Nearshore mixing and nutrient delivery along the western Antarctic Peninsula

D. REIDE CORBETT1,2, JARED CRENSHAW1, KIMBERLY NULL1,3, RICHARD N. PETERSON4, LEIGHA E. PETERSON4 and W. BERRY LYONS5

1East Carolina University, Greenville, NC 27858, USA
2University of North Carolina Coastal Studies Institute, Wanchese, NC 27981, USA
3Current address: Moss Landing Marine Laboratories, Moss Landing, CA 95039, USA
4Department of Coastal and Marine Systems Science, Coastal Carolina University, Conway, SC 29526, USA
5School of Earth Sciences and Byrd Polar and Climate Research Center, Ohio State University, Columbus, OH 43210, USA
corbettd@ecu.edu

Abstract: The surface waters of the Southern Ocean play a key role in the global climate and carbon cycles by promoting growth of some of the world’s largest phytoplankton blooms. Several studies have emphasized the importance of glacial and sediment inputs of Fe that fuel the primary production of the Fe-limited Southern Ocean. Although the fertile surface waters along the shelf of the western Antarctic Peninsula (WAP) are influenced by large inputs of freshwater, this freshwater may take multiple pathways (e.g. calving, streams, groundwater discharge) with different degrees of water-rock interactions leading to variable Fe flux to coastal waters. During the summers of 2012–13 and 2013–14, seawater samples were collected along the WAP, near Anvers Island, to observe water column dynamics in nearshore and offshore waters. Tracers (223,224Ra, 222Rn, 18O, 2H) were used to evaluate the source and transport of water and nutrients in coastal fjords and across the shelf. Coastal waters are compared across two field seasons, with increased freshwater observed during 2014. Horizontal mixing rates of water masses along the WAP ranged from 110–3600 m² s⁻¹. These mixing rates suggest a rapid transport mechanism for moving meltwater offshore.

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Introduction

Over the past one hundred years, average global temperatures have been increasing. The most drastic warming has taken place in climate-sensitive regions at high latitudes, including the western Antarctic Peninsula (WAP; Schofield et al. 2010; Fig. 1) where annual mean atmospheric temperatures have increased by ~3°C over the past 60 years (King 1994, Turner et al. 2005). Due to recent warming, glaciers between 64°S and 66°S are currently receding at an average rate of ~57 ± 25 m y⁻¹ (Cook et al. 2005). Warming air temperatures have been concurrent with sea ice reduction, with many glaciers having near-frontal surface lowering of 0.28 ± 0.03 m y⁻¹ over the past 40 years (Kunz et al. 2012). This in turn has increased glacial melt inputs that have the potential to impact salinity and temperature dynamics in surface waters along the shoreline (Ducklow et al. 2012). The input of glacial melt can also lead to increases in many biogeochemically important elements (e.g. N, P, C, Si, Fe) into the coastal ocean (Wadham et al. 2013). Surface waters of the Southern Ocean play a key role in global climate and carbon cycles by hosting some of the world’s largest phytoplankton blooms, which sequester up to one billion metric tons of CO₂ per year (Takahashi et al. 2012).

Continently derived freshwater inputs often enter coastal waters during meltwater pulses along the WAP. This meltwater can enter the nearshore coastal zone through calving, periglacial stream flow, basal melt and subglacial melt, which passes over or through sediments or other geologic material before reaching the ocean. These pulses create large freshwater lenses that can extend as deep as 50 m into the water column, and as far as 100 km offshore (Dierssen et al. 2002). Meltwater pulses have the potential to develop density gradients that can drive surface currents, resulting in the lateral transport of continentally derived nutrients and micronutrients (e.g. Fe) offshore (Dierssen et al. 2002, Moffat et al. 2008). Dierssen et al. (2002) found that >70% of phytoplankton blooms observed offshore of Anvers Island occurred shortly after pulses of meltwater discharged from local glaciers into the surface ocean.

Recent studies (Bhatia et al. 2013, Hawkings et al. 2014, Lyons et al. 2015) have emphasized the potential importance of direct and indirect glacial inputs to Fe budgets in polar
regions, and how these inputs may fuel primary productivity in Fe-limited environments such as the Southern Ocean (Takahashi et al. 2012). Fertile surface waters along the shelf of the WAP are influenced by large inputs of glacial melt from the peninsula (Dierssen et al. 2002). The objective of this study was to evaluate the transport of nutrient-rich meltwaters from the Antarctic continent into the biologically productive coastal waters along the WAP utilizing a suite of naturally occurring stable- and radioisotopes as tracers of terrestrially derived water masses.

Study area

The WAP extends northward from the western Antarctic continent. This region is the warmest in the Antarctic and one of the most rapidly warming regions in the world (Vaughan et al. 2003, Steig et al. 2009). The rugged topography and warmer maritime climate of the peninsula differentiates the WAP from other regions of the continent. Air temperatures along the western coast of the WAP are dominated by the influence of the Bellingshausen Sea, causing air temperatures to be ~ 7°C warmer than the eastern coast of the peninsula, which is influenced by the cooler Weddell Sea (Morris & Vaughan 2003). This region has seen an increase in July (i.e. midwinter) temperatures of 7.2°C from 1951–2008 (0.124 ± 0.044°C yr⁻¹; Ducklow et al. 2012).

The continental shelf of the WAP is ~ 200 km wide near Anvers Island and has an average depth of ~ 450 m. The aforementioned depth excludes many of the deep canyons that incise the continental shelf, such as the Palmer Deep that reaches a maximum depth of ~ 1500 m. Several

Fig. 1. Sample collection sites and transects (dashed line) during the 2012 (blue squares; T1, 2, 3) and 2014 (red circles; T4, 5, 6) cruises offshore of the western Antarctic Peninsula. Dotted line represents the delineation of nearshore versus offshore samples. Box encompassing sites within Flandres Bay and the T2,T5 line were used to produce the density cross-section (see Fig. 4). Yellow triangle denotes the approximate location of Palmer Station.
glacially carved fjords line the coast of the WAP and were also sampled as part of this research. Flandres Bay is protected from the open ocean by Anvers Island and several smaller islands. The 16 km wide fjord is flanked by mountains on both sides and reaches depths > 600 m. The mouth of Flandres Bay opens to the Gerlache Strait. Barilari Bay and Beascochea Bay are both similarly flanked by rugged mountains, and open to the Grandidier Channel (Fig. 1).

Surface water characteristics along the WAP vary widely on a seasonal basis. During the summer, Antarctic Surface Water (AASW) found above the permanent thermocline (~ 150 m), ranges in temperature from -1.5 – 1.0°C and in salinity from 33.0 – 33.7 (Klinck et al. 2004). During the winter, a relatively cooler and saltier water mass develops near the surface (Winter Water), generally cooler than -1.8°C with a salinity of ~ 34.1 (Klinck et al. 2004). Subpycnocline waters are classified as Circumpolar Deep Water (CDW), subdivided into upper CDW and lower CDW. CDW ranges in temperature

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**Fig. 2.** Air temperature measured at Palmer Station from 2012 through to the end of 2014. Offshore sampling took place in December 2012 (squares) and March 2014 (circles). Hatched boxes are used to illustrate temperatures two months prior to the cruise. Note temperatures below freezing prior to 2012 cruise and above freezing prior to 2014.

**Fig. 3.** Spatial distribution of salinity and temperature in surface waters during the 2012 and 2014 cruises. Note the lower salinities and higher temperatures nearshore during the 2014 relative to the 2012 cruise.
from 1.0–2.0°C and has salinities of 34.6–34.7. Upper CDW has frequent incursions into coastal waters along the continental shelf, mixing with surface waters (Klinck et al. 2004, Meredith et al. 2013).

Cyclonic circulation patterns along the WAP were identified in early oceanographic studies, and further described by Savidge & Amft (2009). Long-term in situ measurements along the peninsula suggest that a cyclonic cell rotates around Anvers Island. Surface waters offshore of Anvers Island flow south and form part of the Antarctic Peninsula Coastal Current (APCC), an along-shore surface current that occurs on the west coast of the peninsula during the summer months (Moffat et al. 2008). The development of the APCC is thought to be initiated by meltwater inputs along the peninsula, although the origin of these waters is poorly understood. The APCC has been observed as far north as Brabant and Anvers islands (Savidge & Amft 2009) and flows southward, eventually making its way into Marguerite Bay and farther south (Moffat et al. 2008). In the Gerlache Strait, surface currents have been observed moving north toward the Bransfield Strait (Savidge & Amft 2009).

Fig. 4. Density cross-section starting inside Flandres Bay and moving across the shelf on T2 (2012) and T5 (2014). Refer to box area in Fig. 1 for cross-section sampling sites. Dashed lines are isopycnals, vertical lines and black circles refer to CTD cast sites and sampling depths. Note increased depth and gradient during 2014.
Groundwaters were collected using a custom drive-point piezometer lined with Teflon tubing. Surface runoff was collected from several different streams via a peristaltic pump near Palmer Station prior to discharge into surface waters. Samples were analysed for a suite of parameters, including radium isotopes (223Ra, 224Ra), δ18O and δD, and nutrient and trace metal concentrations (NO3−, NO2−, NH4+, HPO42−, H4SiO4, total dissolved Fe).

At each nearshore and offshore site, the CTD-rosette (Seabird SBE911) was lowered multiple times from the ship to within a few metres of the sea floor. Temperature (SBE3), salinity (SBE4), density, beam transmission (Wet Labs C-Star), oxygen (SBE43) and fluorescence (Wet Labs ECO) were recorded. Along the cruise track during each field season, a hull mounted acoustic doppler current profiler (ADCP) was used to measure current velocities throughout the water column.

**Methods**

**Sample collection**

During the summers of 2012–13 and 2013–14 (14–19 December 2012 and 19–24 February 2014, respectively; Fig. 2), large volume seawater samples (280–360l) were collected along the WAP aboard the RV Lawrence M. Gould to observe water column and tracer dynamics in nearshore and offshore environments (Fig. 1). Groundwater and surface runoff from glacial melt (e.g. streams) samples (20–150l) were collected while at Palmer Station following similar shipboard measurements. During each field season, offshore and nearshore seawater was collected from one to three depths (surface, mid-depth and bottom) along three transects (2012: T1, T2, T3; 2014: T4, T5, T6) and within glacial fjords. Due to limited ship time in 2014, sample collection was focused on surface water collection. Surface water samples were collected in the upper 10 m of the water column. Mid-depth and bottom water samples were collected just above the pycnocline (30–60m) and ~10 m above the bottom, respectively.

**Radium isotopes**

Radium was quantitatively concentrated from water samples by gravity filtering (rate of < 11 min−1) onto MnO2-coated acrylic fibre (i.e. Mn fibres). Fibres were then rinsed with radium-free water, partially dried with compressed air and immediately analysed for total 223Ra and 224Ra using a radium delayed coincidence counter (RaDeCC; Moore & Arnold 1996). At least four weeks (eight half-lives of 224Ra) after this initial count, fibres were recounted via the RaDeCC for 224Ra supported by 228Th sorbed to the fibres. Excess, or unsupported 224Ra (presented herein), is taken as the difference between the initial count (total) and the second measurement (supported). 223Ra is presented as total, not corrected for 227Ac (negligible in most instances). Detector efficiencies and analytical precision (0.08 dpm m−3 for 223Ra and 0.15 dpm m−3 for 224Ra) were determined from multiple analyses of prepared standard solutions (acquired from the National Physical Laboratory) and

| Tracer/nutrient | Groundwater (n = 29) | Surface runoff (n = 16) |
|----------------|----------------------|-------------------------|
| Salinity       | 1.3 ± 0.3            | 0.03 ± 0.01             |
| Temperature (°C) | 2.6 ± 0.2            | 2.6 ± 0.5               |
| δ18O (%)       | -11.4 ± 0.4          | -10.0 ± 0.6             |
| δD (%)         | -89.4 ± 3.0          | -78.2 ± 4.7             |
| 223Ra (dpm m−3) | 83 ± 22              | 57 ± 15                 |
| 223Ra (dpm m−3) | 3.2 ± 0.9            | 8.2 ± 4.4               |
| 224Ra (dpm m−3) | 33 ± 19              | 66 ± 34                 |
| NO3− + NO2− (μM) | 57.5 ± 3.5          | 33.2 ± 10.6             |
| NH4+ (μM)      | 23.5 ± 7.0           | 13.9 ± 2.4              |
| HPO42− (μM)    | 2.6 ± 0.5            | 1.2 ± 0.2               |
| H4SiO4 (μM)    | 18.6 ± 3.0           | 37.9 ± 12.2            |
| dFe (μM)       | 35.4 ± 7.1           | 0.47 ± 0.12             |

n = 12, b = 18, c = 6.
intercalibrated among laboratories participating in GEOTRACES using a similar standard prepared from an International Atomic Energy Agency solution. Analytical uncertainty for individual samples were based on propagated counting statistics, and averaged 7% and 21% for $^{224}$Ra and $^{223}$Ra, respectively.

**Radon**

Radon analyses were completed by collecting 6 l of water at each location and depth prior to any other sample collection to avoid potential degassing from the Niskin bottles. Seawater was siphoned into a 6 l Nalgene bottle.
with a filling–venting cap to minimize radon evasion. Each seawater sample was analysed with a RAD-7 radon-in-air monitor (Durridge) via procedures outlined in Lee & Kim (2006). The RAD-7 uses a silicon semiconductor at ground potential to measure the short-lived Po isotopes emitted by $^{222}$Rn decay.

**Nutrients, $\delta^{18}O$ and $\deltaD$**

A 60 ml seawater sample was collected at each sampling location and depth for nutrient analyses ($NO_3^- + NO_2^-, NH_4^+, HPO_4^{2-}, H_4SiO_4$). Each sample was syringe filtered on-site through a 0.45 µm filter into a 60 ml polypropylene bottle. All sampling materials and bottles were acid-cleaned prior to use. Samples collected were stored at -20°C until analysis on a Lachat Quickchem Autoanalyzer.

Dissolved iron (dFe) was sampled from freshwater endmembers using acid-cleaned material and trace metal clean protocols. Samples were pumped immediately from source waters (e.g. groundwater, stream, etc.) through an inline 0.2 µm membrane filter into a polyethylene sample bottle using a peristaltic pump to avoid exposure to air. All sampling materials and bottles were acid-cleaned prior to use. Samples were preserved with Optima ultrapure nitric acid and shipped to Skidaway Institute of Oceanography for analysis on a Perkin Elmer NexION 300D ICP-MS.

Water isotope samples were collected at the same time as nutrient samples. To minimize degassing, water was allowed to overflow a 20 ml scintillation vial for $>10$ sec. After the scintillation vial was completely filled with seawater with no head space, vials were capped, sealed with parafilm and stored inverted at 4°C. Samples were shipped to Ohio State University for analysis on a Picarro L2130 Wavelength Scanned-Cavity Ring Down Spectrometer water isotope analyser. Calculated values are relative to Vienna Standard Mean Ocean Water (VSMOW) with precision of 0.03‰ and 0.1‰ for $\delta^{18}O$ and $\deltaD$, respectively.

**Fig. 6.** Spatial distribution of water isotope ($\delta^{18}O$, $\deltaD$) surface water concentrations from the 2012 and 2014 cruises. All surface samples are reported, including those that are probably outliers leading to a shelf ‘bullseye’ in 2012. Note more depleted isotopic signatures nearshore during the 2014 cruise.
Results

Physical parameters

Air temperature measured at Palmer Station provides context for differences observed between the two cruises. Average air temperature for one and two months prior to the 2012 cruise (Fig. 2, hatched area) were -0.3 ± 1.3°C and -2.2 ± 3.2°C, respectively. Average air temperature for one and two months prior to the 2014 cruise (Fig. 2, hatched area) were 1.1 ± 1.1°C and 1.1 ± 1.4°C, respectively. Generally, air temperatures were below freezing and more variable prior to the 2012 cruise, and

Fig. 7. Spatial distribution of nutrient (NH$_4^+$, NO$_3^-$, HPO$_4^{2-}$) surface water concentrations from the 2012 and 2014 cruises. Concentrations nearshore are more depleted in 2014.
above freezing and less variable prior to the 2014 cruise (Fig. 2). Temperatures were significantly higher \((P < 0.01)\) during the 2014 cruise.

During the cruises, surface water temperatures varied from -1.5 to 2.8°C, with an average of 0.5°C. The warmest waters were observed in Flandres Bay and regions closer to shore. The coldest surface temperatures were measured along T3 in 2012 and in Barilari Bay during the February 2014 cruise (Fig. 3). Temperature minima (-2°C) existed during both field seasons throughout the study area at 50–100 m depth. These temperature minima were not present closer to the shoreline, where density gradients were less prevalent across the permanent pycnocline at a depth of ~ 100–150 m (Fig. 4). Surface salinities ranged from 31.2–34.1 with an average of 33.2. Fresher waters were found closest to shore in Flandres Bay, Beascochea Bay and Barilari Bay (Fig. 3). The surface mixed layer did not extend below ~ 40 m depth throughout the study area. The coldest surface temperatures were measured from patterns observed in orthophosphate and nitrate/nitrite, although ammonium was similarly depleted in some areas (Beascochea and Barilari bays). Increased ammonium concentrations in surface waters were observed in the vicinity of Grandidier Channel and in Flandres Bay. Silicate concentrations in surface waters varied from 38–88 μM (Fig. 7). Surface waters in Flandres Bay and in the Gerlache Channel were enriched relative to offshore surface waters and fjords to the south. In deeper waters, silicate concentrations range from 87–108 μM.

**Isotopic tracers**

During the 2012 and 2014 field campaigns, short-lived isotopes (i.e. Ra, Rn) were measured in groundwaters and streams near Palmer Station. In addition, surface waters offshore of Anvers Island and in nearshore fjords were sampled aboard the RV Laurence M. Gould. Groundwater and stream waters had elevated activities of all radiotracers (Table I) relative to nearshore and offshore surface water (Table II). Activities of these freshwater endmembers were as much as three orders of magnitude higher than nearby surface waters.

Surface 223,224Ra activities ranged from 0.1–1.0 dpm m⁻³ and 0.1–12.8 dpm m⁻³, respectively (Fig. 5). Similarly, deep water 223,224Ra activities ranged from 0.2–1.7 dpm m⁻³ and 0.2–16.1 dpm m⁻³, respectively. Surface 222Rn activities ranged from 0.3–0.8 dpm l⁻¹, while deeper waters ranged from 0.1–1.1 dpm l⁻¹. Surface waters were found to be enriched in short-lived Rn and Ra isotopes in the vicinity of Anvers Island and Flandres Bay in 2012, and nearshore in Barilari and Flandres Bays in 2014 (Fig. 5).

Deuterium values in surface and deep waters ranged from -12.1 – -3.7‰ and -11.2 – -0.5‰, respectively. 18O values in surface and deep waters ranged from -1.9–0.4‰ and -1.36–0.02‰, respectively. The distribution of δ18O and δD co-vary. Generally, surface samples nearshore were more depleted relative to stations offshore (Fig. 6, Table II). Gradients in surface water δ18O and δD were most prominent in the fjords (Fig. 6). Offshore surface samples showed high variability in the isotopic values, probably an indication of advective mixing.

**Nutrients**

Orthophosphate concentrations in surface waters ranged from 0.12–2.3 μM. Nitrate/nitrite concentrations in surface waters ranged from 4.0–32 μM. Nitrate/nitrite and phosphate concentrations showed similar patterns (Fig. 7). Generally, bottom water concentrations were higher than surface water values, and surface water concentrations along the coast were lower than offshore waters. Deep water orthophosphate and nitrate/nitrite concentrations ranged from 1.8–2.4 μM and 16.8–37 μM, respectively.

Ammonium concentrations in surface waters ranged from below detection to 6.5 μM (Fig. 7). In deeper waters, ammonium concentrations range from below detection to 1.6 μM. Patterns in the distribution of ammonium differ from patterns observed in orthophosphate and nitrate/nitrite, although ammonium was similarly depleted in some areas (Beascochea and Barilari bays). Increased ammonium concentrations in surface waters were observed in the vicinity of Grandidier Channel and in Flandres Bay. Silicate concentrations in surface waters varied from 38–88 μM (Fig. 7). Surface waters in Flandres Bay and in the Gerlache Channel were enriched relative to offshore surface waters and fjords to the south. In deeper waters, silicate concentrations range from 87–108 μM.

**Discussion**

**Tracer signatures of freshwater sources**

Along the shoreline of the WAP and nearby islands, fresher water masses were observed in the upper layers of the water column (Figs 3 & 4). These salinity differences in the upper parts of the water column co-varied with physical and geochemical parameters that have been used to identify glacial melt, such as δ18O and our radiotracers (Figs 5 & 6). The light isotopic signature of glacial ice has been used previously to trace meltwater pulses along the Antarctic continental shelf (Schlosser et al. 1990, Weppernig et al. 1996, Meredith et al. 2008). Groundwaters and surface runoff from glacial melt (i.e. streams) had an average δ18O signature of -11.4‰ and -10‰, respectively, while the average δD was -89.4‰ and -78.2‰, respectively (Table I).

Groundwaters and streams were enriched in Ra and Rn relative to seawater, and depleted in 18O. Groundwater samples from Anvers Island had average activities of 3.2 and 83 dpm m⁻³ for 223Ra and 224Ra, respectively, which are high relative to average surface water values observed offshore (Tables I & II, Fig. 5). Streams were also enriched in Ra isotopes relative to surface waters along the WAP. On average, streams had activities of 8.2 and 57 dpm m⁻³ for 223Ra and 224Ra, respectively (Table I). Similarly, Rn was enriched in groundwaters and streams.
relative to nearshore and offshore surface waters (Tables I & II). The relative enrichment of Ra and Rn in terrestrially derived waters can be used to identify and evaluate the freshwater (e.g. meltwater) inputs observed along the WAP, assuming other potential sources (e.g. bottom sediments) are taken into account. However, this study has focused on surface waters above the pycnocline in water depths (200–600 m) that far exceed the surface mixed layer (~ 30 m) therefore limiting the potential inputs of these short-lived tracers from bottom sediments or deep water sources.

Tracer variability with proximity to the shoreline
Changes in surface salinity, temperature and tracer activities vary with the amount of meltwater entering the nearshore environment. These parameters and others change with the progression of the summer as more meltwater enters the nearshore environment. The averages of surface temperature and salinity values (Table II) remain within the limits of AASW defined by Klinck et al. (2004) with the exception of nearshore salinity in 2014 (32.7 ± 0.1). These waters were significantly fresher on average than AASW and the other waters measured during this study, which suggests that local glacial melt inputs are mixing with nearshore surface waters. Data from the December 2012 cruise suggest little change in surface seawater characteristics with distance from the shoreline. This cruise took place during the early weeks of the summer when the melt season was just beginning. A notable difference is that offshore waters were much cooler than surface waters nearshore. Glacial melt signals were not as prominent across the shelf compared to gradients in nearshore surface waters. The 2014 cruise took place during the latter phases of the summer. At this point in the season, there were notable differences in the physical and geochemical characteristics of surface seawater nearshore relative to offshore waters. Nearshore surface waters had lower salinity, temperature, δ18O and δD values, and higher $^{223,224}$Ra activities relative to offshore surface waters. These changes probably document an increased meltwater influence with the progression of the summer and the utility of these geochemical tracers in identifying meltwater input. The location and intensity of meltwater inputs can impact and influence the prolific phytoplankton blooms that take place along the WAP (Dierssen et al. 2002). In order to gain an understanding of how quickly these water masses can be distributed along the WAP, mixing models were employed to evaluate coastal and cross-shelf mixing.

Nearshore mixing
Radon and short-lived Ra were used to estimate mixing in nearshore waters. Assuming the source of these tracers is nearshore (e.g. meltwater) and changes in activity are a function of mixing with offshore water of lower activity and radioactive decay, then the change in activity can be modelled via:

$$A_x = A_0 e^{-x \sqrt{\frac{\lambda}{K_h}}}$$  \hspace{1cm} (1)

where $A_x$ is the activity at distance $x$ from the coast, $A_0$ is the activity at the coast ($x = 0$), $K_h$ is the mixing coefficient and $\lambda$ is the decay constant (Moore 2000). Additional assumptions include: i) the system is at steady state, ii) a constant source of tracer only at the landward boundary ($A_0$) and iii) advection is not significant across the distance modelled. We are confident in these assumptions given that no influence of advection was observed (i.e. no deviation from the linear trend of ln (tracer) with distance) and the water depth (200–600 m) far exceeds the surface mixed layer (~ 30 m) limiting new tracer inputs from bottom sediments. Although short-lived Ra isotopes have typically been employed with this method (Moore 2000, Annett et al. 2013), assuming

![Fig. 8. The a. ln$^{222}$Rn and b. ln$^{224}$Ra profiles from Flandres Bay in 2012 and 2014, and well as Beascochea Bay in 2014. Lines represent the best-fit trend line used to calculate mixing rates for each tracer during each sampling year.](image-url)
atmospheric evasion is minimal, radon should provide similar results as the source of the tracers is assumed to be the same (e.g. meltwater). Mixing rates were estimated within two fjords (i.e. Flandres and Beascochea bays) where degassing was reduced (< 0.1 dpm m\textsuperscript{-2} min\textsuperscript{-1}) due to low winds (< 5 m s\textsuperscript{-1}) based on the calculated flux of Rn from the sea surface to the atmosphere for the 24 hours prior to sample collection.

In Flandres Bay, 2\textsuperscript{222}Rn-derived mixing rates were 1020 ± 300 and 570 ± 160 m\textsuperscript{2} s\textsuperscript{-1} for the 2012 and 2014 cruises, respectively, while 2\textsuperscript{234}Ra-derived mixing rates were 570 ± 60 and 110 ± 20 m\textsuperscript{2} s\textsuperscript{-1}, respectively, for the same period (Fig. 8). Both tracers show a decreased mixing rate during 2014 and may represent seasonal variability. Mixing in Beascochea Bay was estimated at 3600 ± 200 m\textsuperscript{2} s\textsuperscript{-1} (Fig. 8) using measurements spanning a distance of ~ 31 km. Similar to Flandres Bay, Beascochea Bay is flanked by glaciers but opens to the relatively unprotected Grandidier Channel at its mouth.

Calculated mixing rates were within the range of other rates calculated along the WAP to the north where Dulaiova et al. (2009) estimated mixing rates via Ra isotopes along the South Shetland Islands of 596–63 000 m\textsuperscript{2} s\textsuperscript{-1} in open ocean waters. This location is where the Antarctic Circumpolar Current approaches the northern edge of the peninsula. However, south of our study area, Annett et al. (2013) calculated mixing rates using Ra isotopes within Marguerite Bay of 0.13–1.0 m\textsuperscript{2} s\textsuperscript{-1}, which are several orders of magnitude lower than those calculated in this study and probably reflect the low-velocity circulation dynamics of the more protected embayment of Marguerite Bay.

Estimating cross-shelf mixing via 2\textsuperscript{233},2\textsuperscript{24}Ra and 2\textsuperscript{222}Rn proved to be more problematic. Mixing models using terrestrially derived tracers assume that there is a single source of tracer emanating from the continental margin and that vertical and lateral (e.g. along-shelf) advective transport is negligible. Surface currents and circulation dynamics across the continental margin are much more complex than the models used here to evaluate them. The ADCP data collected as part of this study and composite data presented by Savidge & Amft (2009) show that circulation patterns are complicated, and vary over different timescales. Recent surface current measurements via high-frequency radar demonstrates the spatial and temporal complexity of currents in this region, with large short-term areas of convergence and divergence, intermixed with cross-shelf and along-shelf advection (Kohut et al. 2016). However, unlike the shelf, the fjords provide an area where the assumptions essential for this type of unidirectional circulation model are more plausible. Because the offshore areas are more susceptible to advective mixing in the along- and cross-shore directions, mixing rates are probably much higher than those estimated here for the fjords.

**Seasonal variability: nutrient transport and shelf productivity**

As noted previously, the December 2012 survey was conducted at the beginning of the summer when the surface waters ranged in salinity from 32.6–34.8 with a mean of 33.7. Radium values in surface waters averaged 1.5 dpm m\textsuperscript{-3}. The average atmospheric temperature for November and December 2012 was -2.2°C (Fig. 2). The 2014 cruise took place during the latter part of February, as the summer season was coming to an end. Surface waters ranged in salinity from 31.2–33.6 and averaged 32.9. For surface waters in 2014, Ra values averaged 3.03 dpm m\textsuperscript{-3}. January and February 2014 atmospheric temperatures at Palmer Station averaged 1.1°C (Fig. 2). As the season progresses, air temperatures > 0°C are more common and there are fewer hours of darkness. The increased temperatures and exposure to sunlight enhance glacial melting, progressively releasing vast quantities of meltwater into the surface ocean.

The highest fluorescence values (6–8 mg m\textsuperscript{-3}) reported in this study were measured in the Grandidier Channel seaward of Barilari Bay. These waters were enriched in 2\textsuperscript{222}Rn, 2\textsuperscript{233}Ra and 2\textsuperscript{24}Ra relative to surface waters offshore where fluorescence (e.g. primary productivity) was lower (0–2 mg m\textsuperscript{-3}). These nearshore waters were clearly fresher, colder, and depleted in \textsuperscript{18}O and D relative to adjacent surface waters (Fig. 6), suggesting that glacial melt comprises a higher percentage of surface waters in the Grandidier Channel near Barilari Bay. Given the higher macro and micronutrient concentrations measured in these terrestrially derived waters, it is hypothesized that nutrient transport via glacial melt plays a role in fuelling the observed primary productivity (e.g. higher fluorescence).

Several studies (Moore 1996, Dierssen et al. 2002, Windom et al. 2006) have linked the terrestrial environment and primary productivity in the surface ocean through submarine groundwater discharge with respect to macro and micronutrients. Unlike the temperate and tropical environments where groundwater discharge studies have focused, the WAP is quite different in that it has a vast reservoir of frozen water covering the majority of the land surface that partially melts during summer months. This study suggests that primary productivity along the WAP could be linked to nutrients provided by these seasonal meltwater inputs following different delivery pathways. Mixing rates in the surface waters observed in nearby Beascochea Bay and within Flandres Bay are within the range of those calculated along the northern reaches of the WAP (Dulaiova et al. 2009), and higher than those calculated in the more isolated Marguerite Bay (Annett et al. 2013). Although mixing rates are probably much higher across the shelf proper, nutrients appear to be quickly taken-up nearshore prior to export offshore, with potential cross-shelf transport via particulate organic forms rather than dissolved inorganic forms of those nutrient species.
Our analysis indicates that the nitrate, ammonium and orthophosphate concentrations are statistically different between the nearshore and offshore samples in 2014, with the offshore samples having higher concentrations (Table II and Fig. 7; Fig. 1 dotted line represents nearshore/offshore delineation). This may suggest either enhanced glacial flux from more far afield, nearshore biological uptake or the influence of upwelled, more nutrient-rich water added offshore. The 2014 nearshore waters have a DIN:P:Si molar ratio of 17:1:78, while the offshore waters have 18:1:47. There has been little research on either subaerial or subglacial melt geochemistry. Surface runoff from glacial melt from Livingston Island to the north of the study area at 62°40′S has DIN:P:Si molar ratios of 5.3:1:54 (Toro et al. 2007), while the only other subglacial water analysis from the West Antarctic Ice Sheet has a ratio of 1.1:1:41 (Christner et al. 2014, Michaud et al. 2016). In comparison, surface runoff from glacial melt collected near Palmer Station has an average DIN:P:Si molar ratio of 39:1:32, while groundwater is 31:1:7 (Table I). Clearly, our surface water data are much more DIN depleted relative to both P and Si of the groundwaters and stream waters around Palmer Station. Whether this suggests preferential uptake of N in these waters or just significant geochemical variabilities in glacial melt requires further investigation. Given the observation that the means of the N:P for both the nearshore and offshore waters are close to the Redfield ratio of 16:1, these waters have probably been modified by biological processes even in close proximity to the glacial sources.

In this study, higher surface fluorescence values (2–8 mg m⁻³) were observed close to the shoreline where isotopic tracers suggest glacial melt inputs are present. Farther offshore, surface fluorescence values were lower (0–2 mg m⁻³). Given this spatial distribution, it is possible that the extent and location of phytoplankton blooms are controlled by the location, intensity and mixing of meltwater inputs along the coast. Circulation dynamics, especially those that control the distribution of meltwater, are impacted by changes in surface water temperature and salinity. As atmospheric and oceanic temperatures along the WAP rise, the volume of meltwater that enters the surface ocean will certainly increase. This change in meltwater dynamics with respect to the duration of warmer atmospheric temperatures is observed in this study over the changing seasons, although the same concepts could be applied over longer time scales.

Conclusions
Terrestrially derived water masses introduce and transport macro and micronutrients along and across the shelf of the WAP. As atmospheric temperatures along the WAP rise, the increasing frequency and intensity of meltwater pulses along the WAP have the potential to profoundly impact global carbon and climate cycles. Similar to other coastal systems around the globe, terrestrial water can be identified via natural tracer concentrations (²²³Ra, ²²²Rn, δ¹⁸O, δD). Groundwater and surface runoff from glacial meltwaters were significantly enriched in Ra and Rn relative to seawater, and depleted in ¹⁸O and D.

These tracers should vary in concentration as a function of meltwater delivery to coastal waters. Given the difference in season between the two cruises (early summer in 2012 and late summer in 2014), the higher activities in Ra and Rn observed nearshore in 2014 are attributed to an increase in meltwater production and discharge. This increased freshwater delivery was also seen in the salinity and water isotopes nearshore compared to the offshore values in 2014.

Calculated horizontal mixing rates of water masses within glacial fjords along the WAP via ²²⁴Ra and ²²²Rn ranged from 110–3600 m² s⁻¹. Mixing rates estimated here were within the range of other rates calculated along the WAP and suggest a rapid transport mechanism for moving meltwater offshore. Based on data collected offshore of Anvers Island during the 2012–13 and 2013–14 summers, the flux of biolimiting nutrients from the Antarctic continent across the continental shelf was evaluated. More work is needed to further evaluate the flux and important biogeochemical elements such as Fe to the surface waters along the WAP to gain a true understanding of how continental inputs play a role in this globally important ecological system.

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Author contributions
DRC and KN conceived the project and led field efforts. JC, RNP and LEP managed the radionuclide analysis in the field and at their respective universities. WBL analysed deuterium and ¹⁸O of all water samples at Ohio State University. All authors contributed to the discussion of results and writing of the manuscript.
