*, 2011), and 2) diatom numerical abundance exceeded 1,000 cells  
106  $\text{L}^{-1}$ , consistent with threshold bloom abundances previously applied in this region (e.g.,  
107 Brzezinski *et al.*, 1998; Villareal *et al.*, 2011).

108 During PB08 a single bloom feature was sampled near the subtropical front northeast of  
109 Hawaii (Fig. 1). The bloom was intensifying as the ship left port in Hawaii and satellite  
110 chlorophyll levels were above the 0.12  $\mu\text{g L}^{-1}$  bloom threshold when the ship arrived at the  
111 bloom, but while the area was being sampled the satellite-observed chlorophyll levels fell below  
112 the bloom threshold. As a result of the declining biomass, only one of the 18 stations sampled in  
113 the area met the diatom-bloom criteria (Fig. 1). Two bloom features were sampled in 2009. In  
114 mid-June of 2009 a high chlorophyll feature developed east of Hawaii at ~26°N, 146°W (Fig. 1),  
115 but remotely-sensed chlorophyll concentrations had dropped significantly by the time of  
116 sampling in mid-July (Fig. 1). However, satellite images revealed another bloom feature  
117 developing to the north of Hawaii at ~25°N, 155°W (Fig. 1). Two stations were sampled within  
118 this feature while it was still active (or developing), and both met the bloom-station criteria. This  
119 bloom will be referred to as the active bloom from PB09 to distinguish it from observations  
120 taken in the faded bloom further to the east.

121 Sampling consisted of CTD casts throughout the day. Pre-dawn casts (04:00-06:00 h, local  
122 time) were done to obtain samples for nutrient concentration, Chl *a* concentration, bSiO<sub>2</sub>  
123 concentrations and bSiO<sub>2</sub> production rates. Samples for diatom abundance and composition  
124 were taken on the rate casts or on the cast immediately before/after. On additional casts during  
125 the day, all samples were collected except those for rate measurements. Due to time constraints  
126 for sampling the second bloom region examined in 2009, rate profiles at PB09 stations 22 and 23  
127 were conducted after 6:00 h local time, but all incubations ran 24 hours to integrate through a  
128 complete photoperiod (see below). For rate measurements water samples were taken at nine  
129 depths within the euphotic zone where the percent of irradiance was determined to be 100, 59,  
130 31, 19, 10, 6, 3.4, 0.6 and 0.1% of that just below the surface (%I<sub>0</sub>). At non-rate stations fixed  
131 sampling depths (surface: i.e. 3 – 5 m, 10, 20, 40, 60, 80, 100, 125, 140, 160 m) were used.  
132 Seawater was collected using 12 L PVC sampling bottles on a rosette equipped with a Seabird  
133 CTD and a photosynthetically active radiation sensor to determine light attenuation. Mixed layer

134 depths were calculated using 1-m binned CTD data based on a  $0.125 \text{ kg m}^{-3}$  change in potential  
135 density from the 0-1 m bin.

136

137 *2.2. Nutrient, taxonomy, and biogenic particulate analysis*

138 Dissolved nutrients were collected and analyzed using standard methods. Unfiltered  
139 seawater samples for  $[\text{Si(OH)}_4]$  determination were refrigerated at  $4^\circ\text{C}$  until analysis at sea using  
140 a sensitive manual colorimetric method (Brzezinski and Nelson, 1995). Nitrate ( $<10 \text{ nM}$ ) and  
141 soluble reactive phosphorus ( $<35 \text{ nM}$ ) were also taken from sampling bottles and analyzed using  
142 both high-sensitivity methods (e.g. Church *et al.*, 2009) and standard colorimetric methods (e.g.  
143 Villareal *et al.*, 2012). The nutrient data are discussed in detail elsewhere (Duhamel *et al.*, 2010;  
144 2011; Villareal *et al.*, 2012).

145 Samples for biomass and diatom-assemblage measurements were analyzed by methods used  
146 previously in subtropical-gyre regions. Seawater for biogenic ( $\text{bSiO}_2$ ) and lithogenic silica  
147 analysis was collected in 2.8 L polycarbonate bottles, filtered through  $0.6 \mu\text{m}$  polycarbonate  
148 filters, and analyzed using the sequential  $\text{NaOH} - \text{HF}$  digestion procedure (Brzezinski and  
149 Nelson, 1995) with reactions carried out in Teflon® tubes which provide low and stable blanks  
150 (Krause *et al.*, 2009). Lithogenic silica concentrations were very low (e.g.  $3 - 8 \text{ nmol Si L}^{-1}$ ) and  
151 are not discussed in detail.  $[\text{Chl } a]$  was determined for two size fractions:  $>0.4 \mu\text{m}$  (e.g., total  
152 phytoplankton community) and  $>10 \mu\text{m}$  (e.g., diatoms, dinoflagellates) by filtering 250 mL and  
153 500 mL, respectively, through polycarbonate membrane filters, extracting in methanol, and  
154 quantifying fluorometrically without acidification (Welschmeyer, 1994). Samples for diatom  
155 abundance and taxonomy were preserved with formalin and enumerated using inverted  
156 microscopy (Villareal *et al.*, 2012).

157

158 *2.3. Measurement of biogenic silica production and export rates*

159 Measurements of gross  $\text{bSiO}_2$  production rates ( $\rho_P$ ) from each sampling depth were  
160 conducted in polycarbonate bottles (300 mL) using the radioisotope  $^{32}\text{Si}$ . Each sample received  
161 360 Bq of high specific activity  $^{32}\text{Si(OH)}_4$  ( $>40 \text{ kBq } \mu\text{mol Si}^{-1}$ ) which had been cleaned of trace  
162 metals by passage through Chelex resin (BioRad). Samples were incubated on the ship deck for  
163 24 hours in acrylic incubators, screened to the relative light level corresponding to each sample  
164 collection depth, and continuously cooled by flowing surface seawater. After incubation, all

silica production rate samples were processed as described in Krause et al. (2012) and  $^{32}\text{Si}$  activity was quantified using gas-flow proportional counting at secular equilibrium of  $^{32}\text{Si}$  and its daughter isotope  $^{32}\text{P}$  (Krause *et al.*, 2011). Specific rates of silica production ( $V_b$ ) were calculated by normalizing  $\rho_p$  to  $[\text{bSiO}_2]$  as in Brzezinski and Phillips (1997). The average  $V_b$  for the upper water column was calculated by vertically integrating  $V_b$  to a specific depth (e.g. 150 m) then dividing by the depth of integration.

The net  $\text{bSiO}_2$  production rate ( $\rho_N$ ) was measured by the net change in biogenic silica concentrations during incubations. During PB08, two 2.8 L polycarbonate bottles were filled from each of the nine light depths sampled on rate casts and processed following the protocol from Demarest et al. (2011). Briefly, one was assigned as the initial and immediately processed, while the other was incubated for 24 hours and then processed. To account for  $\text{bSiO}_2$  that adsorbed to the bottle wall (Krause *et al.*, 2010b) each sample bottle was rinsed with three aliquots of 0.2- $\mu\text{m}$ -filtered seawater and the rinses passed through the same filter as the main sample. Because of the low  $[\text{bSiO}_2]$  and the small net changes observed during PB08, the protocol of Krause et al. (2010b) was used during PB09, where four replicates were run as initial samples and another four incubated as final samples. The increased processing time limited experiments to four depths (59, 31, 10, and 3.4%  $I_0$ ). All samples were analyzed for both  $\text{bSiO}_2$  and lithogenic silica, using NaOH – HF serial digestion described above, and corrected for bias assuming that 10% of the lithogenic silica dissolves during the NaOH digestion (Ragueneau and Treguer, 1994).

The export of  $\text{bSiO}_2$  ( $\rho_E$ ) at 150 m and 300 m was measured on PB08 using a free-drifting Multitrap sediment trap array (e.g. Karl *et al.*, 1996). During the PB09 cruise the combined factors of the passage of hurricane Felicia and occupying an active bloom area for <24 hours, meant that sediment traps could not be deployed. On PB08, collector tubes fitted with entrance baffles were filled with brine (50 g L<sup>-1</sup> NaCl in 0.2  $\mu\text{m}$  filtered surface seawater) and formalin preservative was added to half the tubes. Three tubes of each type were deployed at 150 m and 300 m for approximately 5 days inside and outside of the bloom region. Samples were taken to measure the amount of  $\text{bSiO}_2$  which dissolved during deployment (~3% of particulate flux), as described in Brzezinski et al. (2011), and used to correct the total measured  $\text{bSiO}_2$  flux. Dissolution was not measured in the preserved traps, but there was no significant difference

195 between the particulate flux in preserved and non-preserved traps deployed at the same depth and  
 196 time (t-test, p values ranged 0.27 – 0.91 for pairwise comparisons).

197

### 198 **3. Results**

#### 199 *3.1. Hydrography and nutrients*

200 The hydrography observed during each cruise is described briefly for context; however, a  
 201 more thorough description can be found in Villareal *et al.* (2012). During PB08, MODIS  
 202 imagery revealed a bloom feature developing to the northeast of Hawaii, near the subtropical  
 203 front. CTD casts in the bloom region revealed significant small-scale salinity variations in  
 204 vertical profiles (Fig. 2A, see also Villareal *et al.*, 2012) that are characteristic of the front  
 205 (Shcherbina *et al.*, 2009, 2010). Isotherms in the upper 100 m showed little vertical variability at  
 206 most stations on both transects, but some shoaling of the isothermal surfaces was observed in the  
 207 lower-salinity stations presumably associated with the front (Fig. 2A). The mixed-layer depth in  
 208 the bloom region averaged  $21 \pm 7$  m (SD).

209 Prior to PB09, MODIS satellite data revealed a bloom feature near  $26^{\circ}\text{N}$  and  $145^{\circ}\text{W}$ , which  
 210 began developing in mid-June. By the time of our occupation, the satellite ocean color signal  
 211 faded considerably and in situ measurements indicated that the bloom had collapsed. During a  
 212 zonal transect across the former bloom region the water column had less temperature and salinity  
 213 variability (Fig. 3A) than was observed during the PB08 transects, and the mixed layer was  
 214  $40 \pm 13$  m. A smaller and higher [Chl *a*] bloom was observed to develop closer to Hawaii, and  
 215 reached the threshold bloom [Chl *a*] value by 7 August. Two rate stations (Sta. 22, 23) were  
 216 occupied within this active bloom on 12 August, and satellite imagery showed it collapsed by 24  
 217 August. The physical conditions were similar to those observed in the collapsed-bloom region  
 218 sampled during PB09, but the mixed layer depth was deeper, 48 – 50 m.

219 Consistent with gyre conditions, the concentrations of inorganic nutrients were very low at  
 220 all stations.  $[\text{Si(OH)}_4]$  in the surface waters was consistently one or two orders of magnitude  
 221 higher than the high-sensitivity analysis concentrations for nitrate ( $<0.01\text{ }\mu\text{M}$ ) and soluble  
 222 reactive phosphate ( $<0.035\text{ }\mu\text{M}$ ). On PB08,  $[\text{Si(OH)}_4]$  was generally  $< 2\text{ }\mu\text{M}$  in the upper 50 m  
 223 and the silicicline shoaled towards the north (Fig. 2C). At the single PB08 bloom station, the  
 224 average mixed layer  $[\text{Si(OH)}_4]$  showed a depletion to  $1.22\text{ }\mu\text{M}$  and the vertical gradient in silicic  
 225 acid from the base of the mixed layer to the 1% light depth was similar to that at the northern-

226 transect stations (Fig. 4A). Mixed-layer  $[Si(OH)_4]$  at non-bloom stations was  $1.69 \pm 0.20 \mu M$ ,  
 227 implying a biological drawdown of  $\sim 0.5 \mu M$   $[Si(OH)_4]$  in the bloom.

228 Dissolved silicate was consistently lower during the PB09 cruise (Fig. 3C).  $[Si(OH)_4]$  in the  
 229 upper 50 m on the collapsed-bloom transect was  $< 1.4 \mu M$ , and surface  $[Si(OH)_4]$  was  $< 1 \mu M$  at  
 230 some stations. In the active bloom, the average mixed-layer  $[Si(OH)_4]$  was  $0.87 \mu M$ ,  $\sim 0.3 \mu M$   
 231 less than that at non-bloom stations (Table 1). The vertical structure in  $[Si(OH)_4]$  below the  
 232 mixed layer was similar at all stations during PB09 (bloom and non-bloom, Fig. 3C).

233

### 234 *3.2. Particle concentrations and diatom abundance*

235 Proxies for phytoplankton biomass showed expected low-levels outside of blooms and  
 236 enhancement within blooms. We describe water-column  $[Chl\ a]$  (data not shown) for context  
 237 and report integrated concentrations (Table 1), for a more thorough analysis of the  $[Chl\ a]$   
 238 dataset see Villareal et al. (2012). Total  $[Chl\ a]$  (i.e.  $> 0.4 \mu m$  size fraction) was typically  $0.15 -$   
 239  $0.25 \mu g\ L^{-1}$  in the deep chlorophyll maxima ( $\sim 80 - 120\ m$ ) during both years. Higher values in  
 240 the overlying waters were observed in PB08 ( $0.08 - 0.10 \mu g\ L^{-1}$ ) vs. PB09 ( $< 0.08 \mu g\ L^{-1}$ ). At the  
 241 PB08 bloom station, the total  $[Chl\ a]$  showed relatively minor enhancement over concentrations  
 242 at non-bloom stations.  $[Chl\ a]$  was  $> 0.39 \mu g\ L^{-1}$  in the active PB09 bloom, with exceptionally  
 243 high values of  $0.54$  and  $1.03 \mu g\ L^{-1}$  observed between  $45$  and  $55\ m$  at stations  $22$  and  $23$ ,  
 244 respectively. On average, the  $> 10 \mu m$  fraction was  $\leq 10\%$  of the total  $[Chl\ a]$  in non-bloom  
 245 stations during both years. However, this larger size fraction represented  $30 - 40\%$  of the total  
 246  $[Chl\ a]$  for PB08 bloom and  $80 - 99\%$  at station  $23$  in the active PB09 bloom.

247 Biogenic silica concentrations were consistent with previous studies in this region.  $[bSiO_2]$   
 248 at non-bloom stations during PB08 were  $20 - 40\ nmol\ Si\ L^{-1}$  in the upper  $50\ m$  (Fig. 2D).  
 249  $[bSiO_2]$  maxima, up to  $100\ nmol\ Si\ L^{-1}$ , were observed between  $25 - 50\ m$  at some stations on  
 250 both transects. At the PB08 bloom station, the  $[bSiO_2]$  surpassed  $200\ nmol\ Si\ L^{-1}$  in the surface  
 251 (Fig. 4B).  $[bSiO_2]$  along the PB09 transect of the faded bloom were between  $10$  and  $20\ nmol\ Si\ L^{-1}$   
 252 with little vertical structure in the euphotic zone (Fig. 3D). But in the active PB09 bloom, the  
 253  $[bSiO_2]$  was an order of magnitude higher than at non-bloom stations, with subsurface  
 254 maximums of  $153$  and  $240\ nmol\ Si\ L^{-1}$  at stations  $22$  and  $23$ , respectively (Fig. 4B).

255 Vertical integrals of biomass measurements reveal large differences between bloom and non-  
 256 bloom stations during the PB cruises. During PB08  $\int Chl\ a$  within the  $> 10 \mu m$  size fraction nearly

doubled at the single bloom station ( $2.5 \text{ mg m}^{-2}$ ), while [ $\text{Chl } a$ ] in the  $>0.4 \mu\text{m}$  size fraction at the bloom station ( $18.3 \text{ mg m}^{-2}$ ) was similar to the non-bloom station average ( $18.8 \text{ mg m}^{-2}$ , Table 1). During PB09, [ $\text{Chl } a$ ] in the  $>10 \mu\text{m}$  size fraction was  $1.1 \text{ mg m}^{-2}$  at the non-bloom stations (includes profiles in the faded bloom) and significantly lower than the  $>0.4 \mu\text{m}$  size fraction value ( $16.1 \text{ mg m}^{-2}$ , Table 1). But within the active PB09 bloom, substantial increases were observed in both size fractions; the  $>10 \mu\text{m}$  and  $>0.4 \mu\text{m}$  fractions averaged  $18.3 \text{ mg m}^{-2}$  and  $53.5 \text{ mg m}^{-2}$ , respectively (Table 1). Vertical integrals of [ $\text{bSiO}_2$ ] showed relative changes similar to those of  $>10 \mu\text{m}$  [ $\text{Chl } a$ ], with significant differences between bloom and non-bloom stations. During PB08, the non-bloom  $\int \text{bSiO}_2$  averaged 40% of the bloom station value ( $13.3 \text{ mmol Si m}^{-2}$ , Table 1). Similarly, the non-bloom  $\int \text{bSiO}_2$  in PB09 was only 26% of the  $\int \text{bSiO}_2$  at the two active bloom stations during PB09 ( $7.7 \text{ mmol Si m}^{-2}$ , Table 1).

Diatom abundance and species dominance differed between cruises. During PB08, *Hemiaulus hauckii* was the numerically dominant diatom at all stations (Table 1). The vertical structure in diatom abundance (Fig. 2B, 3B, 4F) was similar to that for [ $\text{bSiO}_2$ ] (Figs. 2D, 3D, 4B) and  $>10 \mu\text{m}$  [ $\text{Chl } a$ ] (Fig. 4C). Surface maximums of  $\sim 250 \text{ cells L}^{-1}$  were observed during the PB08 transects, with abundances  $<50 \text{ cells L}^{-1}$  persisting deeper than 50 m (Fig. 2B). Diatom abundance increased to  $>10,000 \text{ cells L}^{-1}$  in the surface at the bloom station, with  $>100 \text{ cells L}^{-1}$  at depths shallower than 60 m (Fig. 4F). The integrated diatom abundance within the mixed-layer varied by an order of magnitude between non-bloom ( $5.9 \times 10^6 \text{ cells m}^{-2}$ ) and bloom stations ( $8.2 \times 10^7 \text{ cells m}^{-2}$ , Table 1). *M. woodiana* dominated abundance at 11 of the 14 stations where abundance data was obtained during PB09, with *H. hauckii* and *Chaetoceros* spp. being most abundant diatoms at the other stations (Table 1). Non-bloom station cell abundance was very low, e.g.  $<50 \text{ cells L}^{-1}$ , with little vertical structure (Fig. 3B). Within the active bloom, *M. woodiana* accounted for 98% of total diatom abundance, despite maximum numerical abundances for all diatom genera being observed in this bloom (Table 1). Maximum diatom cell abundances in the station-22 mixed layer were  $38,000 \text{ cells L}^{-1}$ , and maximum mixed-layer abundances were higher at station 23 ( $159,000 \text{ cells L}^{-1}$ , Fig. 4F). The mixed-layer integrated diatom abundance at non-bloom stations during PB09 ( $1.1 \times 10^6 \text{ cells m}^{-2}$ ) was three orders of magnitude lower than at the active bloom stations ( $1.4 \times 10^9 \text{ cells m}^{-2}$ ; Table 1).

286

287 *3.3. Biogenic silica production and export rates*

288 Gross biogenic silica production rates ( $\rho_P$ ) varied by an order of magnitude between bloom  
 289 and non-bloom stations. Surface  $\rho_P$  ranged between  $2 - 5 \text{ nmol Si L}^{-1} \text{ d}^{-1}$  at non-bloom stations  
 290 during PB08 (Fig. 2E) and increased to  $>20 \text{ nmol Si L}^{-1} \text{ d}^{-1}$  at the bloom station (Fig. 4D).  
 291 Vertical trends in  $\rho_P$  below 50 m were similar in and out of the bloom area, with rates declining  
 292 from  $<1 \text{ nmol Si L}^{-1} \text{ d}^{-1}$  at 50 m to analytically zero at 175 m, indicating that the full vertical  
 293 extent of  $\rho_P$  in the water column was sampled (Fig. 2E). At PB09 non-bloom stations, only two  
 294 samples (out of 59) had rates  $>2 \text{ nmol Si L}^{-1} \text{ d}^{-1}$ . Rates in the upper 50 m were generally  $1.2 -$   
 295  $1.8 \text{ nmol Si L}^{-1} \text{ d}^{-1}$  and below 50 m, deep  $\rho_P$  declined to rates similar to rates observed in PB08  
 296 (Fig. 3E). Within the active PB09 bloom,  $\rho_P$  values were  $>15 \text{ nmol Si L}^{-1} \text{ d}^{-1}$  throughout the  
 297 upper 50 m, with a subsurface maximum of  $54 \text{ nmol Si L}^{-1} \text{ d}^{-1}$  (Fig. 4D). Deeper than 60 m,  $\rho_P$   
 298 at the active bloom stations were nearly identical to those rates measured at non-bloom stations.  
 299 Vertical integrals of  $\rho_P$  ( $\int \rho_P$ ) demonstrate the strong disparity between bloom and non-bloom  
 300 rates. Non-bloom station average  $\int \rho_P$  during PB08 was 32% of the bloom-station rate ( $0.65$   
 301  $\text{mmol Si m}^{-2} \text{ d}^{-1}$ , Table 1). A larger disparity was observed in PB09, as the non-bloom average  
 302  $\int \rho_P$  was only 9% of the active bloom rate ( $1.33 \text{ mmol Si m}^{-2} \text{ d}^{-1}$ ).

303 As with  $\rho_P$ ,  $[\text{bSiO}_2]$ , and diatom abundance, specific uptake rate  $V_b$  was generally highest in  
 304 the upper water column and declined to analytical zero values at the  $0.1\%I_0$  isolume (Fig. 2F, 3F,  
 305 4E). During PB08,  $V_b$  in the upper 15 m were  $\sim 0.05 - 0.08 \text{ d}^{-1}$  at non-bloom stations (Fig. 2F),  
 306 but at the bloom station  $V_b$  in the upper 15 m was  $\sim 0.11 \text{ d}^{-1}$  (Fig. 4E), implying doubling times of  
 307  $\sim 6$  and  $\sim 10$  days inside and outside bloom, respectively. Vertically-weighted average  $V_b$  within  
 308 the upper 150 m showed no difference between the non-bloom ( $0.05 \text{ d}^{-1}$ ) and bloom ( $0.04 \text{ d}^{-1}$ )  
 309 rates (Table 1), indicating an average doubling time for  $\text{bSiO}_2$  in the upper 150 m of  
 310 approximately two weeks during the PB08 cruise.  $V_b$  on the faded-bloom transect during PB09  
 311 was  $>0.05 \text{ d}^{-1}$  in the upper 50 m at two stations and values  $\geq 0.05 \text{ d}^{-1}$  throughout the upper 150 m  
 312 at the easternmost station (Fig. 3F). At the active-bloom stations during PB09,  $V_b$  averaged  
 313  $\sim 0.35 \text{ d}^{-1}$  in the upper 10 m with a maximum of  $0.75 \text{ d}^{-1}$  at the base of the mixed layer ( $\sim 50$  m,  
 314 Fig. 4E); the latter infers a doubling time of  $<1$  day. For PB09, the vertically-averaged  $V_b$  within  
 315 the active bloom ( $0.17 \text{ d}^{-1}$ ) was nearly three-fold higher than for non-bloom stations ( $0.06 \text{ d}^{-1}$ ),  
 316 implying doubling times of 4 and 11 days for bloom and non-bloom stations, respectively.

317 The net production of biogenic silica ( $\rho_N$ ) is defined as the difference between  $\rho_P$  and gross  
 318 silica dissolution, and thereby represents an upper limit for bSiO<sub>2</sub> export. Because  $\rho_N$  was not  
 319 sampled at every rate station during PB08, we do not have a measurement at the single bloom  
 320 station. When integrated within the mixed layer,  $\int \rho_N$  was positive at three stations (Fig. 5).  
 321 Integration to deeper depths (e.g. 3.4%I<sub>0</sub> and 0.1%I<sub>0</sub>) caused  $\int \rho_N$  to become negative, indicating  
 322 that within the euphotic zone the dissolution of bSiO<sub>2</sub> exceeded its rate of production. During  
 323 PB09, replication allowed for constraints on the variability in the estimates of  $\int \rho_N$  (Fig. 5B).  
 324 Nearly all stations showed positive  $\int \rho_N$  over the mixed layer and often between the surface and  
 325 the 3.4%I<sub>0</sub> isolume (Fig. 5), but only in the active bloom (stations 22, 23, Fig. 5) was  $\int \rho_N$   
 326 statistically different from zero (i.e. production of bSiO<sub>2</sub> exceeded its loss from dissolution) for  
 327 the mixed layer and to the 3.4%I<sub>0</sub> isolume integrals. Considering the variability among  
 328 replicated samples during PB09,  $\int \rho_N$  measured during PB08 were unlikely to be statistically  
 329 different from zero. These results suggest that the outside of bloom stations, silica production  
 330 and dissolution were approximately in balance leading to no or very little net silica production  
 331 (i.e.  $\int \rho_N \approx 0$ ) within the mixed layer.

332 Due to inclement weather, no sediment trap deployments were made during PB09; however,  
 333 we can make comparisons between bloom and non-bloom regions for bSiO<sub>2</sub> flux ( $\rho_E$ ) during  
 334 PB08. Samples with and without formalin showed no statistically significant difference in the  
 335 measured  $\rho_E$  (t-test, p values ranged 0.27 – 0.91), and we report the pooled mean  $\pm$ SD from both  
 336 trap types.  $\rho_E$  at 150 m within the bloom area was higher than that observed outside the bloom  
 337 ( $0.27 \pm 0.05$  mmol Si m<sup>-2</sup> d<sup>-1</sup> vs.  $0.17 \pm 0.03$  mmol Si m<sup>-2</sup> d<sup>-1</sup>). Unexpectedly,  $\rho_E$  at 300 m in the  
 338 bloom area was  $0.36 \pm 0.03$  mmol Si m<sup>-2</sup> d<sup>-1</sup>, ~33% higher than  $\rho_E$  at 150 m in the bloom, and  
 339 was four-fold higher than 300-m  $\rho_E$  outside the bloom ( $0.09 \pm 0.01$  mmol Si m<sup>-2</sup> d<sup>-1</sup>).  
 340

#### 341 4. Discussion

##### 343 4.1. Siliceous biomass and silica production during summer in the NPSG

344 The blooms sampled during the PB cruises are similar to summer-period stocks and rates  
 345 observed previously in this region. On zonal transects along ~25°N and 31°N in NPSG during  
 346 summer, Brzezinski et al. (1998) observed a  $\int b\text{SiO}_2$  range of 2 – 18 mmol Si m<sup>-2</sup> (Fig. 6A), but

347 by our diatom-bloom criteria only one bloom station was sampled during their cruises. Between  
 348 1997 and 2009, summer blooms occurred most years at station ALOHA, elevating  $\int bSiO_2$  by  
 349 ~50% above the non-summer average (2.5 mmol Si m<sup>-2</sup>; Brzezinski *et al.*, 2011). The PB08  
 350 bloom-station  $\int bSiO_2$  (Table 1) was similar to the highest values observed in all 12 years of  
 351  $bSiO_2$  data at station ALOHA, but about 33% lower than the highest integrals reported by  
 352 Brzezinski *et al.* (1998) (Fig. 6A). Similarly, the  $\int bSiO_2$  at the PB09 active bloom stations (Table  
 353 1) was comparable to the highest values observed in the ALOHA record (Fig. 6A). Non-bloom  
 354 station  $\int bSiO_2$  during the PB cruises (Table 1) was also similar to values during previous studies  
 355 in this region (Fig. 6A). Combining our data with the previous data at station ALOHA and from  
 356 transects by Brzezinski *et al.* (1998) shows that the magnitude of  $\int bSiO_2$  during the summer  
 357 season does not appear to increase from the gyre interior (near ALOHA) to the northeastern  
 358 boundary (near the subtropical front).

359 Reports of biogenic silica production rates in the NPSG are limited. Brzezinski *et al.* (1998)  
 360 reported  $\int \rho_P$  between 0.5 and 2.9 mmol Si m<sup>-2</sup> d<sup>-1</sup> (average 1.2 mmol Si m<sup>-2</sup> d<sup>-1</sup>) during their  
 361 summer cruises in the mid-1990s (Fig. 6B). This range and mean were significantly higher than  
 362 that observed recently at station ALOHA, where non-summer  $\int \rho_P$  ranged between 0.1 – 0.2  
 363 mmol Si m<sup>-2</sup> d<sup>-1</sup> and summer values averaged 0.4 mmol Si m<sup>-2</sup> d<sup>-1</sup> (Fig. 6B; Brzezinski *et al.*,  
 364 2011).  $\int \rho_P$  at the three PB bloom stations was higher than at ALOHA during the summer  
 365 (Brzezinski *et al.*, 2011) and they were similar to the mean rate reported for all stations by  
 366 Brzezinski *et al.* (1998) (Fig. 6B). One other report of biogenic silica production rates in this  
 367 region was for *Rhizosolenia* mats during the summer months (Shipe *et al.*, 1999). Shipe *et al.*  
 368 (1999) estimated that *Rhizosolenia* mat production was up to ~0.3 mmol Si m<sup>-2</sup> d<sup>-1</sup> in the upper  
 369 150 m. This rate surpasses nearly all non-bloom stations during the PB cruises (Table 1) and  
 370 reinforces Shipe *et al.*'s (1999) conclusions that *Rhizosolenia* mats are an important overlooked  
 371 source of silica production in this region.

372

#### 373 4.2. Comparison to other open-ocean diatom blooms

374 Even under bloom conditions, the diatom biomass in the NSPG is lower than coastal zones  
 375 (Brzezinski *et al.*, 1997; 2003; Shipe and Brzezinski, 2001) or the Southern Ocean/Antarctica  
 376 (Nelson *et al.*, 1991) and this results in significantly lower rates of  $bSiO_2$  production (Table 2).  
 377 Consequently, NSPG gross rates are low and similar in magnitude to other subtropical gyres.

378 However, the elevated rates observed in the PB blooms are comparable to those observed in  
 379 subtropical-gyre mesoscale features, where intensified vertical nutrient supply enhances  
 380 phytoplankton rate processes (e.g. McGillicuddy *et al.* 1998). NPSG blooms are distinct for  
 381 being spatially larger (Table 2) and for occurring without significant enhancement of  
 382 macronutrients by obvious mesoscale processes. In the Sargasso Sea, Krause *et al.* (2010b)  
 383 observed greatly enhanced  $[b\text{SiO}_2]$ ,  $\int\rho_P$  and  $\int\rho_N$  in a mode-water eddy (MWE), the same eddy  
 384 type harboring the enhanced diatom community described by McGillicuddy *et al.* (2007). While  
 385  $\int b\text{SiO}_2$  was higher in the MWE than at all PB stations,  $\int\rho_P$  was similar and the total amount of  
 386  $b\text{SiO}_2$  produced in this MWE may have been similar to that observed in the PB09 bloom (Table  
 387 2). But  $\int\rho_N$  was nearly 3-fold higher in the PB09 bloom compared to the MWE. Eddies have  
 388 also been observed to have enhanced diatom biomass relative to normal (e.g. non-bloom)  
 389 conditions in the NPSG (Benitez-Nelson *et al.*, 2007; Fong *et al.*, 2008), but direct comparison  
 390 with the PB blooms is problematic since no measurements of  $[b\text{SiO}_2]$  and silica production were  
 391 made during these studies. However, Benitez-Nelson *et al.* (2007) did report a 150-m  $\rho_E$  of 0.43  
 392 mmol Si m<sup>-2</sup> d<sup>-1</sup> in an NPSG cyclonic eddy (i.e. “Cyclone Opal”), this value is only 20% higher  
 393 than the 300-m  $\rho_E$  measured in PB08 bloom region (Fig. 6C).

394 Compared to other diatom blooms examined to date, the PB09 bloom appeared to have lower  
 395 proportional losses of biogenic silica due to silica dissolution. The ratio of net  $b\text{SiO}_2$   
 396 accumulation relative to the gross rate ( $\int\rho_N \div \int\rho_P$ ) is equivalent to the statistic  $1 - D:P$ , a  
 397 diagnostic of fraction of “new” silica production (Brzezinski *et al.*, 2003) ( $D:P$  is the ratio of  
 398 gross silica dissolution to gross silica production). In the PB09, the  $1 - D:P$  in the active bloom  
 399 was 0.77, this is higher than other blooms in subtropical gyres (e.g. 0.66; Krause *et al.*, 2010b)  
 400 and is similar to bloom values observed in high-diatom biomass systems including the Gulf  
 401 Stream Warm Core Rings, Antarctic Polar Front, and the Amazon River Plume (*see fig. 4 in*  
 402 Brzezinski *et al.* 2003). Thus, in the active PB09 bloom, the much lower proportional loss of  
 403  $b\text{SiO}_2$  due to dissolution allows for a higher fraction of  $\rho_P$  to result in  $\int b\text{SiO}_2$  accumulation and  
 404 potentially export.

405

#### 406 4.3. Bloom drivers

407 The summer blooms in the NPSG have been shown to be quantitatively significant in terms  
 408 of their effect on the annual export of particulate organic matter to the ocean interior (Scharek *et*

409 *al.*, 1999b; Dore *et al.*, 2008; Karl *et al.*, 2012); however, a consensus view as to what is driving  
410 blooms during this period has yet to be realized. A companion study suggested that Si was not  
411 limiting to diatom growth during these blooms (Krause *et al.* 2012); therefore, input of Si can be  
412 eliminated as a potential bloom driver. The increase in available N by enhanced nitrogen  
413 fixation has been suggested to be a mechanism which facilitates blooms in the NPSG (e.g.  
414 Wilson, 2003; Dore *et al.*, 2008); such that blooms of diazotrophs or DDAs would be regulated  
415 more by nutrients such as phosphorus and iron (e.g. Calil *et al.*, 2011). This did not seem to be  
416 the case during PB08. A study of phosphorus cycling by Duhamel *et al.*, (2010) found little  
417 evidence of P limitation during PB08 leading to the suggestion that the autotrophs present were  
418 mainly limited by N, thus giving direct advantage to organisms capable of N<sub>2</sub> fixation. This is  
419 consistent with the finding that *H. hauckii*, a DDA, dominated the PB08 diatom silica biomass  
420 and silica production rates (Krause *et al.*, 2012). However, N<sub>2</sub> fixation rates measured on the  
421 cruise were quite low. Watkins-Brandt *et al.* (2011) report the highest N<sub>2</sub> fixation rates on this  
422 cruise to be ~2 nmol N L<sup>-1</sup> d<sup>-1</sup>, with most values being ≤1 nmol N L<sup>-1</sup> d<sup>-1</sup>. Considering an average  
423 diatom Si:N of 1:1 (Brzezinski, 1985) and the maximizing assumption that all measured N<sub>2</sub>  
424 fixation was solely by DDAs, then the measured N<sub>2</sub> fixation rates by Watkins-Brandt *et al.*  
425 (2011) could potentially support most silica production at non-bloom stations, but <10% of silica  
426 production rates measured at the PB08 bloom station.

427 The dominant diatom in the active bloom on PB09, *M. woodiana*, is not a DDA and thus  
428 does not benefit directly from N<sub>2</sub> fixation. *M. woodiana* has been observed on multiple  
429 occasions to be a numerically dominant diatom in NPSG blooms (see table 1 in Dore *et al.* 2008)  
430 despite its small size, it also has been observed to be high abundance in local sediment traps  
431 (Scharek *et al.*, 1999a; b), possibly because of its ability to form large aggregates (Villareal *et*  
432 *al.*, 2012). Both *M. woodiana* and DDAs are part of the same near-surface flora (Venrick, 1988),  
433 suggesting that *M. woodiana* benefits from an indirect linkage with coexisting DDAs where food  
434 web processes, or direct N excretion, transfer N from DDAs to non-nitrogen fixers, but the role  
435 of such processes in bloom initiation and/or development remains unclear (Villareal *et al.*, 2012).

436 Other mechanisms that may drive blooms are physical, including mesoscale features  
437 (Church *et al.*, 2009) and fronts (Calil and Richards, 2010). In a synthesis of four blooms  
438 sampled in the northeastern sector of the NPSG (including the two blooms reported here),  
439 Wilson *et al.* (in review) suggest that bloom development requires a subsurface stratification

440 minimum intersecting the nutricline and that this is close to the base of the mixed layer.  
441 However, this assumes that biological productivity in blooms is driven by N<sub>2</sub> fixation (i.e.  
442 solitary/colonial diazotrophs or DDA) making biological production limited by P availability.  
443 Wilson (2011) recently presented a hypothesis to explain blooms occurring to the northeast of  
444 Hawaii near the subtropical front where internal waves, generated at the Hawaiian islands,  
445 propagate to the northeast and break near a “critical latitude” (e.g. 30° N), thereby enhancing  
446 vertical mixing across the nutricline. The PB08 bloom was located very near the “critical  
447 latitude.”

448

#### 449 *4.4. Bloom dynamics*

450 Multiple lines of evidence suggest the PB08 bloom region was sampled while in a declining  
451 state. The MODIS satellite data showed that surface [Chl *a*] had started declining in this feature  
452 prior to our sampling. The bloom had [Chl *a*] above the 0.12 µg L<sup>-1</sup> threshold through at least 8  
453 July 2008, although due to cloud coverage much of the feature was obscured from MODIS  
454 during its decline. The bloom station was sampled on 9 July; thus, most of our stations were  
455 sampled after the bloom was below the ocean-color bloom threshold. Evidence of bloom decline  
456 was also apparent by a lack of significant differences in V<sub>b</sub> between the PB08 bloom and those at  
457 some non-bloom stations (Fig. 2, 3, Table 1), indicating low diatom activity in the declining  
458 bloom. Lastly, there was a lack of vertical attenuation of ρ<sub>E</sub> in the bloom. The export at 300 m  
459 was 33% higher than that at 150 m, whereas outside the bloom the bSiO<sub>2</sub> flux at 300 m was half  
460 of the flux at 150 m. Given the temporal lag for particles exported at 150 m to reach 300 m, the  
461 increase in flux with depth suggests that a large pulse of silica export occurred prior to our  
462 occupation.

463 In contrast to the PB08 bloom, nearly every measurement taken supports that the active PB09  
464 bloom was sampled in a state of rapid autotrophic-biomass accumulation. The MODIS data  
465 indicates that stations 22 and 23 were sampled at the peak biomass achieved by this bloom  
466 (Villareal *et al.*, 2012). The ρ<sub>P</sub> and V<sub>b</sub> observed in this bloom were the highest of all stations  
467 during both PB cruises and were comparable to the highest rates observed at station ALOHA  
468 during two years of study (2008 – 2009; Brzezinski *et al.*, 2011). The high specific production  
469 rates imply doubling-time estimates within the PB09 active bloom to be <1 – 2 days in the upper  
470 55 m, relative to the 10 – 14 day doubling times elsewhere. This evidence suggests a highly

471 dynamic diatom community, despite the predominant condition of very low  $[NO_3]$ ,  $[SRP]$  (i.e. no  
 472 different than non-bloom stations), which was still in a state of positive net  $bSiO_2$  production  
 473 during our sampling, notwithstanding the even lower  $[Si(OH)_4]$  than non-bloom station  
 474 (presumably from biological drawdown).

475 A distinction among the three bloom stations sampled on the PB cruises is the shift in  
 476 numerical dominance by different diatom genera among blooms and their implied contribution to  
 477 silicon dynamics. Recently, we used a biovolume scaling approach (sensu Conley *et al.*, 1989)  
 478 to determine the Si quota of the dominant diatoms in the PB blooms to estimate which diatom  
 479 species likely dominated  $\rho_P$  and the community  $V_b$  (Krause *et al.*, 2012). We observed that *H.*  
 480 *hauckii* and *M. woodiana* dominated the PB08 and active PB09 blooms, respectively, and they  
 481 likely dominated  $\rho_P$  due to the combined effects of each group having the highest contribution to  
 482  $[bSiO_2]$  and highest net growth rates, compared to the other diatoms present. Thus, in these  
 483 blooms it appears that numerical dominance by a particular diatom group also resulted in those  
 484 groups having the most significant role in local silicon biogeochemistry. Such a condition is not  
 485 always true among diatom assemblages, as a recent analysis in the eastern equatorial Pacific  
 486 suggested that up to 40% of the  $\int\rho_P$  in the euphotic zone was done by diatoms representing <10%  
 487 of the total abundance (Krause *et al.*, 2010a).

488 The export of biogenic silica in the PB08 bloom was significant in the context of the regional  
 489 dataset. Biogenic silica flux at 150 m has been regularly sampled by the HOT program since  
 490 1997 (Brzezinski *et al.*, 2011) where  $\rho_E$  is  $\sim 0.07$  mmol Si  $m^{-2} d^{-1}$  and  $\sim 0.13$  mmol Si  $m^{-2} d^{-1}$   
 491 during non-summer and summer months, respectively (Fig. 6C). During the PB08 cruise,  $\rho_E$   
 492 within and outside of the bloom region were both enhanced, relative to average rates observed at  
 493 station ALOHA (Fig. 6C). The 300-m  $\rho_E$  of  $0.36$  mmol Si  $m^{-2} d^{-1}$  within the PB08 bloom area is  
 494 higher than all but two of the 87 cruise values reported for 150 m  $\rho_E$  at ALOHA since 2001  
 495 (Brzezinski *et al.*, 2011) and was 85% of the 150-m  $\rho_E$  observed at in a regional cyclonic eddy  
 496 (“Cyclone Opal”) which harbored significantly enhanced diatom biomass (Benitez-Nelson *et al.*,  
 497 2007). Under steady-state conditions, the  $\int\rho_N$  measurement sets an upper limit for export. At  
 498 most stations, positive  $\int\rho_N$  was observed in the mixed layer, with increasingly negative values for  
 499 deeper integral depths (Fig. 5). Such biomass accumulation, and export potential, in the mixed  
 500 layer, suggests that diatoms in the NPSG may operate in reverse of the classical two-layer model

501 (Dugdale, 1967; Eppley *et al.*, 1973), where export originates in the deeper layer from  
 502 phytoplankton having adequate nutrients but are limited by light.

503 During the PB cruises we can adequately resolve differences in net production between the  
 504 bloom and non-bloom stations but the method has high uncertainty due to natural variability in  
 505 bSiO<sub>2</sub> replicates. At station ALOHA, an average of 54% of total integrated silica production is  
 506 available for export (i.e. net production) (Brzezinski *et al.*, 2011); however, the absolute export  
 507 rates were low (see previous paragraph). With the high uncertainty in the  $\int \rho_N$  measurement, such  
 508 a small positive signal in  $\int \rho_N$  (inferred by the station ALOHA dataset) at non-bloom stations  
 509 cannot be resolved. Only in the active PB09 bloom was  $\int \rho_N$  positive and statistically different  
 510 from zero, suggesting that during this bloom there was an increased decoupling between  
 511 production and dissolution of silica, relative to that observed under non-bloom conditions, this  
 512 decoupling lead to higher net accumulation of biomass and high export potential. The  
 513 observation that *M. woodiana* formed visible aggregates in this bloom (Villareal *et al.*, 2012)  
 514 provides a mechanism to enhance the settling rate of these diatoms, relative to the rate for single  
 515 cells, thereby enabling *M. woodiana* to be important in export (*M. woodiana* and *H. hauckii*  
 516 numerically dominate diatom flux at ALOHA during summer; Scharek *et al.*, 1999b). The  
 517 PB09-bloom  $\int \rho_N$  implies a maximum potential for export of 1.1 mmol Si m<sup>-2</sup> d<sup>-1</sup>, which ~40%  
 518 higher than the highest  $\rho_E$  observed in this region (Fig. 6C), and consistent with diatom export  
 519 during summer blooms being important to regional carbon export (Karl *et al.*, 2012).

520

#### 521 4.5. The potential summer bloom effect on the annual Si budget in the NPSG

522 During 2008 and 2009 at station ALOHA, blooms accounted for 29% of annual  $\int \rho_P$   
 523 (Brzezinski *et al.*, 2011). This percentage is based on average daily rates of 0.4 mmol Si m<sup>-2</sup> d<sup>-1</sup>  
 524 during two summer-period blooms. Incorporating the  $\int \rho_P$  from the active PB09 bloom (~ same  
 525 latitude as ALOHA; Fig. 1) into the Brzezinski *et al.* (2011) average, increases the bloom-  
 526 average  $\int \rho_P$  from 0.4 mmol Si m<sup>-2</sup> d<sup>-1</sup> to 0.63 mmol Si m<sup>-2</sup> d<sup>-1</sup> or by nearly 50%. Based on the  
 527 limited number of diatom blooms sampled at or near station ALOHA, the contribution of blooms  
 528 to annual silica production rates calculated by Brzezinski *et al.* (2011) may be conservative.

529 Calculating the contribution of blooms to annual budgets requires their total silica  
 530 production, frequency, and areal extent. Bloom frequency and areal extent are best obtained  
 531 from satellite ocean color recognizing the limitations of linking DDA-diatom blooms to ocean

532 color (Villareal *et al.*, 2011; 2012). The PB08 bloom had remotely-observed [Chl *a*] of >0.12 µg  
 533 L<sup>-1</sup> over an average area of 100,000 km<sup>2</sup>. The active PB09 bloom had significantly higher silica  
 534 production but covered only 30,000 km<sup>2</sup>. Both blooms persisted ~16 days but during this time  
 535 period cloud cover sometimes obscured the bloom, introducing a likely error of at least a few  
 536 days. Using the  $\int \rho_P$  for the bloom stations (e.g. PB09 average 1.33 mmol Si m<sup>-2</sup> d<sup>-1</sup>), we estimate  
 537 that the PB08 and PB09 blooms produced 1.04 and 0.64 Gmoles of bSiO<sub>2</sub>, respectively (Table  
 538 2). For comparison, non-bloom rates expressed over this same area and duration would yield  
 539 0.21 and 0.12 Gmoles of bSiO<sub>2</sub> for PB08 and PB09, respectively. Due to their tremendous areal  
 540 extent, these blooms also are significant when considering blooms in other high diatom biomass  
 541 regions (e.g. Monterey Bay, Santa Barbara Channel; Table 2). And, while the PB09 bloom was  
 542 much more biologically active by all our proxies, the larger size of the PB08 bloom more than  
 543 offset the difference in production rates and suggests even diffuse blooms significantly affect  
 544 regional Si budgets. A caveat to these estimations is that the blooms were clearly sampled in  
 545 different stages of their development; therefore, the amount of biogenic silica produced in the  
 546 PB08 bloom is likely underestimated, especially given the higher reported  $\int \rho_P$  by Brzezinski *et*  
 547 al. (1998) in this sector of the NPSG during summer (Fig. 6B).

548 Wilson *et al.* (2008) demonstrated that blooms in the northeastern quadrant of the NPSG,  
 549 near the subtropical front, are significantly larger in area (up to five-fold) and have longer  
 550 duration than blooms near station ALOHA (i.e. gyre-interior). MODIS [Chl *a*] data between  
 551 June and October of 2008 and 2009 show strong differences in the extent of blooms in each year  
 552 of our field study (Fig. 7). A caveat to our analysis is the large amount of cloud cover obscured  
 553 the study region; therefore, the days a particular location was above our threshold [Chl *a*] (color  
 554 scale, Fig. 7A, B) are conservative for all areas. By averaging the  $\int \rho_P$  from all three bloom  
 555 stations during the PB cruises (0.99 mmol Si m<sup>-2</sup> d<sup>-1</sup>), and assuming a 10-day duration (minimum  
 556 estimate based on MODIS data, Fig. 7A, B) we can extrapolate for the bSiO<sub>2</sub> produced during  
 557 the 2008 and 2009 bloom seasons in three latitudinal bands (Fig. 7C, Table 2). In both years, the  
 558 majority of silica production occurred between 30 – 35°N (Table 2). Overall, in 2008 and 2009  
 559 the northern latitude blooms (e.g. 30 – 35 °N) produced 3x and 25x more bSiO<sub>2</sub> than blooms in  
 560 the gyre interior (e.g. 20 – 25 °N), as a function of the larger bloom area during both years.  
 561 Given the differences in the size and duration between blooms in the northeastern portion of the  
 562 gyre and those in the interior near station ALOHA (e.g. Fig. 7), the gyre-interior blooms would

563 have to sustain  $\int \rho_P$  nearly an order of magnitude faster than rates in the north to produce more  
564 bSiO<sub>2</sub> than the northeastern-gyre blooms (Table 2). Considering that the PB08 bloom rate, even  
565 under declining conditions, was 49% of the rate observed in the active PB09 bloom, this seems  
566 unlikely. While the contribution of diatoms to silica and organic matter production and export is  
567 well documented at station ALOHA in the gyre interior (Scharek *et al.*, 1999a; b; Brzezinski *et*  
568 *al.*, 2011; Karl *et al.*, 2012) this analysis suggests that the role of diatom blooms to these  
569 processes is even greater in the northeastern reaches of the gyre near the subtropical front.

570 The temporal resolution of studies reporting Si biogeochemical data in the NSPG outside of  
571 station ALOHA is very poor, and not conducive for constructing an annual budget. The  
572 northeastern region of the NSPG near the subtropical front zone may operate differently in  
573 regards to annual rates of silica production and export than in the gyre interior near station  
574 ALOHA as a consequence of spatially-larger and longer-persisting blooms (Wilson *et al.*, 2008).  
575 Here we demonstrate that there is significant spatial variation in siliceous biomass and  
576 production rates in the NSPG during the summer, which may be of biogeochemical importance  
577 for a regional Si budget and especially to annual rates diatom silica and carbon export. The  
578 northeastern region of the NSPG near the subtropical front, where the most expansive blooms  
579 occur, is not amenable to monthly sampling as is done at station ALOHA. However, future  
580 studies in this and other sectors of the NSPG, could examine processes such export by using  
581 passive (e.g. sediment traps) methods or autonomous vehicles (e.g. glider with fluorometer  
582 and/or transmissometer) which sample with higher temporal resolution.

583

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735 Figure Captions:

736

737 Fig. 1: Station locations during the PB08 and PB09 cruises, station ALOHA (HOT program) is  
738 shown for reference. Symbols are distinguished by year (PB08, circles; PB09, squares) and  
739 bloom stations (filled symbols). Outlined regions are the maximum areal extent of the MODIS-  
740 defined bloom regions ( $>0.12 \mu\text{g L}^{-1}$  [Chl *a*]). Transect direction is denoted by arrows.

741

742 Fig. 2: Section plots showing hydrography, nutrients and biological properties with depth during  
743 the PB08 transects near the subtropical front. A) Salinity and temperature (contours), B) diatom  
744 abundance, C) dissolved silicate concentration, D) biogenic silica concentration [ $\text{bSiO}_2$ ], E)  
745 biogenic silica production,  $\rho_p$  and F) specific biogenic silica production,  $V_b$ . The bloom station  
746 data is not shown as it was situated slightly west of transect 1 (see Fig. 1).

747

748 Fig. 3: Section plots showing hydrography, nutrients and biological properties with depth during  
749 the PB09 transect. Panels and scales are as in Figure 2. Bloom stations were located far west of  
750 this transect and are not shown (see Fig. 4).

751

752 Fig. 4: Vertical profiles of nutrients and biological properties at the PB bloom stations. Each  
753 bloom station is distinguished by symbol type: PB08 (black), PB09 St. 22 (gray crossed), PB09  
754 St. 23 (gray). For comparison, non-bloom (N.B.) station data were binned by depth, averaged,  
755 and plotted (depth is the midpoint of the bin) for PB08 (black line) and PB09 (gray line).

756

757 Fig. 5: The net production of  $\text{bSiO}_2$  ( $\int \rho_N$ ) during A) PB08 and B) PB09. Bars represent vertical  
758 integrals based either in the mixed layer or to a specific light depth. \*Indicates bloom station  
759 during PB09, the PB08 bloom station was not sampled. Error bars are propagated from the  
760 measured variance in the replicates, and summed with depth.

761

762

763 Fig. 6: Comparison of A)  $\int bSiO_2$ , B)  $\int \rho_p$ , and C) 150-m  $bSiO_2$  export in the NPSG. The square  
764 symbols and error bars are the average and standard deviation, respectively, filled symbols  
765 represent data range. Data are arranged by non-summer (September through May) and summer  
766 (bloom season, June through August) months. Data is from Brzezinski et al. (2011) (station  
767 ALOHA), Brzezinski et al. (1998) (transects north of ALOHA during August 1995, July 1996),  
768 and this study; “nd” indicates no data. The PB08 bloom export value at 300 m is averaged with  
769 the flux values at 150 m from in and out of the bloom area (total of three data points).

770

771 Fig. 7: Comparison of 2008 and 2009 bloom seasons in the NPSG using the MODIS dataset. A,  
772 B) days where daily [Chl  $a$ ] was  $>0.12 \mu g L^{-1}$  for 2008 (A) and 2009 (B) between June and  
773 October. C) The area ( $km^2$ ) where [Chl  $a$ ] was  $>0.12 \mu g L^{-1}$ ; values were binned by latitude and  
774 represent the same time period and data from A, B. The longitude considered was between 180–  
775 130° W, except for the 30 – 35° N bin where the only data between 180–135°W was used (to  
776 avoid the California Current boundary). Note: because these use daily MODIS values, the  
777 duration of each bloom is potentially underestimated due to cloud cover (i.e. days with no  
778 recorded value), this is the case for the PB09 bloom feature which was only successfully viewed  
779 during five different days, albeit over a 16-day span.

780

781 Highlights:

- 782 • Bloom biogenic silica stocks and production were enhanced by an order of magnitude  
783 • Bloom biogenic silica export at 300 m was among the highest values in the region  
784 • 77% of gross silica production in one bloom accumulated as biomass (54% is normal)  
785 • Northeastern blooms potentially produced 3–25x more silica than gyre-interior blooms  
786

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Table 1: Mixed layer depth (MLD), dissolved silicate, biological and particulate data for the PB cruises. All integrations are to 150 m (common depth for all stations and generally shallower than the 0.1% light depth) except for the total diatoms (MLD integration).  $V_b$  is average in the upper 150 m (see Methods). Note: integration depths for diatoms, Chl  $a$  and bSiO<sub>2</sub> differ from those presented in Villareal et al. (2012), as their study focused on surface flora (i.e. 0 – 60 m).

Cruise	Station	Lat. (°N)	Long. (°W)	MLD (m)	MLD Ave. [Si(OH) <sub>4</sub> ] (µM)	MLD [Si(OH) <sub>4</sub> ] (10 <sup>6</sup> m <sup>-2</sup> )	% Diatoms	Dominant Diatom (%)	bSiO <sub>2</sub> (mmol m <sup>-2</sup> )	Chl $a$ , >10 µm (mg m <sup>-2</sup> )	Chl $a$ , >0.4 µm (mg m <sup>-2</sup> )	$\int \rho_p$ (mmol Si m <sup>-2</sup> d <sup>-1</sup> )	Ave. $V_b$ (d <sup>-1</sup> )
PB08	1	30.00	150.00	23.3	1.90	6.27	Hemiaulus (69)	2.34	17.1	1.21	0.19	0.08	
PB08	2	30.00	146.50	35.0	1.60	10.6	Hemiaulus (64)	3.06	14.4	1.10	0.19	0.05	
PB08	3	32.35	140.86	11.5	1.67	4.90	Hemiaulus (89)	7.18	17.9	1.35	0.25	0.03	
PB08	4	32.06	140.67	22.0	1.63	—	—	9.16	20.1	1.71	—	—	
PB08	5	31.77	140.47	11.8	1.93	8.32	Hemiaulus (46)	6.16	17.7	1.26	0.18	0.03	
PB08	6	31.48	140.28	15.0	1.95	—	—	3.80	21.4	1.61	—	—	
PB08	*7	31.77	140.71	11.3	1.22	82.9	Hemiaulus (97)	12.83	18.3	2.50	0.65	0.04	
PB08	8	31.67	140.92	13.0	1.39	—	—	6.85	21.7	2.01	—	—	
PB08	9	31.19	140.08	27.3	1.55	8.29	Hemiaulus (90)	6.27	17.1	1.36	0.22	0.04	
PB08	10	30.90	139.89	25.0	1.62	—	—	5.33	19.0	1.33	—	—	
PB08	12	30.61	139.70	24.5	1.53	3.46	Hemiaulus (83)	5.48	17.3	1.35	0.10	0.02	
PB08	13	30.32	139.51	14.0	1.66	—	—	5.23	19.1	1.44	—	—	
PB08	14	28.75	141.25	21.3	1.50	2.26	Hemiaulus (89)	5.11	15.8	1.81	0.25	0.06	
PB08	15	31.52	140.58	24.7	1.68	—	—	6.74	18.4	1.54	0.30	0.04	
PB08	16	32.00	139.00	24.0	2.06	1.06	Hemiaulus (81)	3.67	19.2	1.75	0.20	0.06	
PB08	17	31.67	139.33	22.0	2.10	0.25	Hemiaulus (99)	2.79	20.8	1.63	—	—	
PB08	18	31.33	139.67	26.0	1.68	10.6	Hemiaulus (98)	3.69	19.9	1.63	—	—	
PB08	19	31.00	140.01	21.0	1.49	12.5	Hemiaulus (97)	5.08	20.1	1.69	—	—	
PB08	20	30.67	140.34	12.0	1.58	4.06	Hemiaulus (99)	2.66	19.4	1.52	—	—	
PB08	21	30.33	140.67	22.0	1.60	3.93	Hemiaulus (~100)	—	20.2	1.59	—	—	
PB09	1	20.50	153.69	34.5	1.19	3.79	Mastogloia (73)	2.12	21.0	1.88	0.15	0.07	
PB09	2	25.20	147.00	28.0	1.04	1.01	Mastogloia (36)	1.80	16.5	1.37	—	—	
PB09	*3	26.10	144.70	43.0	0.99	0.99	Mastogloia (67)	2.11	17.7	1.09	0.10	0.05	
PB09	*4	26.10	144.35	37.0	1.02	0.49	Mastogloia (64)	1.91	18.8	1.71	—	—	
PB09	*7	26.10	144.00	49.0	0.86	0.25	Mastogloia (49)	2.03	4.68	0.31	0.10	0.05	
PB09	*8	26.10	143.65	42.0	1.03	0.69	Hemiaulus (34)	1.57	16.0	0.95	—	—	
PB09	*11	26.10	143.30	43.5	1.32	0.56	Mastogloia (74)	1.77	16.4	0.93	0.12	0.07	
PB09	*12	26.10	142.95	8.0	1.34	0.20	Mastogloia (~100)	1.80	15.3	0.80	—	—	
PB09	*13	25.48	143.00	36.8	1.02	1.27	Hemiaulus (55)	1.99	14.7	0.93	0.13	0.06	
PB09	*15	26.10	142.60	40.0	1.37	—	—	2.04	15.0	0.76	—	—	

PB09	‡16	26.10	142.25	58.0	1.02	—	—	2.24	17.1	1.20	—	—
PB09	‡17	26.10	141.90	34.5	1.28	2.46	<i>Mastogloia</i> (56)	2.34	20.9	1.39	—	—
PB09	‡18	26.10	141.55	48.0	1.26	0.89	<i>Mastogloia</i> (86)	2.05	15.2	0.94	—	—
PB09	20	27.50	142.00	37.0	1.30	—	—	2.54	15.7	1.00	—	—
PB09	21	23.50	140.00	58.6	1.30	6.96	<i>Chaetoceros</i> (74)	1.69	17.6	1.18	0.10	0.06
PB09	* ,‡22	25.10	154.70	47.7	0.91	509	<i>Mastogloia</i> (98)	6.13	63.3	18.05	0.91	0.16
PB09	* ,‡23	25.18	154.60	52.7	0.82	2410	<i>Mastogloia</i> (99)	9.19	43.7	18.53	1.74	0.17

\* Bloom stations (i.e. fulfilled data-based criteria)

† Faded bloom stations (PB09; 26°N, 145°W)

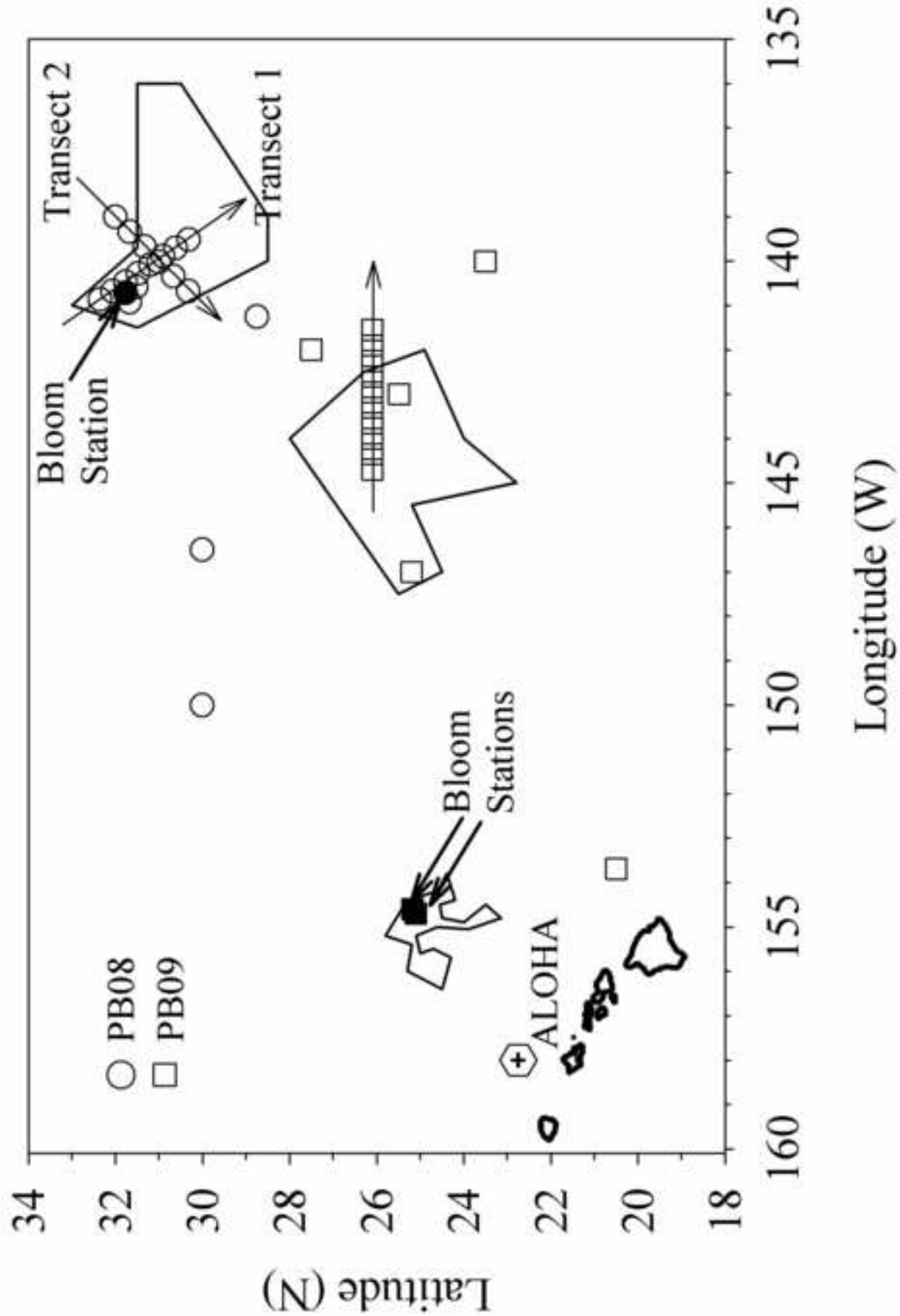
‡ Integration depth (0.1%I<sub>0</sub>) was only 131 and 134 m for stations 22 and 23, respectively.

Table 2: Estimates for the amount of bSiO<sub>2</sub> produced by the individual blooms sampled in 2008 and 2009 (active bloom), and for the bSiO<sub>2</sub> produced during the bloom season (June-October) in three latitude bins, assuming a duration of 10 days (see Fig. 7). For comparison, we have included bSiO<sub>2</sub> produced in other regions using published bloom rates and also assuming a 10-day duration.

Bloom or Region	$\int \rho_p$ (mmol Si m <sup>-2</sup> d <sup>-1</sup> )	Duration (days)	<sup>†</sup> Bloom Area (km <sup>2</sup> )	bSiO <sub>2</sub> Produced (Gmoles)	Reference
PB08 bloom	0.65	16	100,000	1.04	This study
PB09 bloom	1.33	16	30,000	0.64	
NPSG 2008 bloom season					
20-25°N	0.99	10	234,798	2.32	This study
25-30°N	0.99	10	114,425	1.13	
30-35°N	0.99	10	758,114	7.51	
NPSG 2009 bloom season					
20-25°N	0.99	10	15,052	0.15	This study
25-30°N	0.99	10	57,156	0.57	
30-35°N	0.99	10	386,566	3.83	
Sargasso Sea: mode-water eddy	0.98	10	24,200	0.24	Krause <i>et al.</i> , 2010b
Monterey Bay	60	10	1,200	0.72	Brzezinski <i>et al.</i> , 1997, 2003
Santa Barbara Channel	40	10	4,200	1.68	Shipe and Brzezinski, 2001
Ross Sea polynya	34	10	<sup>‡</sup> 160,000	54.4	Nelson <i>et al.</i> , 1991

<sup>†</sup>Bloom rate is assumed over the *entire* regional area

<sup>‡</sup>Average open water area (October - May) over nine years (Arrigo and van Dijken, 2004)



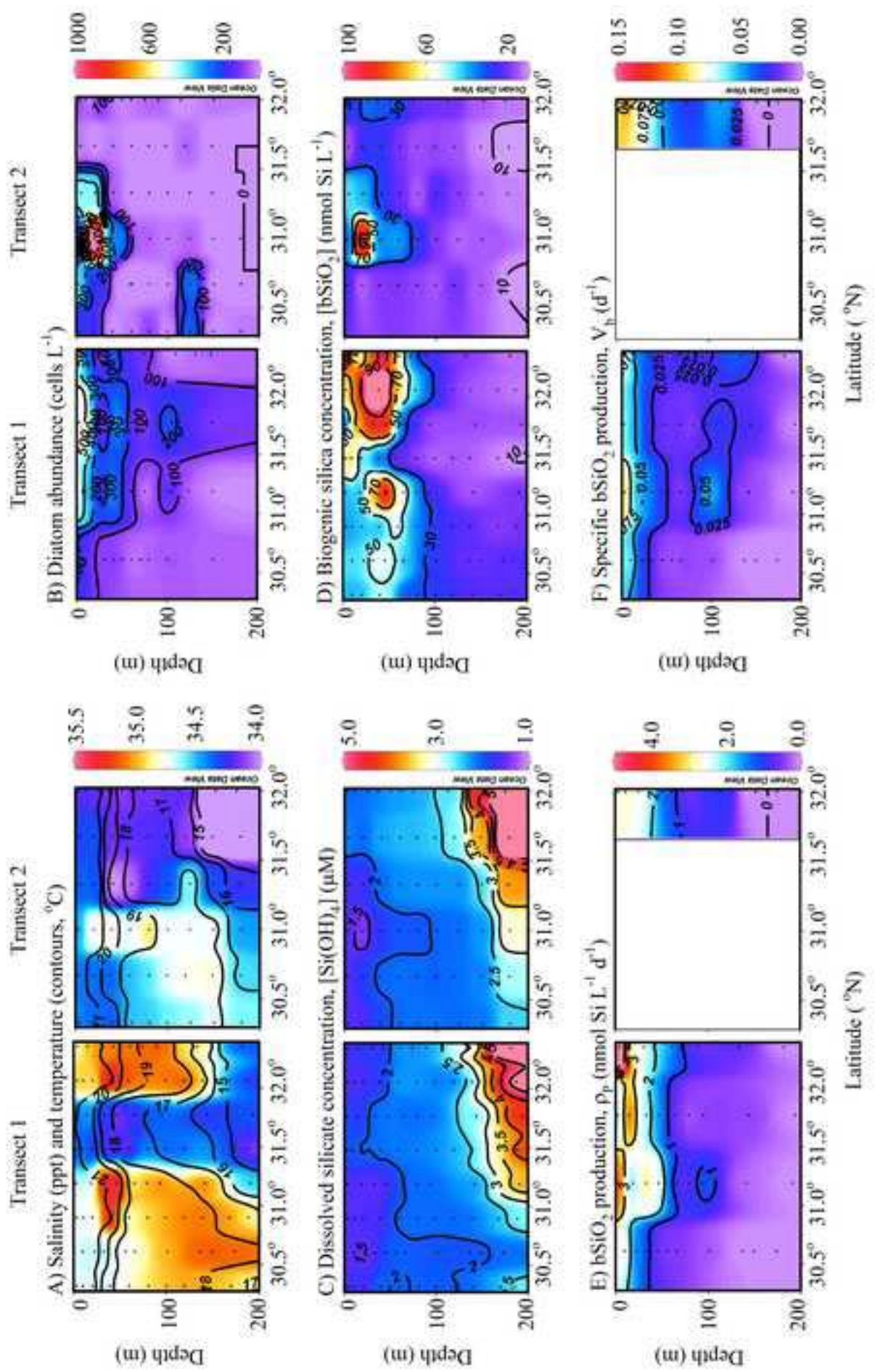
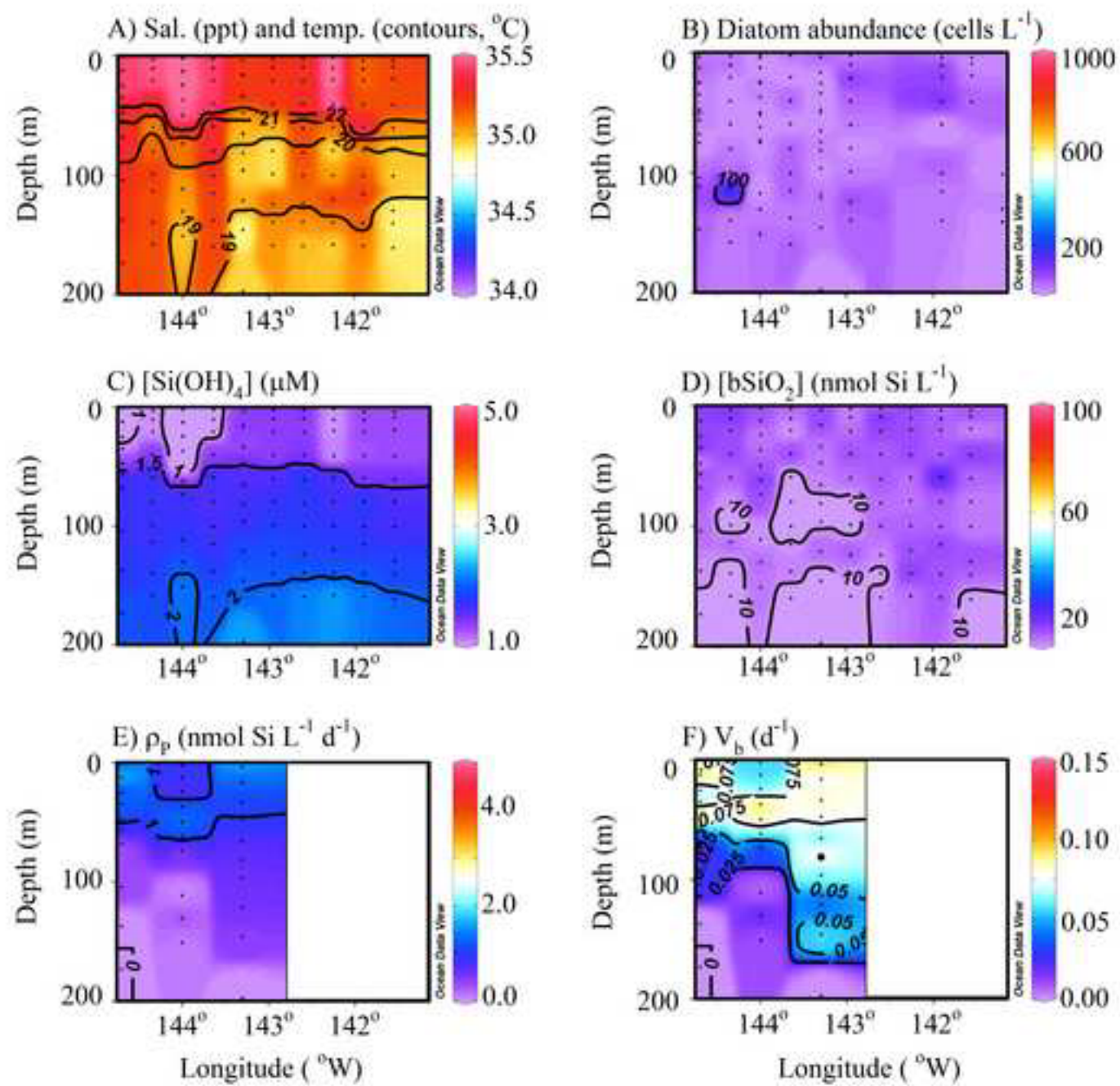


Figure 2

Figure 3



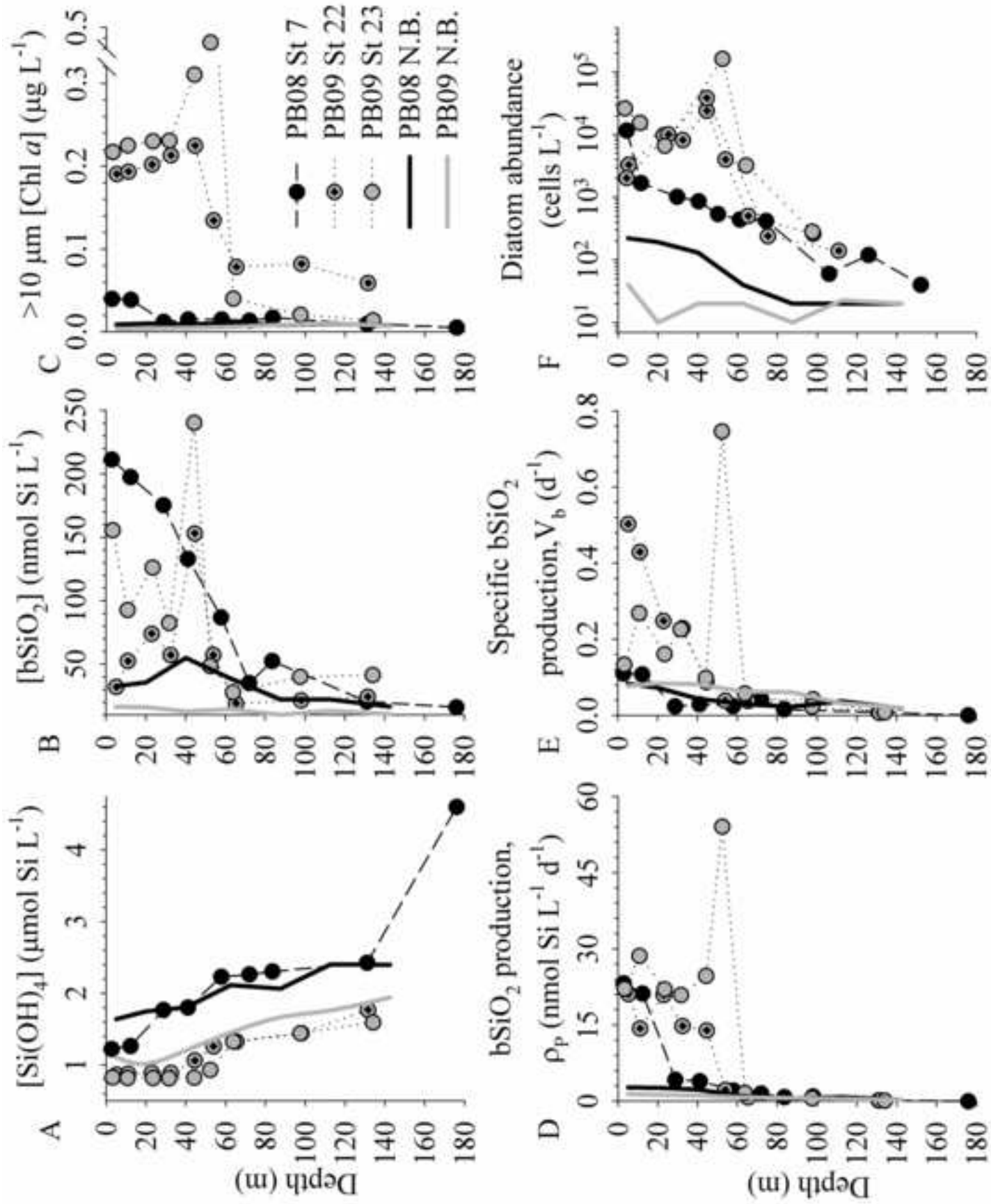
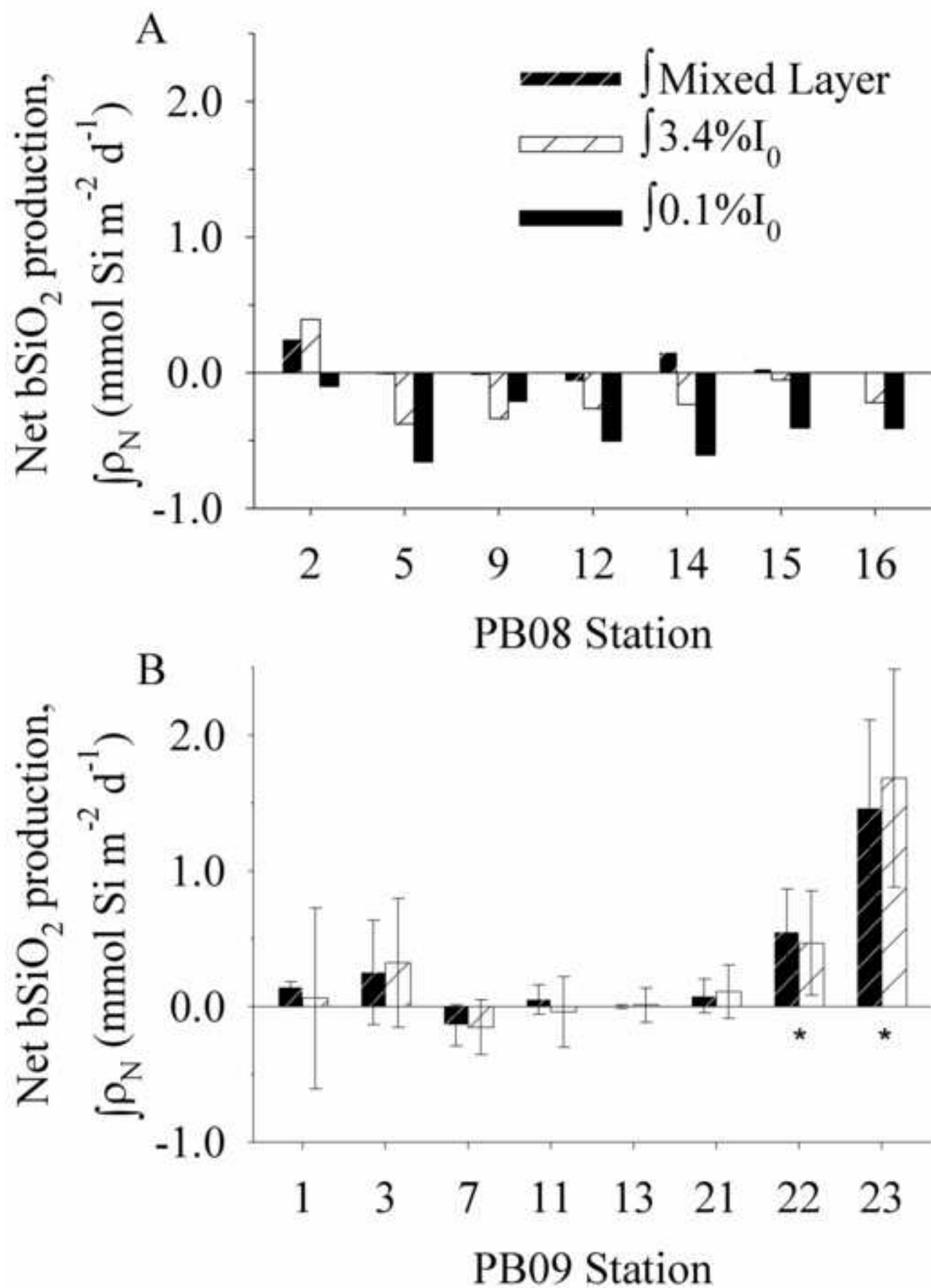
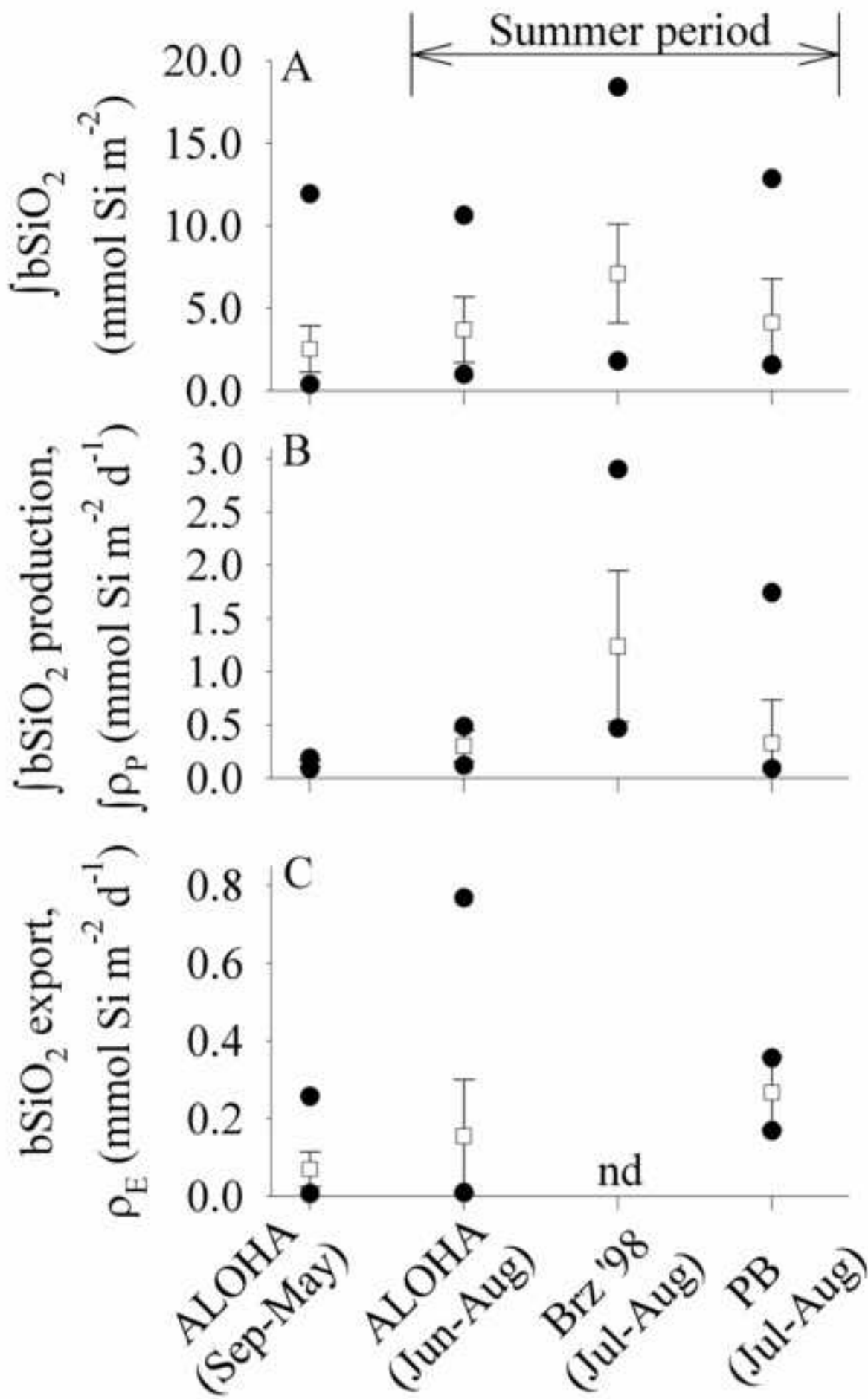


Figure 4





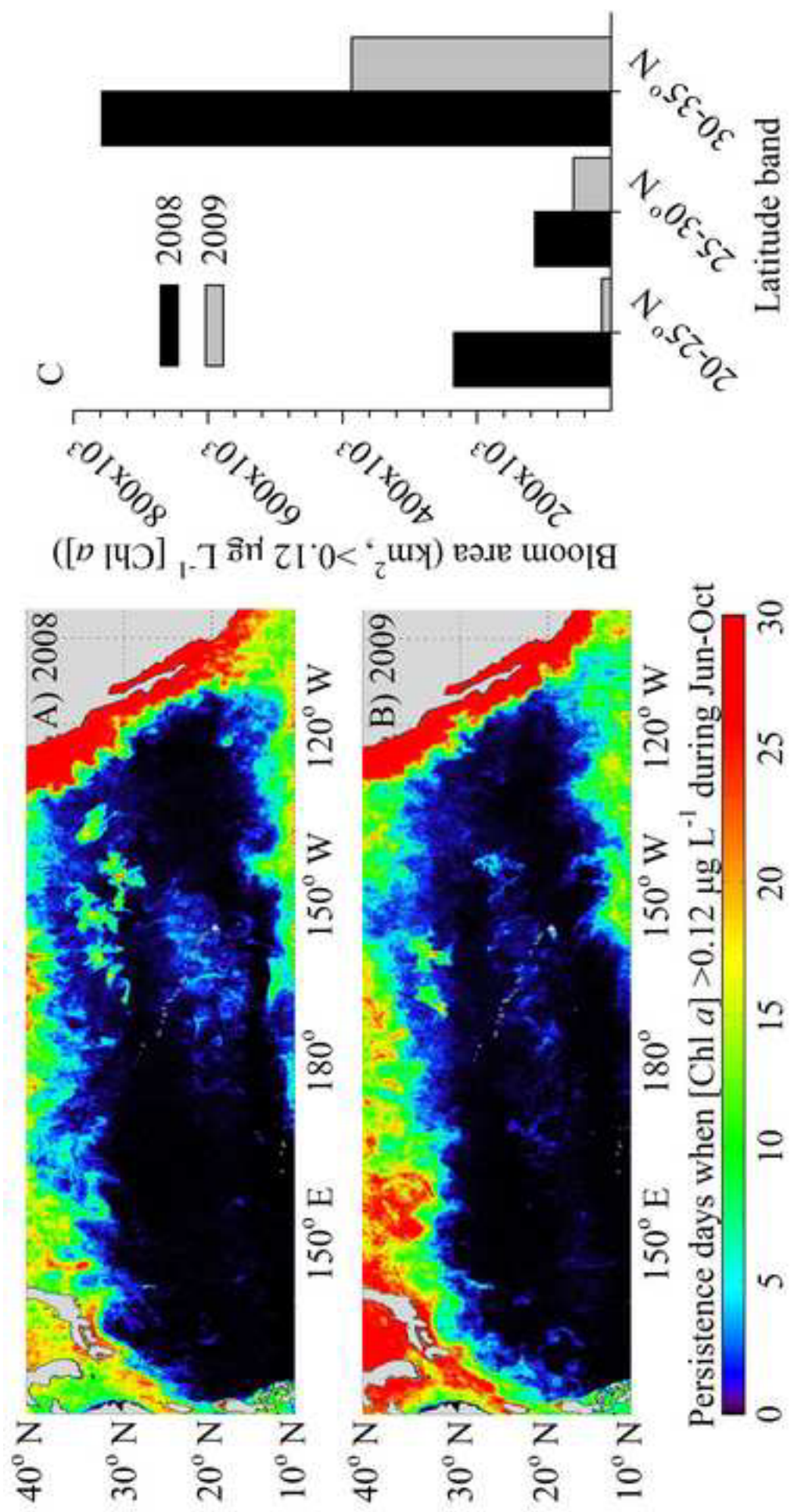


Figure 7