Physical forcing of phytoplankton community structure and primary production in continental shelf waters of the Western Antarctic Peninsula

by B. B. Prézelin¹, E. E. Hofmann², M. Moline³ and J. M. Klinck²

ABSTRACT

Analyses of a multidisciplinary data set, collected in continental shelf waters of the Western Antarctic Peninsula (WAP) during austral summer of January 1993, identified a previously unrecognized forcing mechanism that sets up a physical and chemical structure that supports and assures site-specific diatom-dominated communities and enhanced biological production (Prézelin et al., 2000). This forcing is active when the southern boundary of the Antarctic Circumpolar Current (ACC) flows along the shelf edge, thereby facilitating onshelf bottom intrusions of nutrient-rich Upper Circumpolar Deep Water (UCDW), which then is upwelled or mixed into the upper water column. At times or locations where UCDW is not introduced to the upper water column, diatoms no longer dominate phytoplankton assemblages over the mid- to outer WAP continental shelf. This analysis extends the area and seasons studied through similar analyses of multidisciplinary data sets collected on four additional cruises to the WAP that cover all seasons. Results show that onshelf intrusions of UCDW: (1) occur in other regions of the WAP continental shelf; (2) are episodic; (3) are forced by nonseasonal physical processes; and (4) produce areas of diatom-dominated phytoplankton assemblages and enhanced primary production. At times, multiple intrusions are observed on the WAP continental shelf, and each event may be in a different stage. Further, the occurrence of an intrusion event in one area does not necessarily imply that similar events are ongoing in other areas along the WAP shelf. The UCDW bottom intrusions originate along the outer shelf but they can extend into the inner shelf region because the deep troughs that transect the WAP shelf provide connections between the inner and outer shelf. The boundary between the intruded water and the shelf water is variable in location because of the episodic nature of the onshelf intrusions, and is moved farther inshore as an event occurs. These observations show clearly that the phytoplankton community structure on the WAP shelf is determined by physical forcing and that primary production is likely to be considerably greater than previously believed. Moreover, variability in this physical forcing, such as may occur via climate change, can potentially affect the overall biological production of the WAP continental shelf system.

1. Marine Science Institute and Department of Ecology, Evolution and Marine Biology, University of California, Santa Barbara, California, 93106, U.S.A. email: prezelin@lifesci.ucsb.edu
2. Center for Coastal Physical Oceanography, Old Dominion University, Norfolk, Virginia 23529, U.S.A.
3. Biological Sciences Department, California Polytechnic State University, San Luis Obispo, California, 93407, U.S.A.
1. Introduction

Coincident environmental and biological data sets collected in January 1993, in continental shelf waters of the Western Antarctic Peninsula (WAP) (Fig. 1), were used to describe a previously unrecognized mechanism that sets up a physical and chemical structure that supports enhanced biological production (Prézelin et al., 2000). This mechanism is provided by onshelf subsurface intrusions of warm (>1.5°C) Upper Circumpolar Deep Water (UCDW), which then upwells and/or is mixed upward, introducing elevated concentrations of nutrients and potentially other constituents, such as trace metals, into the upper water column. Diatom-dominated phytoplankton assemblages, forming subsurface chlorophyll $a$ maxima above the pycnocline, characterize the regions where UCDW is mixed into the surface waters. Other phytoplankton taxa dominate outside of these areas (Fig. 1). In addition, the distribution of higher trophic level consumers, such as Antarctic krill (*Euphausia superba*), is influenced by UCDW intrusions (Prézelin et al., 2000). The diatom-dominated communities develop largely below optical depths used in satellite detection and do not conform to remote sensing assumptions of correlation between detectable surface chlorophyll $a$ and deeper chlorophyll $a$ (Moline and Prézelin, 2000). The UCDW-influenced diatom-dominated communities that develop over the outer continental shelf are subject to advection off the shelf during intrusion events and to aggregation and sedimentation following events. These observations led Prézelin et al. (2000) to suggest that the physical environment is a primary factor structuring the phytoplankton community composition in WAP continental shelf waters and that bio-optical methods may underestimate primary production of regions influenced by UCDW. The dynamics of the coupled bio-optical and biological-physical controls of primary production in this physically forced system during January 1993 has been modeled and indicates that the production potential of diatom-dominated communities could be underes-

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Figure 1. (a) Map of the Western Antarctic Peninsula and sites where comparable data sets were collected over the same region of the continental shelf during five cruises that occurred in November 1991, January 1993, March 1993, August 1993, and January 1994 (shaded box parallel to coastline). The findings are placed in a larger context by comparison with observations from other regions of the Antarctic, including cross-shelf transects made in the vicinity of the Bransfield Strait during the Korean survey cruise in the austral summer of 1993 (Kang and Lee, 1995) and north-south transects across the marginal ice zone in the Bellingshausen Sea at 72°W, beyond the continental shelf, surveyed on the austral spring cruise of Icecolors '90 (Prézelin et al., 1994). (b) Summary of dominant phytoplankton community composition overlaid on the temperature maxima ($T_{max}$) distribution below 250 m for the January 1993 cruise. The diatom-dominated communities are comprised of $>60\%$ diatoms; phytoflagellate-dominated communities are comprised of $>60\%$ prymnesiophytes + pelagophytes; and mixed communities are those in which no algal group was $>60\%$. Sites where intrusions of Upper Circumpolar Deep Water (UCDW) were observed in January 1993 are indicated by the gray arrows. Wind-induced upwelling and bathymetrically-induced mixing sites of UCDW are indicated by the black and clear arrows, respectively (from Prézelin et al., 2000).
timated by more than two fold in the absence of consideration of carbon loss due to across-shelf and vertical advection, especially off the shelf on the outer shelf, and due to zooplankton grazing, especially on the inner shelf (Kim et al., 2003).

The mechanisms by which UCDW intrusions occur were examined using a circulation model developed for the WAP continental shelf (Dinniman and Klinck, 2004). Analyses of the simulated circulation fields showed a significant correlation between the curvature of the shelf break and the volume of UCDW transported across the shelf break. The simulation results show that onshelf movement of UCDW occurs at sites where the Antarctic Circumpolar Current (ACC), which flows along the outer WAP shelf, encounters variations in the topography of the shelf break.

Intrusions of UCDW onto the Antarctic continental shelf can set up regions of enhanced localized biological production. However, the biological effects of UCDW intrusions described in Prézelin et al. (2000) refer to a time in the austral summer when the WAP continental shelf is free of sea ice, light is available, herbivore grazing is potentially at a maximum, surface mixed layers and a seasonal pycnocline exist, and surface water temperatures are at an annual maximum. Thus, for this physical forcing mechanism to be considered a primary determinant of annual biological production in Antarctic continental shelf waters, its occurrence at other times of the year and at other sites needs to be demonstrated.

In this study, hydrographic, nutrient, phytoplankton community composition, and primary production distributions from four additional cruises to the WAP shelf (Table 1) are used to describe UCDW intrusions and their effects. These additional data sets extend the spatial domain along the WAP continental shelf, cover all seasons, and provide a repeat realization of the austral summer. Four of the five cruises occurred in the same year, which allows assessment of environmental and biological changes on time scales of a few weeks to months. Section 4 of the paper summarizes the characteristics of UCDW intrusions, considers the effect of intrusions on nonseasonal succession in phytoplankton communities and highlights the importance of these events for biological production and carbon partitioning in Antarctic continental shelf waters. Section 6 provides suggestions for why UCDW intrusions and associated biology have been unrecognized on the Antarctic continental shelf and addresses the potential implications of these events for biological production of Antarctic coastal waters.

2. Methods

a. Study area

The same region of the WAP continental shelf was sampled during four cruises that took place between January 1993 and January 1994 and a fifth cruise that occurred in November 1991 (Table 1). Hydrographic and biological samples were collected at fixed station locations aligned in across-shelf transects from the inner shelf to offshore of the shelf break (Fig. 2). Station spacing was 20 km along transects and 100 km between transects. From
north to south, transects were numbered from 800 to 000. Details of sampling strategies for individual cruises are given in Lascara et al. (1993a,b), Smith et al. (1993a,b), and Prézelin et al. (2000).

For the analyses presented in the following sections, the study area was divided into outer, mid- and inner shelf regions. The outer region includes the shelf area that is within about 50 km of the continental shelf break at 500 to 700 m depth and is directly affected by the ACC. The mid-shelf region is between 50 km and 100 km of the continental shelf break. The inner shelf refers to those regions more than 100 km from the continental shelf break.

b. Hydrographic samples

Conductivity-temperature-depth (CTD) measurements were made using a SeaBird 911+ CTD system. On November 1991, January 1993, August 1993 and January 1994 cruises, the CTD system was mounted on a bio-optical profiling system, which limited the vertical profiles to 500 m, but for most stations, this depth exceeded or was near the bottom. For the March-May cruise, the CTD system was separate from the bio-optical profiling

<table>
<thead>
<tr>
<th>Cruise dates and season</th>
<th>Cruise length (days)</th>
<th>Daylength (hrs)</th>
<th>Across-shelf transects sampled*</th>
<th>Area covered (km²)</th>
<th>CTD stations</th>
<th>Biological sampling stations</th>
<th>HPLC samples</th>
<th>Nutrient samples</th>
<th>P-I curves</th>
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<tbody>
<tr>
<td>8 Jan. to 8 Feb 93</td>
<td>31</td>
<td>20 to 17.3</td>
<td>200–600</td>
<td>80,000</td>
<td>47</td>
<td>40</td>
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<td>25 Mar to 15 May 93</td>
<td>51</td>
<td>7.8 to 11</td>
<td>000–800</td>
<td>180,000</td>
<td>155</td>
<td>79</td>
<td>643</td>
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<td>186</td>
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<td>(fall)</td>
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<tr>
<td>23 Aug. to 30 Sept 93</td>
<td>38</td>
<td>9.8 to 11.8</td>
<td>200–600</td>
<td>60,000</td>
<td>42</td>
<td>39</td>
<td>291</td>
<td>360</td>
<td>90</td>
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<td>(winter)</td>
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<tr>
<td>11 Jan to 7 Feb 94</td>
<td>27</td>
<td>19.3 to 16.4</td>
<td>300–600</td>
<td>54,000</td>
<td>48</td>
<td>32</td>
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<tr>
<td>(summer)</td>
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<tr>
<td>7 Nov to 21 Nov 91</td>
<td>14</td>
<td>17.9 to 18.6</td>
<td>500–700</td>
<td>20,000</td>
<td>27</td>
<td>26</td>
<td>244</td>
<td>265</td>
<td>250</td>
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<tr>
<td>(spring)</td>
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*See Figure 2.
Figure 2. Comparison of the horizontal distribution of the temperature maximum (Tmax) below 200 m for (a) January–February 1993, (b) March–May 1993, (c) August–September 1993, (d) January–February 1994, and (e) November 1991. Filled circles indicate the distribution of the stations at which the CTD measurements were made. Temperatures ranged from <1.0°C (palest gray) to ≥1.8°C (yellow). (f) Bathymetry (2000 m, 1000 m, 500 m) of the study region.
system and casts were made to within a few meters of the bottom at all stations. Complete
descriptions of the sensor calibrations and data processing are given in Lascara et al.
(1993a,b), Smith et al. (1993a,b), and Klinck and Smith (1994).

c. Nutrient samples

Water samples were collected at discrete depths using either 5-L Go-Flo bottles or 10-L
Niskin bottles, filtered through a 4 μm polycarbonate filter and the filtrate frozen at −20°C.
Nutrient determinations for nitrate, silicate and phosphate samples collected during
November 1991, January 1993 and 1994, and August 1993 were done at the Marine
Science Analytical Laboratories, University of California Santa Barbara, according to the
methods of Johnson et al. (1985). Nutrient samples collected during March–May 1993
were analyzed during the cruise using the same methods.

d. Pigment samples and estimation of phytoplankton community structure

Pigment determinations were made at sea using reverse-phase HPLC procedures
(Prézelin et al., 2000). The regression and validation procedures used for calculations
of the distribution of chlorophyll (Chl a) biomass among the different phytoplankton
groups known to be present are the same as those detailed in Prézelin et al. (2000).
Chlorophyll b (Chl b), 19′-hexanoyloxy-fucoxanthin (HEX-FUCO), 19′-butanoyloxy-
fucoxanthin (BUT-FUCO), fucoxanthin (FUCO), peridinin (PER), and alloxanthin
(ALLO) are the biomarkers for chlorophytes, prymnesiophytes, pelagophytes, dia-
toms, dinoflagellates and cryptophytes, respectively. In this analysis, HEX-FUCO was
combined with BUT-FUCO to provide a single chemotaxonomic marker for what is
termed “phytoflagellates,” which represents the combined populations of prymnesio-
iphytes + pelagophytes; nanoplanckton that generally co-occur throughout much of
Southern Ocean.

e. Photosynthesis-irradiance relationships

Photosynthesis-irradiance (P-I) curves for carbon fixation were determined as detailed in
Boucher and Prézelin (1996) and P-I parameters derived using equations of Neale and
Richerson (1987). Measurements of in situ Q_{PAR} are not available and thus analyses are
presently limited to assessing patterns of mid-day and daily integrated potential carbon
production (volumetric P_{MAX}) and assimilation rates (Chl a-specific P_{MAX}), a proxy for
photosynthetic growth potential. Only findings from the P-I determinations measured as
radiolabel ^{14}C uptake in the upper 60 m around solar noon are compared and the potential
impact of diel variations in P-I are not considered here (however, see Kim et al., 2003;
Mizerek and Prézelin, 2004). Our preliminary estimates of daily rates of integrated primary
production are based upon knowledge of day-length for each station and assuming that the
average hourly production over the day is about 0.6 that of mid-day rates of primary
production.
3. Results

a. Temperature maximum below 200 m

As UCDW approaches the Antarctic continental shelf, it shoals to 150–200 m along the outer shelf edge (Sievers and Nowlin, 1984; Jacobs, 1991; Whitworth et al., 1998). Along the WAP where conditions are favorable, this water mass moves onshore and floods the continental shelf below 200 m (Hofmann and Klinck, 1998; Smith et al., 1999) (Fig. 2a). The movement of UCDW across the WAP continental shelf can be traced by the distribution of the temperature maximum (Tmax) below 200 m (Fig. 2). Temperatures >1.5°C are associated with newly intruded UCDW and the southern boundary of the ACC is associated with the 1.8°C isotherm at 200 m (Orsi et al., 1995). Temperatures of 1.5°C–1.3°C are associated with modified UCDW, which is produced by mixing of UCDW and Antarctic Surface Water (AASW) (Smith et al., 1999). The rate of cooling of UCDW is such that it takes 2–3 months to reach temperatures characteristic of modified UCDW (Klinck, 1998; Smith et al., 1999; Smith and Klinck, 2002). Temperatures <1.3°C represent inner shelf water. In January 1993 (Fig. 2a), the southern boundary of the ACC was located along the outer edge of all transects with the exception of Transect 300 where the ACC was displaced farther offshore. Extending onto the shelf ≥100 km on either side of Transect 300 was a large intrusion of UCDW that reached to the inner shelf along Transect 400. The isotherm pattern during the austral summer of 1993 (Fig. 2a) suggested that the UCDW intrusion to the mid-shelf region of Transects 300–400 displaced modified UCDW from a prior event to the inner shelf.

The temperature maximum distribution approximately two months later (Fig. 2b) showed the southern boundary of the ACC along the outer shelf and onshore movement of UCDW over a 400 km region of shelf between Transects 000 and 400. Modified UCDW was found over most of the mid- and inner shelf. Colder water (<1.3°C) on the inner part of Transects 700–800 is derived from outflow from the Gerlache Strait and is associated with a southerly flowing coastal current that is found along the inner portion of the WAP continental shelf (Smith et al., 1999).

Sea ice was present over the entire region sampled in mid-winter, August 1993 (Fig. 2c). The southern boundary of the ACC was located just off the edge of the continental shelf with a small region of onshore movement of UCDW toward the mid-shelf occurring once again at Transect 300. Most of the remaining outer, mid- and inner shelf of Transects 200–600 were covered with modified UCDW. A small isolated region of temperatures >1.5°C was observed at mid-shelf on Transect 500.

By January 1994 (Fig. 2d), the next austral summer, UCDW was found only along the 300 km of the outer shelf sampled between Transects 300 and 600. No UCDW intrusions were evident on the WAP continental shelf at this time, in striking contrast to physical conditions the previous summer (Fig. 2a). Rather, with the exception of a small patch of 1.5°C water in a part of the mid-shelf region on Transect 500, modified UCDW covered the entire mid-shelf regions between Transects 300 and 600 in the austral summer of 1994.

Observations during the austral spring, November 1991, were limited and the across-
shelf transects did not extend far enough offshore to resolve the location of the southern boundary of the ACC (Fig. 2e). Heavy sea ice covered the study region, with the exception of a region of open water in Dallmann Bay on the inner part of Transect 700. UCDW was present at the outer shelf of Transect 700, near the edge of the marginal ice zone (MIZ). Within the MIZ, the shelf was covered by modified UCDW as were the mid- and inner shelf regions.

b. Silicate to nitrate ratio at 200 m

Unmodified UCDW has a silicate to nitrate [Si(OH)\(_4\):NO\(_3\)] ratio between 3.0 and 3.5 (Smith et al., 1999) and this ratio might be expected below 200 m when UCDW intrusions are present on the shelf of the Western Antarctic Peninsula. However, observations show considerable variability in the 200 m distribution of the Si(OH)\(_4\):NO\(_3\) ratio in regions where the temperature distributions indicate that UCDW intrusions have occurred (Fig. 3). Reduction in the ratio is an indication of prior and/or ongoing biological processing of this UCDW. The lowered Si(OH)\(_4\):NO\(_3\) ratio is attributable to preferential utilization of silicate with respect to nitrate (e.g. Prézelin et al., 2000), which takes place near the pycnocline as low light adapted diatoms respond to the UCDW (see below).

In January 1993, lowest Si(OH)\(_4\):NO\(_3\) ratios at 200 m (<2.0, Fig. 3a) coincided with the onshore intrusion of UCDW between Transects 200 and 400 (Fig. 2a). In areas where UCDW intrusions were present and diatoms dominated, the Si(OH)\(_4\):NO\(_3\) ratios in the upper 100 m averaged about 1.7 (Table 2). The lowering of the Si(OH)\(_4\):NO\(_3\) ratio from a base value of ca. 3.0 for UCDW was due to preferential removal of Si(OH)\(_4\) relative to NO\(_3\) (Prézelin et al., 2000). The highest water column Si(OH)\(_4\):NO\(_3\) ratio detected was 3.05 at a mid-shelf station on Transect 500 sampled two days after a major storm event (Prézelin et al., 2000) and before significant biological processing occurred. The newly upwelled waters were characterized by the uncommon prevalence of chlorophytes mixed with an abundance of diatoms (Prézelin et al., 2000). For five stations on Transects 500 and 600, where chlorophytes were >15% of the chlorophyll \(a\), recent UCDW upwelling was indicated by the hydrographic measurements and the Si(OH)\(_4\):NO\(_3\) ratios >2.6 for the upper 100 m (Table 2). Four stations were sampled at the outflow of the Gerlache Strait that were not influenced by UCDW and at these sites phytoflagellate-dominated communities were observed and water column Si(OH)\(_4\):NO\(_3\) ratios averaged ca 2.0 (Table 2).

In March–May 1993, the Si(OH)\(_4\):NO\(_3\) ratio at 200 m along the outer WAP shelf (Fig. 3b) was about 2.0, with lowest values associated with the onshore intrusion of UCDW along Transects 200 and 300. The 200 m waters of all the nearshore stations of the WAP had Si(OH)\(_4\):NO\(_3\) ratios >2.6 while those over the mid-shelf region ranged between 2.0–2.6. Mid-shelf ratios were the lowest, ca 2.3, on Transect 800 (Fig. 3b) where the mixed layer depths (MLD) were shallower and Si(OH)\(_4\) concentrations were lower than for other regions (Table 3, no. 6). Mid-shelf ratios of Si(OH)\(_4\):NO\(_3\) were highest, ca 2.6, on
Figure 3. Comparison of the horizontal distribution of the Si(OH)₄:NO₃ ratio at a depth of 200 m for (a) January–February 1993, (b) March–May 1993, (c) August–September 1993, (d) January–February 1994, and (e) November 1991. Filled circles indicate the distribution of the sampling sites. The progression from darkest to lightest shaded areas represents Si(OH)₄:NO₃ ratios of >2.8, 2.4 to 2.8, 2.0 to 2.4, 1.6 to 2.0 and <1.6, respectively. (f) Bathymetry (2000 m, 1000 m, 500 m) of the study region.
Table 2. **January–February 1993.** Comparison of mean and standard deviation for mixed layer depths (m), integrated water column Chl $a$ biomass (mg m$^{-2}$) and nutrient concentrations (m mol m$^{-2}$), as well as averaged ratios of nutrient concentrations, for the upper 100 m. Data are sorted by percent dominance by diatoms or phytoflagellates within phytoplankton assemblages surveyed between Transects 200–600 along the west Antarctic Peninsula. Listed is the number of stations where the criteria for each subset is met. Not included are stations where the vertical sampling did not extend to 100 m. Water column characteristics are also summarized for chlorophytes and dinoflagellates at stations where they comprised >15% of the integrated Chl $a$ biomass. Community comparisons expressed as the ratio of diatoms: phytoflagellate and diatom: chlorophyte water column characteristics are provided only if the difference between assemblages was significant at the >95% confidence level (t-test). ns, not significant; na, not applicable; —, not measured.

**Diatoms dominated biologically processed waters of UCDW-influenced sites; Phytoflagellates dominated nearshore waters at outflow from the Gerlache Strait; Chlorophytes were distributed in waters upwelled within two days of sampling; Dinoflagellates were localized at one site within the zone of phytoflagellate-dominated communities.**

<table>
<thead>
<tr>
<th>Phytoplankton</th>
<th>Stations</th>
<th>MLD</th>
<th>$\Sigma$ Chl $a$</th>
<th>$\Sigma$ Si(OH)$_4$</th>
<th>$\Sigma$ NO$_3$</th>
<th>$\Sigma$ NH$_3$</th>
<th>$\Sigma$ PO$_4$</th>
<th>Si(OH)$_4$:NO$_3$</th>
<th>NO$_3$:PO$_4$</th>
<th>Si(OH)$_4$:NH$_3$</th>
<th>NO$_3$::PO$_4$</th>
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<tbody>
<tr>
<td>Diatoms &gt;50%</td>
<td>15</td>
<td>71 ± 11</td>
<td>30 ± 10</td>
<td>3.06 ± 0.39</td>
<td>1.81 ± 0.11</td>
<td>—</td>
<td>.131 ± .013</td>
<td>1.69 ± 0.2</td>
<td>13.9 ± 1.6</td>
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<tr>
<td>Diatoms &gt;60%</td>
<td>6</td>
<td>68 ± 11</td>
<td>25 ± 4</td>
<td>2.80 ± 0.30</td>
<td>1.85 ± 0.14</td>
<td>—</td>
<td>.135 ± .011</td>
<td>1.53 ± 0.1</td>
<td>13.8 ± 0.9</td>
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<tr>
<td>Phytoflag. &gt;50%</td>
<td>15</td>
<td>absent</td>
<td>43 ± 14</td>
<td>3.81 ± 0.85</td>
<td>1.78 ± 0.23</td>
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<td>.135 ± .026</td>
<td>2.14 ± 0.4</td>
<td>13.5 ± 1.8</td>
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<td>Phytoflag. &gt;60%</td>
<td>4</td>
<td>absent</td>
<td>38 ± 12</td>
<td>3.84 ± 0.42</td>
<td>1.73 ± 0.16</td>
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<td>.132 ± .008</td>
<td>2.21 ± 0.4</td>
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<td>Chlorophytes &gt;15%</td>
<td>1</td>
<td>absent</td>
<td>50 ± 14</td>
<td>6.58 ± 21</td>
<td>2.16 ± 0.19</td>
<td>—</td>
<td>.442 ± .012</td>
<td>2.61 ± 0.202</td>
<td>11.2 ± 1.0</td>
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<td>aT$<em>{500}$–T$</em>{600}$</td>
<td>5</td>
<td>65 ± 26</td>
<td>50 ± 14</td>
<td>5.02 ± 0.68</td>
<td>2.11 ± 0.19</td>
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<td>.189 ± .012</td>
<td>2.61 ± 0.202</td>
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<td>Dinoflag. &gt;15%</td>
<td>2</td>
<td>absent</td>
<td>39 ± 5</td>
<td>5.40 ± 0.20</td>
<td>2.33 ± 0.10</td>
<td>—</td>
<td>.192 ± .010</td>
<td>2.33 ± 0.1</td>
<td>12.1 ± 0.1</td>
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</table>

**Community Comparisons**

| >60% Diatoms | 0.68 | 0.66 | 0.73 | ns | — | ns | 0.692 | ms |
| >60% Phytoflagellates | 0.68 | 0.66 | 0.73 | ns | — | ns | 0.692 | ms |

ainnermost stations on transect line; bover Marguerite Trough
Transects 000–100 (Fig. 3b). The distribution of Si(OH)_4:NO_3 ratios for the upper 100 m of the water columns generally correlated with those at 200 m.

The Si(OH)_4:NO_3 ratio of 3.0 for inner Transect 000 in March–May 1993, where inorganic nutrients and chlorophytes were also abundant, suggested the presence of unmodified UCDW. However, the temperature distribution (Fig. 2b) was colder than unmodified UCDW and the ratio of NO_3:PO_4 in these waters (Table 3) was higher than observed for newly intruded UCDW rich in chlorophytes (Table 2). Using chlorophytes >15% total chlorophyll a as a marker, this cold water region extended to mid-shelf on Transect 100 where Si(OH)_4:NO_3 ratios were ca 2.8 (Table 3).

The Si(OH)_4:NO_3 ratio distribution for August–September 1993 (Fig. 3c) resembled that of January 1993 (Fig. 3a). Ratios <2.0 occurred along the outer part of all transects, coincident with the UCDW temperature distribution (Fig. 2c). The Si(OH)_4:NO_3 ratios for the rest of the region sampled were between 2.0 and 2.5, with highest values in 1.5°C water along the mid-shelf of Transect 500.

The Si(OH)_4:NO_3 pattern observed in January–February 1994 (Fig. 3d) provided a striking contrast to those seen in the 1993 cruises, including the prior summer (Fig. 3a). Values <2.0 were observed over most of the region, with lowest values along Transects 500–600. Exceptions were two locales on Transect 400 where the Si(OH)_4:NO_3 ratios were >2.0. In November 1991, the sea ice-covered UCDW transitional waters had Si(OH)_4:NO_3 ratios <2.0 at 200 m (Fig. 3e) while the upper 100 m had Si(OH)_4:NO_3 ratios that averaged 2.0 ± 0.12 (Table 6). At 200 m in the ice-free region and in the upper 100 m of sampling stations along the 700 transect, Si(OH)_4:NO_3 ratios were slightly greater than 2.0 (Fig. 3e, Table 6).

c. Mixed layer characteristics

The average temperature and salinity of the mixed layer was used to compare the hydrographic structure of the upper water column during the five cruises. The average values were obtained by integrating the temperature and salinity values measured at each hydrographic station to the depth of the maxima in the Brunt-Väisälä frequency, which is a measure of water column stratification. For this analysis, the maximum was assumed to represent the bottom of the mixed layer. The average MLD values were then used to define the portion of a temperature-salinity diagram that characterized the upper water column during each cruise (Fig. 4).

The temperature and salinity of the MLD in January 1993 were characteristic of Antarctic Surface Water (AASW) that forms in the WAP region in the austral summer (Smith et al., 1999). The temperature range reflected seasonal heating that occurs during the spring and summer and the mixed layer salinity range reflected input of freshwater from melting sea ice. The shallowest MLD was ca 65 m and the deepest was >100 m (Table 2).

The March–May period captured the fall transition as the upper water column cooled and sea ice began to form. Average mixed-layer temperature decreased and reached
Table 3. March–May 1993. Comparison of mean and standard deviation for mixed layer depths (m), integrated water column Chl $a$ biomass (mg m$^{-2}$) and nutrient concentrations (m mol m$^{-2}$), as well as averaged ratios of nutrient concentrations, for the upper 100 m. Data are sorted by percent dominance by diatoms or phytoflagellates within phytoplankton assemblages surveyed between Transects 000–900 along the west Antarctic Peninsula. Listed is the number of stations where the criteria for each subset is met. Not included are stations where the vertical sampling did not extend to 100 m. Water column characteristics are also summarized for chlorophytes where they comprised $>15\%$ of the integrated Chl $a$ biomass. At least four diatom-dominated patches, two phytoflagellate-dominated patches and two regions characterized by abundant chlorophytes were present.

<table>
<thead>
<tr>
<th>Phytoplankton</th>
<th>Stations</th>
<th>MLD</th>
<th>$\Sigma$ Chl $a$</th>
<th>$\Sigma$ Si(OH)$_4$</th>
<th>$\Sigma$ NO$_3$</th>
<th>$\Sigma$ NH$_3$</th>
<th>$\Sigma$ PO$_4$</th>
<th>Si(OH)$_4$:NO$_3$</th>
<th>NO$_3$:PO$_4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Diatoms $&gt;50%$</td>
<td></td>
<td></td>
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<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T$<em>{000}$–T$</em>{900}$</td>
<td>33</td>
<td>84 ± 22</td>
<td>34 ± 17</td>
<td>6.19 ± 1.09</td>
<td>2.71 ± 0.10</td>
<td>—</td>
<td>.197 ± .025</td>
<td>2.28 ± 0.37</td>
<td>14.0 ± 1.4</td>
</tr>
<tr>
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<td></td>
</tr>
<tr>
<td>T$_{000}$, at shelf break</td>
<td>2</td>
<td>absent</td>
<td>16 ± 1</td>
<td>4.86 ± 0.01</td>
<td>2.59 ± 0.08</td>
<td>—</td>
<td>.185 ± .007</td>
<td>1.88 ± 0.06</td>
<td>14.3 ± 0.1</td>
</tr>
<tr>
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<td></td>
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<td></td>
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<tr>
<td>T$_{100,000}$, mid shelf</td>
<td>1</td>
<td>50</td>
<td>17</td>
<td>8.14</td>
<td>2.94</td>
<td>—</td>
<td>.220</td>
<td>2.87</td>
<td>13.52</td>
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<td>3. Diatom $&gt;60%$</td>
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<tr>
<td>T$_{300}$–300, mid shelf</td>
<td>3</td>
<td>82 ± 1</td>
<td>23 ± 8</td>
<td>6.89 ± 0.19</td>
<td>2.73 ± 0.05</td>
<td>—</td>
<td>.183 ± .015</td>
<td>2.53 ± 0.03</td>
<td>15.0 ± 1.2</td>
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<tr>
<td>4. Diatom $&gt;64%$</td>
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</tr>
<tr>
<td>T$_{500}$, outer shelf</td>
<td>3</td>
<td>absent</td>
<td>49 ± 9</td>
<td>5.77 ± 0.19</td>
<td>2.60 ± 0.03</td>
<td>—</td>
<td>.177 ± .006</td>
<td>2.22 ± 0.10</td>
<td>14.9 ± 0.3</td>
</tr>
<tr>
<td>5. Diatom $&gt;64%$</td>
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</tr>
<tr>
<td>T$_{600}$</td>
<td>6</td>
<td>absent</td>
<td>51 ± 10</td>
<td>6.26 ± 0.52</td>
<td>2.64 ± 0.08</td>
<td>—</td>
<td>.197 ± .028</td>
<td>2.37 ± 0.13</td>
<td>13.5 ± 1.3</td>
</tr>
<tr>
<td>6. Diatoms $&gt;64%$</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>T$<em>{800}$–T$</em>{900}$</td>
<td>10</td>
<td>90 ± 12</td>
<td>36 ± 17</td>
<td>5.43 ± 1.26</td>
<td>2.76 ± 0.08</td>
<td>—</td>
<td>.210 ± .030</td>
<td>1.98 ± 0.42</td>
<td>13.4 ± 1.8</td>
</tr>
<tr>
<td>All Phytoflag. $&gt;50%$</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>1. Phytoflag, shore</td>
<td>12</td>
<td>85 ± 25</td>
<td>21 ± 10</td>
<td>6.26 ± 1.41</td>
<td>2.75 ± 0.11</td>
<td>—</td>
<td>.202 ± .021</td>
<td>2.27 ± 0.46</td>
<td>13.6 ± 0.9</td>
</tr>
<tr>
<td>2. Phytoflag, shelf</td>
<td>4</td>
<td>83 ± 22</td>
<td>18 ± 7</td>
<td>7.12 ± 0.72</td>
<td>2.71 ± 0.13</td>
<td>—</td>
<td>.190 ± .012</td>
<td>2.62 ± 0.21</td>
<td>14.1 ± 0.4</td>
</tr>
<tr>
<td>All Chlorophytes $&gt;15%$</td>
<td>15</td>
<td>81 ± 16</td>
<td>27 ± 16</td>
<td>6.42 ± 1.13</td>
<td>2.72 ± 0.13</td>
<td>—</td>
<td>.204 ± .018</td>
<td>2.37 ± 0.38</td>
<td>13.4 ± 1.9</td>
</tr>
<tr>
<td>1. T$_{000,020}$–060 $&gt;35%$</td>
<td>2</td>
<td>58 ± 34</td>
<td>13 ± 2</td>
<td>8.39 ± 0.32</td>
<td>2.76 ± 0.24</td>
<td>—</td>
<td>.213 ± .003</td>
<td>3.04 ± 0.15</td>
<td>13.0 ± 1.3</td>
</tr>
<tr>
<td>T$_{100,060}$–080</td>
<td>2</td>
<td>70 ± 21</td>
<td>17 ± 1</td>
<td>6.19 ± 1.09</td>
<td>2.76 ± 0.24</td>
<td>—</td>
<td>.199 ± .002</td>
<td>2.62 ± 0.07</td>
<td>14.2 ± 0.4</td>
</tr>
<tr>
<td>2. Band across shelf</td>
<td>11</td>
<td>85 ± 9</td>
<td>29 ± 17</td>
<td>6.22 ± 0.91</td>
<td>2.72 ± 0.13</td>
<td>—</td>
<td>.202 ± .019</td>
<td>2.26 ± 0.27</td>
<td>13.5 ± 1.2</td>
</tr>
</tbody>
</table>
freezing at $-1.82^\circ C$. Average salinity of the mixed layer increased in response to brine rejection from freezing surface waters. The MLD increased to about 80 to $>100$ m over the region sampled, except along the outer portions of Transects 100–300 where it was about 67 m (Table 3). These changes were consistent with increased mixing of the upper water from atmospheric forcing as the fall to winter transition occurs.

By winter (August), the upper water column was composed of only Winter Water (WW), which is characterized by temperatures of $<-1^\circ C$ and salinities of 33.8 to 34.0 (Smith et al., 1999). This water mass replaced the AASW and was formed by the intense cooling that occurs during winter. MLDs were $>100$ m over the entire region (Table 4), which was consistent with intense atmospheric forcing (wind mixing) and convective cooling. The mixed layer in austral spring (November) showed the beginning of warming and the presence of lower salinity waters, as expected from melting sea ice. However,
Table 4. **August 1993. Ice covered.** Comparison of mean and standard deviation for mixed layer depths (m), integrated water column Chl $a$ biomass (m mol m$^{-2}$) and nutrient concentrations (m mol m$^{-2}$), as well as averaged ratios of nutrient concentrations, for the upper 100 m. Data are sorted by percent dominance by diatoms or phytoflagellates within phytoplankton assemblages surveyed between Transectss 300–600 along the west Antarctic Peninsula. Community composition data measured on Transect 200 was lost. Listed is the number of stations where the criteria for each subset is met. Not included are stations where the vertical sampling did not extend to 100 m. Community comparisons expressed as the ratio of diatoms: phytoflagellate are provided only if the difference between assemblages was significant at the >95% confidence level (t-test). ns, not significant; na, not applicable; —, not measured. **Diatom-dominated communities covered most of shelf region, with phytoflagellate-dominated communities located off the shelf and along the southern boundary of the study region.**

<table>
<thead>
<tr>
<th>Phytoplankton</th>
<th>Stations</th>
<th>MLD</th>
<th>$\Sigma$ Chl $a$</th>
<th>$\Sigma$ Si(OH)$_4$</th>
<th>$\Sigma$ NO$_3$</th>
<th>$\Sigma$ NH$_3$</th>
<th>$\Sigma$ PO$_4$</th>
<th>Si(OH)$_4$:NO$_3$</th>
<th>NO$_3$:PO$_4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>T$_{200}$, inner shelf</td>
<td>3</td>
<td>absent</td>
<td>—</td>
<td>6.36 ± 0.29</td>
<td>3.03 ± 0.03</td>
<td>0.051</td>
<td>0.235 ± 0.009</td>
<td>2.10 ± 0.12</td>
<td>12.9 ± 0.34</td>
</tr>
<tr>
<td>Diatoms $&gt;50%$</td>
<td>17</td>
<td>absent</td>
<td>15 ± 3</td>
<td>5.58 ± 1.0</td>
<td>2.71 ± 0.28</td>
<td>0.030 ± 0.021</td>
<td>0.189 ± 0.021</td>
<td>2.05 ± 0.27</td>
<td>14.6 ± 1.10</td>
</tr>
<tr>
<td>T$_{300}$, inner shelf $&gt;62%$</td>
<td>2</td>
<td>absent</td>
<td>14 ± 1</td>
<td>6.28 ± 0.28</td>
<td>2.89 ± 0.04</td>
<td>0.050</td>
<td>0.220 ± 0.003</td>
<td>2.17 ± 0.07</td>
<td>13.2 ± 0.78</td>
</tr>
<tr>
<td>T$_{400}$, outer shelf $&gt;55%$</td>
<td>4</td>
<td>absent</td>
<td>15 ± 2</td>
<td>5.16 ± 0.41</td>
<td>2.87 ± 0.04</td>
<td>0.021 ± 0.005</td>
<td>0.200 ± 0.009</td>
<td>1.80 ± 0.14</td>
<td>14.4 ± 0.49</td>
</tr>
<tr>
<td>T$_{500}$, center shelf $&gt;55%$</td>
<td>3</td>
<td>absent</td>
<td>16 ± 0</td>
<td>7.29 ± 0.23</td>
<td>2.89 ± 0.11</td>
<td>0.045 ± 0.015</td>
<td>0.185 ± 0.007</td>
<td>2.52 ± 0.11</td>
<td>15.7 ± 1.22</td>
</tr>
<tr>
<td>ratio T$<em>{300}$:T$</em>{500}$</td>
<td>ns</td>
<td>ns</td>
<td>0.86</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Phytoflag. $&gt;50%$</td>
<td>13</td>
<td>absent</td>
<td>12 ± 2</td>
<td>4.70 ± 0.9</td>
<td>2.77 ± 0.34</td>
<td>0.035 ± 0.026</td>
<td>0.188 ± 0.020</td>
<td>1.67 ± 0.17</td>
<td>14.7 ± 0.9</td>
</tr>
<tr>
<td>T$_{300}$, center shelf $&gt;60%$</td>
<td>2</td>
<td>absent</td>
<td>11 ± 1</td>
<td>5.70 ± 0.24</td>
<td>3.02 ± 0.04</td>
<td>0.148 ± 0.090</td>
<td>0.207 ± 0.003</td>
<td>1.89 ± 0.14</td>
<td>14.6 ± 0.37</td>
</tr>
<tr>
<td>T$_{300}$–400, offshore $&gt;60%$</td>
<td>3</td>
<td>absent</td>
<td>13 ± 3</td>
<td>4.89 ± 0.60</td>
<td>2.92 ± 0.07</td>
<td>0.045 ± 0.028</td>
<td>0.189 ± 0.016</td>
<td>1.67 ± 0.16</td>
<td>15.3 ± 1.08</td>
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<tr>
<td>T$_{600}$, offshore $&gt;55%$</td>
<td>2</td>
<td>absent</td>
<td>14 ± 2</td>
<td>2.99 ± 0.07</td>
<td>2.08 ± 0.14</td>
<td>0.050 ± 0.000</td>
<td>0.155 ± 0.002</td>
<td>1.44 ± 0.07</td>
<td>13.4 ± 0.73</td>
</tr>
</tbody>
</table>

**Community Comparisons**

| >50% Diatoms | ns | 1.25 | 1.23 | ns | ns | ns | 1.25 | ns |
| >50% Phytoflagellates | ns | 1.25 | 1.23 | ns | ns | ns | 1.25 | ns |

MLDs remained deep (Table 6), suggesting that seasonal heating of the upper water column has not started.

The average temperature and salinity values for the second austral summer (January 1994) showed that mixed-layer structure can vary widely from one summer to the next. In January 1994, the upper water column was colder and saltier than in the previous year, although the two summer cruises occurred at approximately the same time (Table 1). The colder temperatures and smaller range of temperature values suggests that seasonal warming was delayed in 1994 and the higher salinity implies less freshwater input. Observations show that sea ice retreat occurred later in the austral spring and summer of 1994 than in 1993 (Moline and Prézelin, 1996a; Lascara et al., 1999), which is consistent with the patterns seen in the average thermohaline properties of the mixed layer. In addition, MLDs were shallow (Table 5), suggesting that seasonal heating had started, but that the water had not been exposed to strong atmospheric forcing.

d. Percent diatom distributions in the upper 100 m

During the austral summer of 1993, diatom dominance was highest (>60%) in the upper 100 m along the outer and mid-shelf of Transect 300 (Fig. 5a), where UCDW was moving onto the shelf (Fig. 2b) and where Si(OH)$_4$:NO$_3$ ratios at 200 m were <1.6 (Fig. 3a). Regions with >50% diatoms were located on the outer part of all the other transects (Fig. 5a), a distribution that closely correlated with the distribution of UCDW on the WAP shelf (Fig. 2a) and with Si(OH)$_4$:NO$_3$ ratios of 1.6–1.8 (Fig. 3a). Just shoreward of these diatom-dominated waters, there was a band of modified UCDW water with 40–50% diatoms (Fig. 5a) and Si(OH)$_4$:NO$_3$ ratios at 200 m of 1.8–2.0. Areas with just 30–40% diatoms were confined to the east of Transect 400 and associated with modified UCDW with Si(OH)$_4$:NO$_3$ ratios at 200 m of 2.0–2.3. On inner parts of Transects 500–600 (Fig. 5), diatoms were <25% of assemblages, 200 m temperature was >1.4°C and Si(OH)$_4$:NO$_3$ ratios were >2.6.

In January 1993, the integrated chlorophyll $a$ in the upper 100 m in diatom-dominated regions was 30% and 50% less, respectively, than for phytoflagellate-dominated and chlorophyte-containing regions (Table 2). The integrated Si(OH)$_4$ concentration was lowest in the diatom-dominated regions. Removal of Si(OH)$_4$ by diatoms was reflected in 30–40% lower Si(OH)$_4$:NO$_3$ ratios than for non-diatom dominated regions. Integrated NO$_3$ and phosphate (PO$_4$) concentrations were similar in diatom-dominated and phytoflagellate-dominated regions and about 20% higher in regions with abundant chlorophytes (Table 2).

In the austral fall of 1993, integrated chlorophyll $a$ for all diatom-dominated regions was greater than those for areas dominated by phytoflagellates and chlorophytes and for diatom-dominated areas sampled during the austral summer two months earlier (Table 3). Integrated nitrate concentrations in the upper 100 m were higher in March–May than in January 1993 (Tables 2, 3). By comparison, Si(OH)$_4$ and Si(OH)$_4$:NO$_3$ ratios were lower in diatom-dominated regions and NO$_3$:PO$_4$ ratios everywhere were generally higher than
Table 5. January–February 1994. Comparison of mean and standard deviation for mixed layer depths (m), integrated water column Chl $a$ biomass (m mol m$^{-2}$) and nutrient concentrations (m mol m$^{-2}$), as well as averaged ratios of nutrient concentrations, for the upper 100 m. Data are sorted by percent dominance phytoflagellates within phytoplankton assemblages surveyed between Transects 300–600 along the west Antarctic Peninsula. Listed is the number of stations where the criteria for each subset is met. Not included are stations where the vertical sampling did not extend to 100 m. Water column characteristics are also summarized for dinoflagellates at stations where they comprised $>15\%$ of the integrated Chl $a$ biomass. High biologically processed waters with shallow MLDs contained no diatom-dominated communities; phytoflagellate-dominated communities accounted for $75\%$ of water columns surveyed; a nutrient-dependent succession from phytoflagellates to dinoflagellates appeared evident.

<table>
<thead>
<tr>
<th>Phytoplankton</th>
<th>Stations</th>
<th>MLD</th>
<th>$\Sigma$ Chl $a$</th>
<th>$\Sigma$ Si(OH)$_4$</th>
<th>$\Sigma$ NO$_3$</th>
<th>$\Sigma$ NH$_3$</th>
<th>$\Sigma$ PO$_4$</th>
<th>Si(OH)$_4$:NO$_3$</th>
<th>NO$_3$:PO$_4$</th>
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</thead>
<tbody>
<tr>
<td>Phytoflag. $&gt;50%$</td>
<td>20</td>
<td>$57 \pm 9$</td>
<td>$34 \pm 9$</td>
<td>$4.53 \pm 0.56$</td>
<td>$2.53 \pm 0.29$</td>
<td>$0.185 \pm 0.064$</td>
<td>$0.170 \pm 0.022$</td>
<td>$1.80 \pm 0.22$</td>
<td>$15.0 \pm 1.9$</td>
</tr>
<tr>
<td>Phytoflag. $&gt;60%$</td>
<td>17</td>
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<td>$34 \pm 10$</td>
<td>$4.47 \pm 0.58$</td>
<td>$2.52 \pm 0.31$</td>
<td>$0.191 \pm 0.064$</td>
<td>$0.170 \pm 0.024$</td>
<td>$1.79 \pm 0.24$</td>
<td>$15.0 \pm 2.1$</td>
</tr>
<tr>
<td>Phytoflag. $&gt;80%$</td>
<td>8</td>
<td>$57 \pm 10$</td>
<td>$37 \pm 6$</td>
<td>$4.60 \pm 0.56$</td>
<td>$2.58 \pm 0.28$</td>
<td>$0.187 \pm 0.030$</td>
<td>$0.177 \pm 0.027$</td>
<td>$1.81 \pm 0.27$</td>
<td>$14.8 \pm 2.2$</td>
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<td>Dinoflag. 20–35%</td>
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</tr>
<tr>
<td>T$_{300}$, outer shelf</td>
<td>4</td>
<td>$51 \pm 3$</td>
<td>$31 \pm 12$</td>
<td>$4.78 \pm 0.34$</td>
<td>$2.66 \pm 0.16$</td>
<td>$0.234 \pm 0.020$</td>
<td>$0.167 \pm 0.010$</td>
<td>$1.81 \pm 0.20$</td>
<td>$16.0 \pm 1.7$</td>
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<tr>
<td>T$_{500–600}$, outer shelf</td>
<td>4</td>
<td>$44 \pm 10$</td>
<td>$33 \pm 6$</td>
<td>$4.38 \pm 0.64$</td>
<td>$2.74 \pm 0.33$</td>
<td>$0.155 \pm 0.074$</td>
<td>$0.187 \pm 0.020$</td>
<td>$1.60 \pm 0.17$</td>
<td>$14.6 \pm 0.3$</td>
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<tr>
<td>Diatoms 35–45%</td>
<td>3</td>
<td>$68 \pm 4$</td>
<td>$37 \pm 7$</td>
<td>$4.75 \pm 0.54$</td>
<td>$2.78 \pm 0.10$</td>
<td>$0.278 \pm 0.010$</td>
<td>$0.176 \pm 0.023$</td>
<td>$1.71 \pm 0.25$</td>
<td>$15.8 \pm 1.5$</td>
</tr>
<tr>
<td>Diatoms &lt;15%</td>
<td>9</td>
<td>$56 \pm 10$</td>
<td>$37 \pm 9$</td>
<td>$4.48 \pm 0.61$</td>
<td>$2.46 \pm 0.38$</td>
<td>$0.200 \pm 0.041$</td>
<td>$0.177 \pm 0.023$</td>
<td>$1.84 \pm 0.27$</td>
<td>$13.9 \pm 1.5$</td>
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</tbody>
</table>
Figure 5. Comparison of the horizontal distribution of the percent diatoms within phytoplankton assemblages that were vertically integrated to 100 m for (a) January–February 1993, (b) March–May 1993, (c) August–September 1993, (d) January–February 1994, and (e) November 1991. Filled circles indicate the distribution of the sampling sites. The progression from darkest to lightest shaded areas indicates diatom percent contributions of >70%, 60 to 70%, 50 to 60%, 30–50% and <30% respectively. (f) Bathymetry (2000 m, 1000 m, 500 m) of the study region.
those observed in the austral summer. In March–May 1993, several groupings of diatom-dominated assemblages were identifiable along 900 km of the WAP (Fig. 5b). At the center of each grouping, diatoms accounted for >70% of the water column chlorophyll a. One grouping was found along inner Transect 200, in proximity to Marguerite Trough near Adelaide Island, and overlaid modified UCDW water that had relatively high Si(OH)₄:NO₃ ratios of 2.6–2.8 (Table 3). There were indications of another patch intruding from the south on to the outer shelf of Transect 000. Two additional diatom patches encompassed most of the shelf between Transects 500–600 that coincided with modified UCDW (1.3–1.4°C); with Si(OH)₄:NO₃ ratios of 2.1–2.3 on outer Transect 500 and 2.4–2.5 on inner Transect 600. The chlorophyll a of these patches was about three times as high as concentrations observed in patches on Transects 000–200 (Table 3). A fifth diatom patch was detected along the continental shelf break between Transects 700–800 and appeared to extend across the shelf to the Gerlache Strait (Fig. 5b). Diatom dominance was evident all along Transect 800 and coincided with the lowest Si(OH)₄:NO₃ ratios observed (<1.9). The mid- to inshore shelf portions of Transect 800 were covered with deep waters <1.2°C and Si(OH)₄:NO₃ ratios at 200 m were 2.2–2.5. The coldest deep water, with Si(OH)₄:NO₃ ratios >2.6 and no diatom-dominated communities, was found at inner shelf of Transect 800, among the many islands present in this region.

By late winter, in August 1993, diatom-dominated communities were uniformly spread over most of the WAP continental shelf between Transects 400–600 and innermost Transect 300 (Fig. 5c). The highest percentage of diatoms (approaching 60%) was located over modified UCDW near the outer edge Transect 400 (Fig. 2c), with Si(OH)₄:NO₃ ratios of 2.2–1.8 (Fig. 3c), and the innermost stations of Transect 300 (Fig. 2c) near Marguerite Trough, with Si(OH)₄:NO₃ ratios of 2.2–2.0 (Fig. 3c). The integrated chlorophyll a in the upper 100 m in August 1993 was low in all areas (Table 4). Integrated silicate concentrations were ca 25% higher in diatom-dominated regions, while concentrations of nitrate and phosphate were similar for all regions. Thus, Si(OH)₄:NO₃ ratios were elevated in diatom-dominated regions in late winter while NO₃:PO₄ ratios were similar for diatom and phytoflagellate dominated areas (Table 4).

By austral summer of January 1994, there were no diatom-dominated communities in the study region (Fig. 5d). Chlorophyll a in the upper 100 m was dominated by phytoflagellates. Communities with >40% diatoms were located at the outer Transect 500, coincident with UCDW and Si(OH)₄:NO₃ ratios <1.8. The vertical distribution of phytoplankton measured in January 1994 (not shown) indicated that diatom-dominated communities were located at depths >100 m at the inner station of Transect 500 (>60% diatoms) and >80 m at mid-shelf on Transect 600 (100% diatoms) with ammonium (NH₃) significantly higher than elsewhere (Table 5). High ammonium may be indicative of a post bloom sedimentation or lateral advection of diatom particles from an earlier UCDW event. It is also indicative of the additional inorganic nitrogen load possible within a recycling diatom ‘seed’ community.

During the austral spring cruise in November 1991 (Fig. 5e), the marginal ice zone
(MIZ) extended to the outer continental shelf between Transects 500–700. Diatoms were not prevalent in the MIZ but abundant in open waters of Dallmann Bay where a subsurface diatom bloom was observed at a MLD of about 67 m (Table 6). The diatom bloom was physically isolated from the modified UCDW evident in the outer shelf regions (Fig. 2e).

e. Phytoplankton community composition

In January 1993, diatoms dominated the outer shelf waters of Transects 200–600 and extended into the mid-shelf of Transects 200–300 (Fig. 6a). Diatoms also dominated communities below the fresher water lens (FWL) that formed on the inner part of Transect 400 from glacial melt. In the FWL, dinoflagellates were prevalent. In diatom-dominated communities, phytoflagellates were the second most abundant phytoplankton group and visa versa. Phytoflagellate-dominated communities extended from inner Transects 200–300 through the mid-shelf of Transect 400 and throughout inner and mid-shelf regions of Transect 600. Phytoflagellates did not dominate similar regions of Transect 500 due to abundant chlorophytes. In January 1993, bands of concentrated chlorophytes were located coincident with modified UCDW at the leading edge of UCDW, e.g. from inner Transect 200 to mid-shelf Transect 400 and inshore of a region of UCDW upwelling on Transect 500 (Fig. 6a). Other conditions also lead to abundant chlorophytes. In the cold waters of Marguerite Bay (Fig. 5f), chlorophytes were ca 30% of water column chlorophyll a.

In March–May of 1993 (Fig. 6b), diatom-dominated communities covered most of the WAP shelf and would have dominated several adjacent sites, except for the high abundance of chlorophytes. Several groupings of diatom-dominated communities were present (Fig. 5b), some separated from each other by phytoflagellate-dominated communities located along the inner and outer portions of most transects (Fig. 6b). Each diatom group seemed to be associated with different UCDW intrusion events. There were confl uences of more than one diatom-dominated group (Fig. 5b) where distinctions were evident only in comparisons of hydrographic (Fig. 2), nutrient (Table 3) and physiological parameters (see below). Chlorophytes were abundant in what appeared to be an across-shelf band of patches that were associated with modified UCDW and UCDW intrusions.

Chlorophytes were also abundant in areas not apparently influenced by UCDW. On the inner shelf of Transect 000 in austral fall 1993, chlorophytes accounted >35% (Fig. 5b) of chlorophyll a in an area where the Si(OH)₄:NO₃ ratios were >3.0 and NO₃:PO₄ ratios were ca. 13 (Table 3). The MLD was ca 60 m and yet there was no indication of UCDW at 200 m where water temperature was 1.2°C. Adjacent diatom-dominated communities, also with shallow MLDs, were almost as enriched in nutrients and contained somewhat higher levels of chlorophyll a, but no chlorophytes.

In March–May 1993, diatom groupings 2 and 3 (Figs. 5b and 6b) appear to have resulted from separate UCDW events, most likely two intrusions that moved across the shelf via Marguerite Trough. The Si(OH)₄:NO₃ ratios of ca. 2.4–2.5 and NO₃:PO₄ ratios approaching 15 (Table 3) indicate that biological processing of the water has occurred. The lower Si(OH)₄:NO₃ ratios of 2.2, and higher chlorophyll a concentrations suggest that diatom
<table>
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<th>Phytoplankton</th>
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<th>MLD</th>
<th>Chl a</th>
<th>Si(OH)&lt;sub&gt;4&lt;/sub&gt;</th>
<th>NO&lt;sub&gt;3&lt;/sub&gt;</th>
<th>NH&lt;sub&gt;3&lt;/sub&gt;</th>
<th>PO&lt;sub&gt;4&lt;/sub&gt;</th>
<th>Si(OH)&lt;sub&gt;4&lt;/sub&gt;/NO&lt;sub&gt;3&lt;/sub&gt;</th>
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<td><strong>Diatoms &gt; 50%</strong></td>
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<td>T&lt;sub&gt;700&lt;/sub&gt; spring bloom &gt;65%</td>
<td>2</td>
<td>67 ± 1</td>
<td>44 ± 4</td>
<td>4.49 ± 0.30</td>
<td>2.14 ± 0.16</td>
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<td>.174 ± .003</td>
<td>2.10 ± 0.02</td>
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<td>13 ± 3</td>
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<td>1.80 ± 0.04</td>
<td>—</td>
<td>.140 ± .001</td>
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<tr>
<td>marginal ice zone &gt;50%</td>
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<td>28 ± 12</td>
<td>4.06 ± 0.55</td>
<td>2.03 ± 0.27</td>
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<td>.176 ± .032</td>
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<td>12 ± 1</td>
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</tr>
<tr>
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<td>2</td>
<td>absent</td>
<td>18 ± 3</td>
<td>4.70 ± 0.34</td>
<td>2.19 ± 0.02</td>
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<td>Diatom bloom: Gerlache diatoms</td>
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<td>1.19</td>
<td>—</td>
<td>1.24</td>
<td>1.80</td>
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<tr>
<td>Diatom bloom: MIZ phytoflag.</td>
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<td>1.57</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
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Table 6. **November 1991.** Comparison of mean and standard deviation for mixed layer depths (m), integrated water column Chl a biomass (mg m<sup>-2</sup>) and nutrient concentrations (m mol m<sup>-2</sup>), as well as averaged ratios of nutrient concentrations, for the upper 100 m. Data are sorted by percent dominance by diatoms or phytoflagellates within phytoplankton assemblages surveyed between Transects 500–700 along the west Antarctic Peninsula. Listed is the number of stations where the criteria for each subset is met. Not included are stations where the vertical sampling did not extend to 100 m. Water column characteristics are also summarized for chlorophytes and dinoflagellates at stations where they comprised >15% of the integrated Chl a biomass. The ratio of diatoms: phytoflagellate and diatom:chlorophyte water column characteristics are provided only if the difference between assemblages were significant at the >95% confidence level (t-test). ns, not significant; na, not applicable; —, not measured. A diatom spring bloom in ice-free waters of Dahlmann Bay; phytoflagellates dominate marginal ice zone located very near the continental shelf break.
group 4 on the outer shelf of Transect 500 resulted from a later stage UCDW intrusion event. Diatom group 5, inshore on Transect 500, had hydrographic, nutrient, and phytoplankton community characteristics similar to group 4, but different photosynthetic physiologies that suggest a younger ‘stage’ (see below). Diatom group 6 resulted from a large UCDW event that influenced the entire across-shelf region along Transect 800 as well as the outer shelf portions of Transects 700–900 (Fig. 6b). The variability in the nutrient concentrations associated with diatom group 6 is sufficient to warrant further subdivision or to regard this as a large UCDW event in which the inshore waters were at a later stage (‘aged’) than outer shelf waters. Diatom groups 5 and 6 were separated from each other by a band of chlorophytes on Transect 700. Inshore, these diatom-dominated regions were constrained by phytoflagellate-dominated communities in the colder waters of the Gerlache Strait outflow (Fig. 2), which was as enriched in inorganic macronutrients as those of diatom-dominated waters (Table 3). Thus, differences in the structure of these phytoplankton communities suggested that UCDW provided necessary diatom growth requirements and that the Gerlache Strait outflow did not support the development of diatom communities.

In the austral spring of August–September 1993, when much of the shelf was covered by sea ice, diatoms dominated communities on the shelf over most of Transects 400–600 (Fig. 6c). Phytoflagellate-dominated communities were found in the southern part of the study region, where the sea ice was thicker along Transect 300, along the entire outer shelf and in the outflow of the Gerlache Strait. Other phytoplankton taxa were largely absent. On average, Si(OH)₄:NO₃ ratios and chlorophyll a concentrations were ca 25% lower in phytoflagellate communities compared with diatom communities (Table 3). Silicate concentrations of outer shelf phytoflagellate communities were ca half those of onshore phytoflagellate communities. While appearing rather homogeneous, diatoms on Transect 400 were in waters where Si(OH)₄:NO₃ ratios at 200 m were <2.0 compared to values ca 2.4 on Transects 500–600. The UCDW intrusion on Transect 300 was associated with reduced prevalence of diatoms (Fig. 5c) but decreased Si(OH)₄:NO₃ ratios, indicating that this may be a relatively new event.

In the following austral summer of January 1994, phytoflagellates dominated the phytoplankton community over the entire study region, often in association with increased
abundances of dinoflagellates in stratified near-surface waters (Fig. 6d). No diatom-dominated communities were evident. No significant concentrations of chlorophytes were present (Fig. 6d). In contrast to the nutrient-rich conditions observed in January 1993, water column nutrients were severely depleted in January 1994, which provides further evidence that UCDW intrusions are not seasonal and that intervals between UCDW events can, at times, be long enough for significant phytoplankton community succession to occur over large portions of the WAP continental shelf.

The only other comparison between ice-covered sea versus non-ice-covered sea regions occurred in the austral spring of 1991, when sea ice covered Transects 500–600 and the inner Transect 700 (Dallmann Bay) was largely ice free. The phytoplankton communities in the MIZ (Fig. 5e) were heavily concentrated in near-surface waters and completely dominated by phytoflagellates, especially *Phaeocystis* spp. A major diatom bloom dominated the subsurface waters of the ice-free regions of Dallmann Bay (Fig. 5d). There was a small intrusion of UCDW at the outer shelf of Transect 700, although it is not thought to be the cause of the diatom bloom. Likewise, the diatoms in Gerlache Strait were in nutrient-poorer water than those associated with the bloom in Dallmann Bay (Table 6). Chlorophytes were found only at depth on Transects 500–600.

**f. Phytoplankton community succession and nutrient ratios**

The distributions of diatom dominance (Fig. 5) and the phytoplankton community composition (Fig. 6) obtained from the five cruises give the impression of succession of the phytoplankton communities. The diatom-dominated communities tend to be associated with regions where UCDW has moved onto the WAP shelf and other taxa dominate outside these areas. A change in dominance may also arise because of sudden advective changes or slower progressive changes in nutrient concentration or changes in nutrient ratios (Moline and Prézelin, 1996a). For example, in the intervals between UCDW intrusion events, when nutrients were not mixed or upwelled into the upper water column, phytoflagellates became increasingly dominant as the surface waters ‘aged’ and nutrients were depleted. Thus, changes in phytoplankton community composition observed during the different cruises were examined within the context of the Si(OH)₄:NO₃ ratio and NO₃:PO₄ ratios (Fig. 7). These ratios are indicative of the water mass present and provide an approach for identifying nutritional regimes that favor certain phytoplankton groups and discourage others.

The March–May 1993 phytoplankton community assemblages were highly variable (cf. Figs. 5b, 6b), creating a mosaic of discrete communities associated with the largest range of Si(OH)₄:NO₃ ratios measured during all five cruises (Fig. 7b). Diatom communities associated with Si(OH)₄:NO₃ ratios of 3.0–3.5 were in newly intruded UCDW and were in the early phases of growth, e.g. communities associated with Marguerite Trough. Those associated with low Si(OH)₄:NO₃ ratios (1.8–2.0) were actively growing, e.g. the northern part of the study region near Transect 800.

In August 1993, the range of Si(OH)₄:NO₃ and NO₃:PO₄ ratios was smaller and the
transition from high to low Si(OH)$_4$:NO$_3$ ratios was associated with the change from diatom to phytoflagellate-dominated communities. In austral spring (November 1991), the lowest NO$_3$:PO$_4$ ratios of all cruises were observed with a bloom of *Phaeocystis* spp. in the MIZ. The large diatom bloom observed in open water was associated with low Si(OH)$_4$: NO$_3$ ratios.
Figure 8.
Comparison of nutrient ratios from the two summers (Fig. 7a) shows differences that result from dominance by diatoms versus phytoflagellates. Diatoms became dominant when UCDW intrusions replenished macronutrients and perhaps necessary trace metals. Diatoms lower the initially high Si(OH)₄:NO₃ ratios (>3.0) by preferential uptake of abundant Si(OH)₄. At intermediate Si(OH)₄:NO₃ ratios (2.5 to 2.0) and NO₃:PO₄ ratios, diatoms and phytoflagellates dominance overlap with phytoflagellates becoming dominant at higher NO₃:PO₄ ratios. Dominance by phytoflagellates occurred when NO₃:PO₄ ratios were high and Si(OH)₄:NO₃ ratios were <2.5.

g. Primary production and growth physiology

Comparisons of the distributions of potential volumetric primary production, integrated chlorophyll a, and Chl-specific rates of primary production, with each overlaid on a background showing the percent diatom distribution, are provided in Figures 8–10, respectively. A comparison of the photosynthetic characteristics of diatom-dominated and phytoflagellate-dominated communities for each cruise is provided in Table 7. Average daytime rates of volumetric production should be considered a zero-order estimate, as temporal corrections for in situ diurnal periodicity of photosynthesis were simplified with the use of a standard curve. Potential-specific rates of primary production (P_MAX/Chl a) are considered a proxy for growth, assuming light-saturated conditions in the upper 60 m of the water column. Particulate organic carbon (POC) concentrations were used in estimates of carbon turnover times.

In January 1993, volumetric production was greatest in the UCDW transition waters between diatom-dominated and phytoflagellate-dominant communities, evident as an across-shelf band of production from near-shore on Transect 300 to off shelf on Transect 600 (Fig. 8a). Compared to phytoflagellate-dominated water columns on a regional scale, diatom-dominated water columns appeared 40% less productive (Fig. 8a), had 50% less Chl a (Fig. 9a), 20% less POC and the same abundance of particulate organic nitrogen (PON) (data not shown). Diatom communities had a higher light requirement for light-saturated photosynthesis (Iₓ) and lower relative photosynthetic quantum efficiency (alpha/Chl) than phytoflagellate communities. The carbon-to-nitrogen (C:N, wt:wt) ratios averaged 5.5 ± 0.5 (n = 14) for diatom-dominated water columns and 6.4 ± 1.6 (n = 11) for
phytoflagellate-dominated water columns. The POC-based turnover times were estimated to be almost three times as fast for diatom-dominated communities as for phytoflagellate-dominated communities e.g. 1.71/10060.88 days and 4.87/10061.02 days, respectively.

In January 1993, the Chl-specific assimilation rates were 3m gC m g Chl a h l a 1 hour 1 in diatom-dominated communities along the outer shelf of Transects 200–400 and across the shelf of Transect 400 (Fig. 10a). The highest assimilation rate for diatoms (4.1 mg C mg Chl a h 1 hour 1) was estimated for low Chl a waters over the UCDW intrusion at the shelf break on Transect 300 (Fig. 2a). Here, the Iks were 106 12 Ein m 2 s 1 and the water column C:N ratios averaged.ca. 5.2. High biomass communities of diatoms (Fig. 9a), located over modified UCDW mid-shelf on Transect 300 (Fig. 2a), had similar C:N ratios but lower assimilation rates (2.5 ± 0.3 mg C mg Chl a h 1 hour 1) (Fig. 10a) and Iks (78 ± 18 µEin m 2 s 1).

Within phytoflagellate communities, PMAX was >4 mg C mg Chl a h 1 hour 1 (Fig. 10a) at two locations in modified UCDW (Fig. 2a) where diatoms accounted for >40% of the community (Fig. 5a). One site was a localized region on the inner shelf of Transect 400 (Fig. 10a) at the frontal boundary of the cold coastal current waters (Fig. 2a). The other was mid-shelf on Transect 600. Communities with >70% phytoflagellate had the lowest photosynthetic potential, averaging 1.2 mg C mg Chl a h 1 hour 1 (Fig. 10a), and Iks (46 µEin m 2 s 1). These observations suggest that phytoflagellates were low-light adapted

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<tr>
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<td>1.42</td>
<td>No significant difference</td>
<td>No significant difference</td>
<td>1.37</td>
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</tbody>
</table>

*—represents diatom bloom in ice free Dallmann Bay compared to prymnesiophytes located in MIZ near the continental shelf.

**—note that diatom measurements were made earlier in the cruise, when days were longer, than phytoflagellates, e.g. diatoms in the northern section and phytoflagellates in the southern section of the study area.

Table 7. Average photosynthetic characteristics of diatom-dominated communities expressed as a ratio of those determined to phytoflagellate-dominated communities during four cruises on the continental shelf of the west Antarctic Peninsula. January 1994 data are not included as there were no diatom-dominated communities in the sampling region at that time. Values are based upon integrated or integrated averages of biomass and productivity, respectively, for the upper 60 m for all sampling stations given in Figure 7.
Figure 9. Comparison of total Chlorophyll a biomass from 0 to 60 m (circles), relative to the distribution of diatom-dominated communities (shown as background, see Fig. 5), for (a) January–February 1993, (b) March–May 1993, (c) August–September 1993, (d) January–February 1994, and (e) November 1991.
Figure 10. Comparison of average Chl-specific primary production for the upper 60 m (circles), relative to the distribution of diatom-dominated communities (shown as background, see Fig. 5), for (a) January–February 1993, (b) March–May 1993, (c) August–September 1993, (d) January–February 1994, and (e) November 1991.
and that much of the chlorophyll $a$ accumulation at this site was not the result of cell growth but of pigment-dependent photoadaptation of slower growing cells. Since recently intruded waters were mixing in this region, it is not possible to use nutrient concentrations or C:N values to estimate turnover times. However, the $>60\%$ phytoflagellate communities at a greater distance from the new intrusion event routinely had C:N ratios of 6–8, values consistent with the view that these communities did not have a high growth potential prior to the event. If true, then new production would be much lower in phytoflagellate-dominated regions than in diatom-dominated regions than estimates based on chlorophyll $a$ and light penetration alone would predict (Moline and Prézelin, 2000).

From March to May of 1993, daylength shortened from 11 to 7.8 hours (Table 1). Production measurements were made only at stations the ship occupied during mid-day hours (Fig. 10b) POC and PON measurements were not made on this cruise. Diatoms dominated most water columns sampled in the northern region (Transects 500–900) and about half those sampled in the southern region (Transects 000–400) (Fig. 6). Chlorophyll $a$ was several fold higher in the northern region than the southern region (Fig. 9b). $I_{45}$ were uniformly higher in the northern ($75 \pm 13 \mu\text{Ein m}^{-2} \text{s}^{-1}$) than in the southern region ($52 \pm 11 \mu\text{Ein m}^{-2} \text{s}^{-1}$), perhaps a reflection on low-light photoadaptation during the season of declining solar radiation. Compared to phytoflagellate-dominated communities, diatom-dominated water columns had, on average, more than twice the chlorophyll $a$ and mid-day volumetric production rates (Table 7). Diatoms clearly accounted for most of the primary production of the WAP continental shelf in the fall of 1993. Chl-specific $P_{\text{MAX}}$ was $>5 \text{mg C mg Chl } a^{-1} \text{hour}^{-1}$ in diatom groupings 1, 5 and 6 (Fig. 10b), indicating enhanced rates of Chl-specific production was supported in WAP continental shelf waters throughout the austral fall and that it was unlikely that primary production was nutrient limited in the diatom patches sampled. Given its higher nutrient concentrations, higher $\text{Si(OH)}_4: \text{NO}_3$ ratios and lower $\text{NO}_3: \text{PO}_4$ ratio (Table 3), diatom group 1 appeared at an earlier stage of an UCDW event or ‘younger’ than diatom groupings 5 and 6. Other areas had estimated assimilation rates of 3–5 $\text{mg C mg Chl } a^{-1} \text{hour}^{-1}$, comparable to rates observed in diatom-dominated communities two months earlier in the austral summer. UCDW intrusions in the fall may play a key role in determining the amount and duration of enhanced primary production as winter approaches.

At the end of winter, in August 1993, chlorophyll $a$ (Fig. 9c), volumetric production (Fig. 8c) and assimilation rates (Fig. 10c) were uniformly low over the entire study region. The presence of an UCDW intrusion along Transect 300 was not coincident with enhanced primary production. Given that the MLD, nutrient availability and nutrient ratios in waters above the UCDW on Transect 300 were identical to those on Transect 400 (Table 4), it appeared the nutrient-rich UCDW water had not mixed upward significantly, perhaps because the presence of pack ice dampened mixing processes. Examination of vertical profiles (not shown) indicated that the identification of phytoflagellate-dominant communities in the upper 100 m was due primarily to their very high biomass in near-surface waters under the pack ice. Diatoms, while less abundant, dominated deeper phytoplankton
communities. The presence of the warmer UCDW may be linked to the release of phytoplankton from the pack ice and, in this case, those ice communities were dominated by phytoflagellates (*Phaeocystis* spp.). If true, UCDW intrusions may also have a previously unrecognized role in the initiation of spring phytoplankton growth.

By the following summer, in January 1994, phytoflagellates dominated all water columns sampled and dinoflagellates had become a significant component of near-surface phytoplankton communities (Fig. 6). High chlorophyll *a* accumulated on the inner part of Transect 600, the outer part of Transect 400, and in Marguerite Bay on Transect 300 (Fig. 9d). There were no UCDW intrusions. With the limited production data available (Fig. 8d, 10d), it appeared that production on Transect 600 was similar to that observed the prior summer when phytoflagellates were also present in this region of the study area.

Due to strong westerly winds, the pack ice was compressed during the November 1991 and the edge of the MIZ was especially sharp. Compared to phytoflagellate-dominated communities in the surface waters of the MIZ, the diatom-dominated communities located below the MIZ had ca 50% more chlorophyll *a* (Fig. 9e) and volumetric production (Fig. 8e). The diatom bloom in the open waters of Dallmann Bay had twice the POC concentrations as MIZ phytoplankton. The C:N ratio of the bloom approached 10 while the other two areas had C:N ratios of 6–7 (data not shown). The lowest C:N ratio of 4.3 was measured at the outer edge of Transect 700, which was influence by UCDW. Chl-specific rates of primary production (Fig. 10e) exceeded 5 mg C mg Chl *a*⁻¹ hour⁻¹ in the ice-free diatom bloom, the MIZ where phytoflagellates dominated, and at the outer edge of Transect 700. Chl-specific rates of production were lowest, e.g. 2–3 mg C mg Chl *a*⁻¹ hour⁻¹, in diatom communities located in deep water under the ice on the inner Transect 600.

4. Discussion

a. UCDW intrusion development

The hydrographic and phytoplankton community distributions observed during the four cruises in 1993 and 1994 showed patterns and progressions that are consistent with episodic forcing that resets the WAP continental shelf at irregular intervals. Periods of no or minimal intrusion effect, such as January 1994, were characterized by average Si(OH)₄:NO₃ ratios, high NO₃:PO₄ ratios, and dominance by phytoflagellates. In contrast, January 1993 was characterized by a new intrusion of UCDW along the outer WAP shelf which resulted in diatom-dominated phytoplankton communities. The effect of forcing by multiple intrusions, in differing stages of development, was represented by the diverse biological structure seen in March–May 1993. The August 1993 period was representative of an early stage intrusion. November 1991 provided a more limited characterization of the WAP during a time characterized by reduced or no UCDW intrusions. This time perhaps best illustrates the effect of the MIZ on phytoplankton community composition.

The space and time coverage of the WAP continental shelf afforded by the 1993 and
1994 observations allowed characterization of the different stages associated with UCDW intrusions. New UCDW intrusions were confined to the outer shelf and were associated with temperatures of 1.8°C to 1.5°C at 200 m, Si(OH)$_4$:NO$_3$ ratios of 2.8 to 3.5 at 200 m, NO$_3$:PO$_4$ ratios of 10 to 12, and low phytoplankton biomass with subsurface diatom communities beginning to develop. Once on the WAP continental shelf, the UCDW intrusion affected the phytoplankton community structure of the upper water column only if it upwelled and/or mixed upward and exchanged with the surface waters. The processes that allow mixing of UCDW include wind and topographic effects as well as diffusion and mixing across the permanent pycnocline (Smith et al., 1999; Smith and Klinck, 2002). The latter may be the dominant process during times when sea ice is present, such as in the winter (Smith and Klinck, 2002).

The start of an intrusion event was indicated by onshelf movement of the temperature maximum, decreasing Si(OH)$_4$:NO$_3$ ratios, increasing NO$_3$:PO$_4$ ratios, increasing diatom dominance, and a potential increase in Chlorophyll $a$. The presence of a well-developed UCDW intrusion on the WAP shelf was identified by movement of the temperature maximum of 1.6°C to 1.4°C to mid-shelf, Si(OH)$_4$:NO$_3$ ratios <2.0 at 200 m, NO$_3$:PO$_4$ ratios of 12 to 14, and decreased diatom dominance with dominance by photoflagellates and other phytoplankton taxa increasing. The final stages of an old or waning UCDW intrusion occurred when the intruded water mass became stranded on the WAP shelf and was characterized by water temperatures of 1.35 to 1.5°C, Si(OH)$_4$:NO$_3$ ratios that increased above 2.0, decreased NO$_3$:PO$_4$ ratios, and decreased diatom dominance with increased dominance by other phytoplankton taxa.

Areas that were not affected by intrusions of UCDW were characterized by temperatures <1.3°C at 200 m, variable nutrient ratios, variable nutrient concentrations, and mixed phytoplankton community assemblages. These conditions occurred in the areas affected by outflow from the Gerlache Strait and along the inner shelf where there is a southward flowing coastal current. Semi-enclosed bays, such as Dallmann Bay, are similar but may be dominated by diatoms or other taxa.

b. Circulation patterns and phytoplankton communities

A recurring pattern seen in the distribution of the temperature maximum below 200 m was onshelf intrusion of UCDW, which began at the shelf break in response to variability in the southern boundary of the ACC. The southern boundary of the ACC, determined by the location of the 1.8°C isotherm, was seen along the outer shelf in all of the times sampled (Fig. 2). The Antarctic Slope Front is absent along the WAP (Jacobs, 1991; Whitworth et al., 1998), which allows the southern boundary of the ACC to flow along the shelf break, thereby, allowing the oceanic water mass, UCDW, to intrude onto the continental shelf.

However, onshelf movement of UCDW did not occur all along the shelf break, rather it occurred at the outer end of Transects 200–300 (Figs. 2b–d) and Transects 600–700 (Figs. 2a–d). These are locations where the curvature of the shelf break is sufficient to allow
onshore movement of UCDW (Dinniman and Klinck, 2004). It is in these regions that diatoms dominated the phytoplankton community structure (Fig. 7) and Si(OH)₄:NO₃ ratios were reduced (Fig. 3).

The intrusions moved onshelf and in some areas extended into the inner shelf region via deep across-shelf troughs that connect the inner and outer shelf regions. This across-shelf movement supports regions of enhanced diatom abundance, such as the one observed in March–May 1993 along the inner portion of Transect 200 (Fig. 4B). Transect 200 crosses Marguerite Trough, which allows UCDW to move from the outer shelf into Marguerite Bay (Klinck et al., 2004). Similarly, the high diatom abundances along the mid-shelf of Transect 600 (Fig. 4b, c) may be sustained by across-shelf movement of UCDW via the trough that extends from the shelf break towards Anvers Island.

The boundary between newly intruded UCDW and modified UCDW that covers the shelf below the permanent pycnocline is variable because of the episodic nature of the onshelf intrusions. This boundary moves further inshore as an intrusion event occurs. The phytoplankton community structure associated with this boundary is distinct and may be characterized by the presence of chlorophyll b, no to low diatoms biomass, and dominance by phytoflagellates (Fig. 5). Thus, this phytoplankton assemblage represents a transitional community between oceanic and shelf populations.

The inner portion of the WAP is characterized by a southerly-flowing coastal current, which is characterized by low salinity and temperature, which produces a frontal boundary with shelf waters (Smith et al., 1999). This inner shelf current was best resolved during the March–May 1993 cruise (Fig. 2b). The phytoplankton assemblages associated with the inner shelf current were mixed, with no pattern of consistent dominance by any group (Figs. 5, 6).

Temperature and phytoplankton observations from the nearby vicinity of Anvers Island (inner-most part Transect 600) tended to be anomalous relative to adjacent observations. This area is affected by outflow from the Gerlache Strait and the phytoplankton communities likely result from different environmental conditions. Similarly, the large diatom-dominated bloom observed in November 1991 in Dallmann Bay along the inner part of Transect 700 may be representative of an environment with special conditions.

c. Temporal variations

An important conclusion arising from the comparisons of the hydrographic and phytoplankton distributions observed in 1993 and 1994 is that phytoplankton community structure on the WAP is seasonally independent. What is observed on this shelf arises from episodic inputs of UCDW, the frequency of which is determined by variability in the southern boundary of the ACC relative to the WAP shelf break. In turn, variability in this boundary is driven by larger scale climate-related processes rather than local seasonal forcing.

The temperature and phytoplankton distributions from January 1994 showed no UCDW intrusions on the WAP shelf, mixed phytoplankton assemblages, and average nutrient
ratios. These distributions likely represent a baseline state for the WAP continental shelf. It should be noted that these conditions were observed in the austral summer when conditions are optimal for biological production. Thus, intrusions of UCDW are needed to stimulate phytoplankton community development/succession and biological production (evidenced by changes in nutrient ratios) on the WAP continental shelf.

The occurrence and progression of an intrusion event in one area of the shelf is independent of intrusions that occur in other regions of the WAP shelf. Thus, at any given time one or more intrusions, in different stages of development, may be present on the WAP continental shelf. This can provide considerable spatial variability in production, which must be taken into account when attempting system-wide production estimates.

d. Implications for non seasonal succession

Phytoplankton community succession certainly occurs in Antarctic coastal waters as water columns stratify and nutrient limitation occurs (Moline and Prézelin, 1996a and references therein; Moline et al., 1997). However, this succession is subject to disruption by physical forcing events such as upwelling, wind mixing, and freshwater lenses produced by seasonal glacial ice melt that seal off the upper mixed layer (Moline and Prézelin, 1996a). In waters over the WAP continental shelf, the occurrence of UCDW intrusions, and subsequent upwelling and/or mixing at specific locations, provides an episodic and major disruption of ongoing phytoplankton community succession.

The UCDW intrusions force a resetting of the chemical system to one that assures the immediate development of diatom-dominated communities over whatever phytoplankton assemblages were present at the time of the event. Over the shelf, phytoplankton community succession from diatom-dominated to phytoflagellate-dominant communities, which eventually contain a strong dinoflagellate component, proceeds as the UCDW intrusion subsides and as water column stratification leads to nutrient limitation within the upper mixed layer. The degree of succession from diatoms to phytoflagellates depends upon the frequency of the physical forcing of UCDW intrusions. Thus, classic seasonal succession from spring diatoms to summer phytoflagellates would not be a good representation of phytoplankton community changes at the regional scale. Recognizing the role that the presence or absence of UCDW intrusion events have in structuring phytoplankton communities makes it possible to explain the complexity of the observed phytoplankton distributions on the WAP continental shelf.

e. Implications for production and carbon partitioning

Episodic resetting of the western Antarctic continental shelf waters to diatom-dominated communities, with enhanced biological production, can alter the partitioning of organic carbon and change food web dynamics. Environmental stress is initially low at the start of an UCDW intrusion event because nutrients are high. Little dissolved organic carbon (DOC) is produced as diatom growth accelerates and cells remain in log-phase growth (Hansell and Carlson, 2002). The DOC that is produced should be near Redfield ratios in
composition and thus quite labile to microbial remineralization (Smith et al., 1998; Carlson et al., 1998, 2000; Carlson and Hansell, 2004). The majority of new organic carbon is as POC. However, this production may go unrecognized if the diatoms are advected off the shelf, if the UCDW intrusion sites are high grazing sites, and/or if the majority of UCDW-induced production occurs at a depth deeper than usually assumed in satellite-remote sensing algorithms (Moline and Prézelin, 2000).

With cessation of an UCDW intrusion event, diatoms face declining nutrient supply, resulting in less POC production and more DOC excretion. The C:N ratio of this DOC should be higher and have reduced lability (Hansell and Carlson, 2002), resulting in declining food quality of the diatom-dominated community. As diatoms enter senescence, they tend to aggregate and sediment out of the mixed layer. Thus, the frequency and duration of UCDW intrusion events can also play an important role in benthic community dynamics on the WAP shelf.

Phytoflagellates achieve dominance in UCDW-influenced waters that have been biologically processed by diatoms and thus contain lower nutrient concentrations. However, they are also abundant in high-nutrient waters not influenced by UCDW and unable to support diatoms, such as the outflow of the Gerlache Strait in March 1993 (Fig. 5b). Both types of nutrient sources may initially sustain phytoflagellate growth but the carrying capacity (biomass and/or potential primary production) of the UCDW-influenced waters may be significantly less. Antarctic phytoflagellates tend be nanoplankton communities where microbial loop dynamics may be of central importance to carbon cycling.

f. Extensions to other regions

This analysis shows that UCDW intrusions, with their subsequent biological effects, occur at other times of year and at other sites than described in Prézelin et al. (2000) for the austral summer of 1993. The extent to which patterns and distributions observed on the WAP continental shelf are general features of Antarctic coastal waters can be partially addressed by observations reported in other studies. Kang and Lee (1995) reported abrupt changes in phytoplankton taxonomic assemblages along two across-shelf transects made in February 1993 that extended from western Bransfield Strait to Drake Passage (Fig. 1). Diatom dominance of the phytoplankton biomass increased abruptly on the outer shelf and shelf break seaward of the islands at the western end of Bransfield Strait. The maximum in diatom abundance was at about 60 m. The observations made by Kang and Lee (1995) only extended to 100 m, but the areas where diatoms dominated coincided with warmer temperatures, high macronutrients, and reduced Si(OH)₄:NO₃ ratios (their Figs. 2 and 3). Kang and Lee (1995) speculated that the dominance of diatoms was related to regional hydrographic features and that it may be possible to use diatoms and other phytoplankton species as markers for specific water masses within hydrographic regimes.

As suggested by Prézelin et al. (2000), places where the Southern ACC Front (SACCF) nears the shelf edge are potential sites for UCDW intrusions. These sites are nonuniformly distributed around the Antarctic (Jacobs, 1991; Whitworth et al., 1998). Direct evidence
for increased biological production in these regions is lacking because of the limited historical observations for Antarctic coastal waters. However, indirect evidence is provided by the distribution of regions of high concentrations of Antarctic krill (Everson and Miller, 1994) and the distribution of cetaceans (Tynan, 1998). These studies provide a qualitative correlation between areas where the SACCFA nears the shelf edge, the shallowing of CDW, and enhanced biological production. These correspondences clearly warrant further study, given the implications of the results from this study.

In oceanic waters off the Antarctic continental shelf (see Fig. 1), UCDW upwelling was observed when leads in the sea ice were present in the MIZ (Prézelin et al., 1994). These open regions within the sea ice became diatom-dominated while Phaeocystis-dominated communities persisted under the sea ice. The Si(OH)₄:NO₃ ratio was initially high, indicating input of UCDW, and then decreased as the nutrients were used by the diatoms. Two UCDW upwelling events occurred within one month at this study area. This open-ocean upwelling of UCDW provides episodic inputs of nutrients to the upper water column, which potentially results in enhanced diatom growth and increased biological production. These observations indicate another mechanism for increasing the biological production of Antarctic waters that has yet to be fully quantified.

5. Conclusion

Beyond observations of the role of UCDW in the reproductive success of Antarctic krill (e.g. Hempel et al., 1979; Hofmann et al., 1992) and its potential importance for cetaceans (Tynan, 1998), the effect of this water mass on biological production of Antarctic continental shelf waters was not considered until the study by Prézelin et al. (2000). The analyses presented here extend those given in Prézelin et al. (2000) to show that UCDW is a primary determinant of phytoplankton community composition and production in WAP continental shelf waters, and that it is potentially important for the larger Antarctic system. These observations and analyses make it surprising that UCDW intrusions are only now being recognized as being important to the productivity of Antarctic waters. However, several factors likely contributed to the lack of recognition of this process.

Previous biological studies did not routinely measure phytoplankton community composition and typically focused on the upper 100 m of the water column (see Priddle et al. (1994) for a discussion), which does not capture the subpycnocline inputs of UCDW. Thus, the sampling strategies were such that the role of the deeper processes in controlling phytoplankton community structure in the upper water column would not be recognized. The weak stratification of Antarctic continental shelf waters allows tight coupling between deep and surface processes and this characteristic of the environment needs to be incorporated into future biological studies.

The phytoplankton communities that develop in response to UCDW intrusions are subsurface and are at depths that are not detectable by satellite-borne ocean color sensors. Thus, these episodic events are not represented in the existing ocean color measurements.
An implication of this is that primary production and chlorophyll $a$ estimates made for Antarctic continental shelf waters from ocean color measurements will be underestimates. The photosynthetic physiological measurements made in 1993 and 1994 show that high rates of primary production are the norm in areas affected by UCDW intrusions. Thus, the observed low biomass with relatively high growth potential likely reflects heavy grazing of a highly productive system and/or advective loss due to the ACC and shelf currents. Thus, chlorophyll $a$ estimates alone, which are the majority of the available measurements for Antarctic coastal waters, are not sufficient to describe the phytoplankton community growth potential. Not surprising, these measurements need to be constrained with coincident measures of primary production, grazing rates and grazer biomass estimates.

Subsurface intrusions and subsequent upwelling or upward mixing of oceanic waters have been observed in other regions, such as the southeastern U.S. continental shelf (Atkinson, 1977, 1985) and the east coast of Australia (Andrews and Gentien, 1982). Estimates of the nutrient and carbon fluxes associated with intrusion events on these continental shelves (e.g. Lee et al., 1991) show that these events account for a significant portion of organic matter production. On the southeastern U.S. continental shelf, phytoplankton production and biomass are $>10$ times higher within intrusions than surrounding resident shelf waters (Yoder, 1985). Lee et al. (1991) estimated that nitrogen flux associated with upwelled Gulf Stream-derived waters on the southeastern U.S. shelf could support production of ca 8 million tons of carbon per year, if all of the nitrogen was used. The potential of biological production associated with UCDW intrusions on the Antarctic continental shelf has yet to be factored into nutrient and carbon budgets constructed for this region. The data sets used in this study, while comprehensive, are not adequate to quantify exchange rates, particle transformations, and total production associated with UCDW events. Multidisciplinary studies that follow an intrusion event through its entire life cycle on the shelf are needed and must be flexible enough to adapt to a variable environment. Failure to do so will result in missing a major portion of the signal that is controlling biological production in Antarctic coastal waters.

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REFERENCES


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