Stable isotopic composition of deep-sea gorgonian corals *Primnoa* spp.: a new archive of surface processes

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ABSTRACT: The deep-sea gorgonian coral *Primnoa* spp. live in the Atlantic and Pacific Oceans at depths of 65 to 3200 m. They have an arborescent growth form with a skeletal axis composed of annual rings made from calcite and gorgonin. Lifespans may exceed several hundreds of years. It has been suggested that isotope profiles from the gorgonin fraction of the skeleton could be used to reconstruct long-term, annual-scale variations in surface productivity. We tested assumptions about the trophic level, intra- and inter-colony isotopic reproducibility, and preservation of isotopic signatures in a suite of modern and fossil specimens. Measurements of gorgonin δ¹⁵N indicate that *Primnoa* spp. feed mainly on zooplankton and/or sinking particulate organic matter (POMsink), and not on suspended POM (POMsusp) or dissolved organic carbon (DOC). Gorgonin δ¹³C and δ¹⁵N in specimens from NE Pacific shelf waters, NW Atlantic slope waters, the Sea of Japan, and a South Pacific (Southern Ocean sector) seamount were strongly correlated with surface apparent oxygen utilization (AOU; the best available measure of surface productivity), demonstrating coupling between skeletal isotopic ratios and biophysical processes in surface water. Time-series isotopic profiles from different sections along the same colony, and different colonies inhabiting the same area were identical for δ¹³C, while δ¹⁵N profiles were less reproducible. Similarity in C:N, δ¹³C and δ¹⁵N between modern and fossil specimens suggest that isotopic signatures are preserved over millennial timescales. These results support the use of *Primnoa* spp. as historical recorders of surface water processes such as biological productivity and the isotopic composition of source nutrients.

KEY WORDS: *Primnoa* · Gorgonin · δ¹³C · δ¹⁵N · Trophic level · Paleoceanography

INTRODUCTION

*Primnoa resedaeformis* (Gunnerson) is a deep-sea gorgonian coral with known occurrences in North Atlantic, Arctic and North Pacific waters at depths of 65 to 3200 m (Breeze et al. 1997, Etnoyer & Morgan 2003). A subspecies, *P. resedaeformis notialis* (Bayer), occurs on seamounts in the Southern Ocean sector of the Pacific Ocean. *P. willeyi* (Hickson) and *P. pacifica* (Kinoshita) are found in the eastern and western North Pacific, respectively (Smithsonian holdings: http://goode.si.edu/webnew/pages/nmnh/iz/Query.php). These corals have an arborescent growth pattern with a skeleton made of calcite and a proteinaceous material called gorgonin (Goldberg 1976) arranged in alternating concentric rings around longitudinal growth axes (Risk et al. 2002, Sherwood 2002). Towards the outer growth surface of older portions of the skeleton the outer cortex may be comprised of just calcite, with gorgonin layers lacking. Based on ²¹⁰Pb-dating, Andrews et al. (2002) inferred that visible growth rings in the skeleton are formed annually. This
was subsequently validated with bomb-14C (Sherwood et al. 2005, this volume). Subannual banding patterns have also been identified using scanning electron microscopy and Nomarski differential interference imaging (Risk et al. 2002, Sherwood 2002). This coral appears to have lifespans on timescales of up to several centuries (Andrews et al. 2002, Risk et al. 2002).

Based on 14C analyses, Griffin & Druffel (1989) originally suggested that the main source of carbon to deep-sea corals was sinking particulate organic matter (POM<sub>sink</sub>). They further suggested (Druffel et al. 1995) that the carbon and nitrogen isotopic composition of the proteinaceous layers of the colonial zoanthid <i>Gerardia sp.</i> could be a recorder of surface ocean processes (productivity, nutrient sources, etc.). <i>Primnoa</i> spp. may also form their gorgonin skeleton from POM<sub>sink</sub> since δ<sup>13</sup>C and δ<sup>15</sup>N in the polyps and gorgonin show similar regional differences to δ<sup>13</sup>C and δ<sup>15</sup>N of surface water POM (Heikoop et al. 1998, 2002). Moreover, the δ<sup>13</sup>C and δ<sup>15</sup>N of the polyps are highly correlated to the δ<sup>13</sup>C and δ<sup>15</sup>N of associated gorgonin (Heikoop et al. 2002). Together, these results suggest that the isotopic composition of annual gorgonin layers could record the temporal history of processes that control the isotopic composition of POM (Heikoop et al. 2002) including plankton productivity and the δ<sup>13</sup>C and δ<sup>15</sup>N of nutrient sources (e.g. Ward-Paige et al. 2005). The δ<sup>13</sup>C and δ<sup>15</sup>N of gorgonin from a suite of corals from Alaskan waters, waters off the eastern shore board of the United States and Canada, and a South Pacific seamount were positively correlated (Heikoop et al. 2002), suggesting that surface ocean productivity relative to nutrient supply may be the primary control on the isotopic composition of gorgonin.

For stable isotope profiles generated from the annual gorgonin layers of <i>Primnoa</i> spp. skeletons to have any meaningful environmental significance, 4 conditions must be met: (1) the trophic position of <i>Primnoa</i> spp. must be known; (2) organic diagenesis must not affect the isotopic composition of the skeleton; isotopic trends must be reproducible among (3) different sections of the same colony and (4) different colonies inhabiting the same area. The purpose of this paper is to test these assumptions using a suite of recently collected live and fossil specimens from the Atlantic and Pacific Oceans.

### MATERIALS AND METHODS

Specimens were obtained during research and fishing expeditions (Table 1). In addition, 7 specimens were obtained from the Smithsonian Institution National Museum of Natural History. <i>Primnoa resedaeformis</i> (Gunnerus) was collected from the NE Channel, SW of Halifax, Nova Scotia (Fig. 1), and from east of Virginia Beach, Virginia, USA. <i>P. willeyi</i> (Hickson) was collected (m) east of Queen Charlotte Islands 1998 54.386° S 119.800° W 549

<table>
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<tr>
<th>Sample</th>
<th>Species</th>
<th>Location</th>
<th>Year collected</th>
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<th>Long.</th>
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<td>S. Pacific seamount</td>
<td>1964</td>
<td>54.817° S</td>
<td>119.800° W</td>
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*Fossil specimen, ca. 1850 AD, based on 210Pb dating; b fossil specimen, ca. 200 AD, based on U-Th dating (D. B. Scott unpubl.); c preserved in ethanol; d exact co-ordinates unknown; these are general co-ordinates for location.
son) was collected east of the Queen Charlotte Islands, Chatham Sound and Knight Inlet, British Columbia, and from the Aleutian Islands and Prince William Sound, Alaska. *P. pacifica* (Kinoshita) was collected in the Sea of Japan. The subspecies *P. resedaeformis notialis* was collected from a seamount in the Southern Ocean sector of the South Pacific. Of the Smithsonian samples, 3 were originally reported in Heikoop et al. (2002); these are included here to highlight geographic trends in stable isotopic composition.

All specimens were collected alive, except for Colonies Fossil-95 and COHPS-2001-1. These dead-collected specimens were dated radiometrically by B. Ghaleb at GEOTOP-UQAM-McGill, Montreal, Canada (Scott unpubl. data). Samples from Colony COHPS-2001-1 are ca. 1850 AD, based on $^{210}$Pb-$^{226}$Ra analyses of the outer calcite cortex region of the coral. Fossil-95 is ca. 200 AD, based on 2 uranium-series dates on the middle and outer regions of a section of the colony.

Colonies for stable isotope analyses (Fig. 2) were sectioned with a rock saw and ground and polished on a diamond lap wheel to a thickness of about 5 mm. Sections were photographed with a Nikon Coolpix digital camera in macro-mode. After some trial and

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**Fig. 1.** Gulf of Maine (GOM) region, NW Atlantic Ocean. Colonies of *Primnoa resedaeformis* were collected from the NE Channel. Symbols show locations of water samples for analysis of suspended POM (POM$_{	ext{susp}}$). GB: Georges Bank

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**Fig. 2.** *Primnoa resedaeformis*. Colony collected from the NE Channel: (a) Colony HUD-2001-055-VG15 (tissue missing); (b) Colony DFO-2002-con5 (tissue present). Boxes outline sections for isotopic time-series profiles, tissue samples. Scale ruler = 15 cm.
error we found that photographing the sections in ultra-violet light gave the clearest image of ring patterns in the horny axis owing to contrast between the calcite-rich (luminescent) and gorgonin-rich (non-luminescent) portions of the annual growth rings (Fig. 3).

Annual gorgonin rings were isolated by dissolving sections in 5% HCl for 1 wk (up to 3 wk for larger sections). Upon dissolution, sections were transferred to a Petri dish filled with distilled water, and the annual rings were teased apart with tweezers and scalpel under a binocular microscope. Photographs of the sections taken before dissolution were used to guide sampling. Individual rings were placed in 5 ml polyethylene vials with 5% HCl for an additional week to ensure that all the calcite had dissolved. After 2 more rinses in HCl, the rings were triple-rinsed in de-ionized water and dried in a low temperature oven. Rings averaged about 5 mg in weight. Tissue material scraped off the skeletal axes was prepared in the same way.

Isotopic and C:N analyses of gorgonin were performed by elemental analyzer/continuous flow isotope ratio mass spectrometry at GEOTOP-UQAM-McGill. Isotope ratios are reported in conventional delta notation, where (example for carbon): δ¹³C = [R_sample/ R_standard - 1] × 1000; and R = ¹³C/¹²C. Standards used were PDB (δ¹³C) and air (δ¹⁵N). Analytical error, as measured by the standard deviation of duplicate measurements, averaged 0.10 ‰ for δ¹³C and δ¹⁵N, and 0.01 for C:N. Some of the corals obtained from the Smithsonian Institute were preserved in ethanol, which may have affected stable isotope compositions (Bosley & Wainright 1999).

Suspended particulate organic matter (POM₄悬浮) in the Gulf of Maine/Georges Bank region (Fig. 1) was sampled as part of the Ecosystems Monitoring Survey at the Northeast Fisheries Science Center, Narragansett, Rhode Island, USA. Samples were collected in spring, summer and autumn, between November 2000 and August 2003. In addition, a vertical transect (depths of 3.7, 100 and 215 m) was sampled in August 2004 in the NE Channel (Fig. 1). Surface samples were obtained with a ship-board, near surface flow-through system. Deeper samples were collected with Niskin bottles attached to a CTD rosette. A volume of 600 to 1000 ml of water was pre-filtered through 300 µm mesh to remove most zooplankton, and then onto a GF/F filter. Samples were immediately frozen and transported to the US Environmental Protection Agency, Atlantic Ecology Division, Narragansett, for isotopic analysis. Carbon and nitrogen isotope composition was determined by continuous flow isotope ratio mass spectrometry using a Carlo-Erba NA 1500 Series II elemental analyzer interfaced to a GV Instruments Optima mass spectrometer. All samples were analyzed

Fig. 3. *Primnoa resedaeformis*. Section A5 from Colony DFO-2002-con5, photographed under ultraviolet light. Gray portion of outer section is the inorganic calcite cortex; darker layers towards middle are annual gorgonin rings isolated for chemical analyses. Based on growth ring counts, specimen is approx. 75 yr old (Sherwood et al. 2005). Scale bar = 0.5 cm
in duplicate with a typical difference of about 0.1‰. Sample material was re-analyzed periodically over a period of several months and exhibited a precision of 0.30‰, calculated as a single sigma standard deviation of all replicate values. This latter estimate of precision is appropriate for POM_{susp} δ^{15}N values determined in this study.

**RESULTS**

**Composition of tissue and skeletal gorgonin**

Stable isotope and C:N data for *Primnoa* spp. are summarized in Table 2. There were large differences in the isotopic and elemental composition between tissue and gorgonin. C:N ratios were higher and more variable in tissue (6.1 ± 1.4, n = 11) than gorgonin (3.3 ± 0.1, n = 8). δ^{13}C values were more negative in the tissue by an average of 3.0 ± 1.7‰ (n = 9). Most of this difference in δ^{13}C may be accounted for by the difference in lipid content of the 2 fractions. Taking C:N as a proxy for lipid content, lipid-normalized values of δ^{13}C (δ^{13}C') were calculated from equations in McConnaughey & McRoy (1979). After normalization, δ^{13}C' values were more negative in the tissue by 1.0 ± 0.6‰ (n = 6). Values of δ^{15}N were more positive in the tissue by 0.9 ± 0.9‰ (n = 6).

The reason for lower δ^{13}C' (even after lipid-normalization) and higher δ^{15}N in the tissue compared with the gorgonin is not clear, but may be related to differences in tissue turnover time (Tieszen et al. 1983). Each gorgonin layer integrates seasonal variations over 1 yr, and the average isotopic compositions listed in Table 2 integrate inter-annual variations over many years. We expect that the tissue, which turns over in something less than 1 yr, represents a unique seasonal signature. Another possibility relates to differences in the amino acid contents of the 2 fractions (O. A. Sherwood unpubl. data). Different amino acids are known to have unique and widely variable δ^{13}C (Keil & Fogel 2001) and δ^{15}N (McClelland & Montoya 2002); therefore, the relative proportion of amino acids between tissue and gorgonin may indicate differences in stable isotope content.

**Geographic and interspecific variability**

There were significant compositional differences among the different species and geographic areas (Table 2). *Primnoa pacifica* and *P. resedaeformis notialis* from the Sea of Japan and South Pacific had the lowest tissue C:N values (4.5 ± 0.4, n = 3). *P. willeyi* from the NE Pacific had intermediate C:N (6.1 ± 1.4, n = 5), and *P. resedaeformis* from the NW Atlantic had the highest C:N (7.5 ± 0.2, n = 3). δ^{13}C' and δ^{15}N were

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### Table 2. *Primnoa* spp. Summary of stable isotope and C:N data (mean ± SD). δ^{13}C': lipid-normalized δ^{13}C; –: insufficient material for analysis

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<th>Sample</th>
<th>n</th>
<th>C:N</th>
<th>δ^{13}C</th>
<th>δ^{13}C'</th>
<th>δ^{15}N</th>
<th>n</th>
<th>C:N</th>
<th>δ^{13}C</th>
<th>δ^{13}C'</th>
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<td>3</td>
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<tr>
<td>Smith-1010257</td>
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<td>−21.87</td>
<td>9.09 ± 0.22</td>
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<td>Smith-1010785</td>
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<td>−19.49 ± 0.15</td>
<td>−17.86</td>
<td>12.75 ± 0.18</td>
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<td>Smith-87624</td>
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<td>−20.41 ± 0.45</td>
<td>−21.56 ± 0.5</td>
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*Isotopic analyses performed at University of New Mexico; all others at GEOTOP; only 3 samples were analyzed for C:N; assumes a C:N ratio of 3.2*
positively correlated (p < 0.001; Fig. 4a), as previously found by Heikoop et al. (2002). Isotope values in $\delta^{15}N$ vs. $\delta^{13}C'$ space showed clear geographic, trends, with highest values in the NE Pacific, intermediate values in the NW Atlantic and Sea of Japan, and lowest values in the Pacific–Southern Ocean Sector (Fig. 4a). The sample from Atka Island in the NE Pacific (Colony Smith-1010257) deviated from this pattern; it had isotope values more similar to the South Pacific samples.

Isotope values were plotted against apparent oxygen utilization (AOU), selected from the Levitus & Boyer (1994) data set as the best available measure of surface productivity for open ocean, slope water and shelf sites alike (Fig. 4b). Surface water AOU data were obtained from the 0.5° latitude × 0.5° longitude grid nearest each of the coral collection locations. Both $\delta^{13}C'$ and $\delta^{15}N$ significantly increased with increasingly negative AOU (i.e. higher productivity; p < 0.0001).

**Intra-colony isotopic reproducibility**

Specimen HUD2001-055-VG15 *Primnoa resedaeformis* was the first colony examined for trends in the isotopic composition of annual gorgonin rings. This colony was collected alive and transferred to an aquarium, where the tissue layer eventually died and sloughed off the skeleton. A 50 cm long main branch was snapped off the colony for geochemical sampling. Sections for stable isotope analyses were cut from the base of the main branch, and from 2 divergent branches 10 cm higher up the colony (Fig. 2a). The basal section measured 14 mm in diameter. Stable isotope and C:N profiles from this specimen are shown in Fig. 5. The $\delta^{13}C$ profiles were identical among the 3 different sections (correlation coefficients, r, ranged 0.84 to 0.97; p < 0.0001), showing trends towards heavier values with increasing age. The amplitude of $\delta^{13}C$ profiles was 3‰, much larger than the analytical error (0.10‰). $\delta^{15}N$ profiles were less reproducible between the different sections (r = 0.38 to 0.48; p = < 0.01 to 0.05). Differences among coeval rings (up to 1.5‰) were equivalent to the amplitude of the profiles, and cannot be explained by analytical error. C:N profiles were the least reproducible among the different sections (r = –0.23 to 0.44; p = 0.02 to 0.78), with differences among coeval rings (up to 0.07) exceeding the analytical error (0.01).

A larger colony of *Primnoa resedaeformis* (DFO-2002-con5) was subsequently obtained from the Canadian Department of Fisheries and Oceans and the same experiment was repeated. This colony was completely covered in live tissue and was frozen immediately after collection (Fig. 2b). Sections for geochemistry were thawed and air dried in the laboratory. The colony measured 70 cm in length and had a diameter of 26 mm at the base, which included a thick accumulation of calcite cortex (Fig. 3). Analyses of 5 tissue samples taken along the length of Specimen DFO-2002-con5 showed no difference in $\delta^{13}C$, within analytical error (Fig. 6). The exception was the tissue sample 20 cm from the top of the colony, which had a value 0.3‰ heavier than the rest. $\delta^{15}N$ was slightly more variable, with values increasing steadily by 0.8‰ from the tip to the base of the colony. C:N varied by up to 2 among the different tissue samples. Stable isotope
profiles (Fig. 7) of gorgonin layers were generated from 2 sections at the base of the colony, separated by a distance of 3 cm. C:N was not measured. Isotope profiles were virtually identical among the 2 different sections (r = 0.69 [δ¹³C] and 0.63 [δ¹⁵N]; p < 0.0001).

Inter-colony isotopic reproducibility

Inter-colony isotope reproducibility was assessed among 3 different colonies (Fig. 8). For each colony, ring numbers were converted to calendar ages as described in Sherwood et al. (2005). Briefly, photographic prints of each section were circulated among 3 amateur counters, and the average of the 3 age determinations was calculated. Ring counts were validated by measurements of bomb-¹⁴C in each of the sections (Sherwood et al. 2005). The inter-colony results paralleled intra-colony results, with excellent reproducibility of δ¹³C and poorer reproducibility of δ¹⁵N (Fig. 8). Correlation of δ¹³C time-series among the different colonies was highly significant (r = 0.67 to 0.86; p < 0.0001). Correlation of δ¹⁵N was significant between Colonies HUD-2001-055-VG15 and ROPOS-639009 (r = 0.44; p < 0.05), but was insignificant in the other 2 cases.

Comparison of modern and fossil specimens

The 2 specimens from ca. 200 AD and 1850 AD were compared to other corals collected alive from the NE
Channel (Table 2). C:N ratios of these older specimens, 3.4 ± 0.1 (n = 9), were slightly higher than the modern ones, 3.15 ± 0.05 (n = 260). The fossil samples were heavier in δ\(^{13}\)C (lipid normalized to account for difference in C:N) by 1.5‰ (Fig. 9). Part of this difference may be due to the Suess effect, caused by the depletion of atmospheric \(^{13}\)C from the burning of fossil fuels, with subsequent depletion of oceanic dissolved inorganic carbon (DIC; Quay et al. 1992). The 1.5‰ difference between the 150 yr old Colony COHPS-2001-1 and the most recent samples from Colony DFO-2002-con5 is consistent with the decrease in atmospheric δ\(^{13}\)C between the mid-1800s and the present (Francey et al. 1999). The lighter values from ca. 1920, however, exceed the magnitude of the Suess effect. Therefore, there may be other oceanographic and/or trophic level changes affecting δ\(^{13}\)C over shorter timescales. There was also a slight trend towards higher δ\(^{15}\)N with increasing age, with the 2 fossil specimens having values similar to the oldest layers from, Colony DFO-2002-con5 (Fig. 9).

δ\(^{15}\)N composition of plankton

To assess the trophic level (TL) of Primnoa resedaeformis collected from the NE Channel (see next subsection), δ\(^{15}\)N at the base of the food web was assessed from measurements of POM\(_{susp}\). Unfortunately, the NE Channel was not targeted for sampling in the years 2000 to 2003 of the Ecosystems Monitoring Survey (Fig. 1). With the exception of the central shoals of Georges Bank, δ\(^{15}\)N in surface water POM\(_{susp}\) was consistent throughout the entire region (4.1 ± 1.2‰, n = 56; Fig. 10). We therefore assume that this value is representative of POM\(_{susp}\) in waters overlaying the NE Channel.

In August 2004, the NE Channel was occupied to collect POM\(_{susp}\) along a depth transect (Fig. 1). Below the euphotic zone δ\(^{15}\)N increased rapidly to a maximum of 18.5‰ at 215 m depth (Fig. 11). While these data represent only 1 snapshot in time, they are consistent with earlier results reported for Wilkinson Basin, located farther inside the Gulf of Maine (Libes & Deuser 1988).
DISCUSSION

Trophic level

It has been demonstrated that deep-sea corals form their organic endoskeletons from sinking POM (POMsink), based on the presence of modern $\Delta^{14}C$ ($>0\‰$) in the gorgonin fraction of their skeletons (Griffin & Druffel 1989, Heikoop et al. 1998, Roark et al. 2005, Sherwood et al. 2005). Reconstruction of bomb-$^{14}C$ from the skeletons of *Primnoa resedaeformis* collected from the NE Channel (Sherwood et al. 2005) was identical to other proxy records derived from a mollusc shell (Weidman & Jones 1993) and haddock otoliths (Campana 1997). Moreover, the timing of the initial rise and peak in bomb-$^{14}C$ was in phase with direct measurements of seawater DIC (Nydal 1998). Therefore, the gorgonin fraction is derived from surface water DIC, transmitted to depth via the plankton food web; but these results do not indicate whether *Primnoa* spp. feed upon sinking phytoplankton, zooplankton or other trophic intermediaries. Because of trophic level fractionation of $\delta^{13}C$ and $\delta^{15}N$, interpretation of time series isotope profiles requires knowledge of the TL.

Further insight into the TL of *Primnoa* spp. is provided by $\delta^{15}N$. This is typically enriched in a consumer relative to its diet by an average enrichment factor of $\Delta \delta^{15}N = 3.4\‰$ (DeNiro & Epstein 1981, Minagawa & Wada 1984, Vander Zanden & Rasmussen 2001). The TL of *Primnoa* spp. may therefore be estimated by comparing our data with the isotopic signatures of other organisms of known TL (e.g. Vander Zanden et al. 1997, Polunin et al. 2001). We used primary consumers as the baseline, and calculated TL by the formula: $\text{TL}_{\text{consumer}} = (\delta^{15}N_{\text{consumer}} - \delta^{15}N_{\text{baseline}}) / 3.4 + 2$ (Vander Zanden & Rasmussen 2001). Wherever possible, we used the gorgonin, rather than tissue results, since these integrate seasonal and inter-annual isotope variations occurring at the base of the food web. Our approach was to reconstruct regionally-specific TL models from literature data.

*Primnoa resedaeformis* from the NE Channel was compared with the TL model of Fry (1988), which was based on taxa collected from nearby Georges Bank. We added to this model isotopic data for POM$_{\text{sink}}$ (Macko 1981) and size-fractionated zooplankton (Fry & Quinones 1994). Most of the invertebrates and fish reported in Fry (1988) and Fry & Quinones (1994) were collected within the 100 m isobath on Georges Bank, where $\delta^{15}N$ of POM$_{\text{sink}}$ is 1.4‰.
heavier than in surrounding waters (Fig. 10), probably as a result of greater use of regenerated ammonium on the bank (Ostrom et al. 1997, Wu et al. 1999a). We assume that isotopic enrichment on Georges Bank is transmitted to higher trophic levels. This difference in δ15N equates to 0.4 trophic levels and must be accounted for in the TL model output.

The NE Channel/Gulf of Maine δ15N-TL model is shown in Fig. 12. We subtracted 1.4‰ from the data for the Georges Bank taxa. Herbivorous scallops were used for baseline δ15N (Fry 1988; 6 – 1.4 = 4.6‰). Among live-collected Primnoa resedaeformis from the NE Channel, the inter-colony average δ15N was 10.0 ± 0.3‰ (n = 5). This value is similar to the δ15N for large benthic isopods, large polychaetes and planktivorous fishes (Fig. 12). The calculated TL is 3.6. This suggests that P. resedaeformis is primarily carnivorous. We have also observed that the polyps on P. resedaeformis point downwards, suggesting that these corals may also feed on resuspended meiofauna.

Another factor which may affect the TL estimate is isotopic modification of particulate matter in deep waters. Significant enrichment of δ15N below the euphotic zone (Fig. 11) rules out the possibility that Primnoa resedaeformis feeds on the highly degraded POMsusp encountered at depth (otherwise δ15N in P. resedaeformis would be much higher; Fig. 12). POMsink, may also become isotopically enriched below the euphotic zone, although to a much lesser extent than POMsusp (Altabet 1988, Altabet et al. 1991, Voss et al. 1996). We estimate that POMsink in the NE Channel has a δ15N signature of 6.5‰ based on sediment data (Macko 1981), and general similarity in δ15N between sediments and POMsink (Altabet & Francois 1994, Voss et al. 1996, Ostrom et al. 1997). Assuming ΔΔδ15N = 3.4, P. resedaeformis could feed on POMsink as well as zooplankton (Fig. 12).

Heavier δ15N in Primnoa willeyi from the NE Pacific (12.8 ± 0.6‰, n = 5), compared to P. resedaeformis from the NW Atlantic, may reflect either a difference at the base of the food web or higher TL. We constructed a TL model based on literature δ15N data from 2 inshore Bays near Juneau (Goering et al. 1990) and Prince William Sound, Alaska (Kline 1999; note that 1 of our specimens [Smith-51283] was collected in Prince William Sound). Baseline δ15N was set to 8‰, using the values reported for herbivorous copepods (Neocalanus cristatus; Kline 1999) and bivalves (Goering et al. 1990). Overlap in δ15N values among similar taxa from these 2 Alaskan bays, despite a separation of over 600 km, lends confidence that δ15N signatures are well-conserved within the coastal NE Pacific ecosystem. We calculated a TL of about 3.4 for P. willeyi. Measurements of POMsink from off Vancouver Island (δ15N = 8 to 9‰, Peña et al. 1999, Wu et al. 1999a) are also consistent with this fraction being a food source for P. willeyi. Lighter δ15N in the sample from Atka Island (9.1‰) may be explained by low δ15N in offshore primary producers (Wu et al. 1997; see next paragraph).

It has been shown that coastal ecosystems often exhibit lower ΔΔδ15N than the globally accepted value of 3.4‰ (Wu et al. 1997, Sherwood & Rose 2005). To validate our TL estimates, we also looked at 4 taxa that were sampled in both the NE Pacific and NW Atlantic: euphausiids, bivalves, pollock and sole. TL outputs...
for the 2 different ecosystems were statistically equal (matched-pairs t-test). Moreover, TL outputs for both ecosystems conformed to expectations: bivalves (TL = 2); euphausiids (TL = 2.6); pollock (3.8); sole (3.5 to 3.8). Therefore, our use of Δδ15N = 3.4‰ for both Georges Bank and coastal NE Pacific ecosystems does not appear to introduce bias in our model outputs.

In summary, isotopic data support the following 3 conclusions about the diet of Primnoa spp. corals: (1) The presence of modern Δ14C in Primnoa resedaeformis rules out DOC and DIC as a significant carbon source for the gorgonin fraction of the skeleton (Sherwood et al. 2005. Measurements of δ15N suggest that (2) zooplankton and/or POMsank constitute the main diet of P. resedaeformis and P. willyei, and (3) the highly degraded fraction of POMsusp found at depth is not a significant food source. This is also supported by the finding that the dark, more gorgonin-rich portion of the annual ring couplets in P. resedaeformis co-incides with the timing of the spring/summer bloom of phytoplankton and zooplankton (Sherwood 2002).

Lack of isotopic data for the Sea of Japan and the South Pacific in the literature prevented a similar analysis of the TL of the remaining species. It is quite probable that Primnoa pacifica and P. resedaeformis notialis share a similar type of diet with P. resedaeformis and P. willyei, since all the corals have similar-sized polyps.

As passive suspension feeders, octocorals feed opportunistically on a wide spectrum of plankton size-classes, from nanoeukaryotes to zooplankton (Ribes et al. 1999, 2003, Orejas et al. 2003). Temperate and boreal asymbiotic species feed mainly on zooplankton and detrital POM in about equal proportions, with smaller plankton (<100 µm) accounting for <10% of energy demand (Ribes et al. 1999, 2003). Among zooplankton, gorgonians ingest smaller, low-motility prey items (Coma et al. 1994, Rossi et al. 2004). Our isotopic data conform to this general pattern, with the exception that the highly degraded POMsusp found at depth does not appear to be a significant food source. Heterogeneity of prey items may explain the differences in the intra- and inter-colony reproducibility of δ13C and δ15N. It may be possible that different parts of the colony are more effective at capturing different-sized prey items, depending on localized current regimes (Coma et al. 1994). Differential feeding on prey size-classes could potentially alter δ15N signatures, due to strong trophic level fractionation of 15N (Fry & Quinones 1994). If this is true, then analyses of more sections per colony and more colonies per site are recommended to calculate the average δ15N. Since trophic level fractionation of 13C is much weaker (generally <1‰; Vander Zanden & Rasmussen 2001), differential feeding should exert less influence on δ13C, thereby lending greater isotopic reproducibility.

### Surface–benthic coupling of isotopic signatures

The overall isotopic signature of a food web is determined by biophysical processes at the level of primary producers. These processes are myriad and complex, especially over shorter timescales. Phytoplankton δ15N mainly depends on the efficiency of nitrogen utilization (Nakatsuka et al. 1992, Altabet & Francois 1994, Wu et al. 1997, 1999a) and on the δ15N signature of the nitrogenous substrate (Ostrom et al. 1997, Altabet et al. 1999). Phytoplankton δ13C mainly depends on [CO2] (Rau et al. 1992, Hoffman et al. 2000), growth rate (Nakatsuka et al. 1992, Hoffman et al. 2000) and cell geometry/plankton species composition (Fry & Wainright 1991, Popp et al. 1998), as well as the δ13C signature of the bicarbonate substrate (Cullen et al. 2001). As a result of these myriad factors, there are large isotopic differences between open ocean, slope water and coastal ecosystems (Wu et al. 1997). Furthermore, different pathways of isotopic fractionation often lead to decoupling of δ13C and δ15N in new production (Ostrom et al. 1997, Wu et al. 1999a,b). In Fig. 4b, our use of surface water AOU is not meant to imply control on isotopic fractionation, but to give some sense of the different surface productivity regimes where the corals lived.

The lightest values were found in the South Pacific, located near a high-nutrient low-chlorophyll (HNLC) domain, where despite high [NO3] and [CO2], primary production is limited by micronutrients such as iron. Similarly low values were observed in a specimen from Atka Island, in the Aleutian Islands, also located near an HNLC domain (Wu et al. 1999b). Heavier values were found in the slope water regions of the NW Atlantic and the Sea of Japan, where primary productivity is typically dominated by blooms of large, fast-growing phytoplankton (Mousseau et al. 1996). These blooms lead to nutrient depletion and heavier δ13C and δ15N values. The heaviest values were in corals from Alaska and British Columbia, consistent with highest productivity rates in these coastal domains. These results demonstrate that isotopic signatures originating from primary producers are transmitted to, and preserved within, the organic endoskeletons of Primnoa spp. It is therefore not surprising that δ13C and δ15N in Primnoa spp. are both so tightly correlated with AOU (Fig. 4b).

### Preservation of original isotopic composition

For isotopic trends from gorgonin to have any paleoceanographic utility, there must not be any diagenetic overprinting. In many of the fossil specimens donated to us by fishermen, the gorgonin appeared to be more susceptible to degradation than the inorganic calcite cortex fraction. On the broken axes of these dead colonies, the
inner horny axis is often worn down in a smooth cup-shaped depression, while the cortex remains intact. Similarity in C:N, δ^{13}C and δ^{15}N between modern and fossil specimens suggests that isotopic signatures are preserved from the time of original formation. This is also supported by identical amino acid abundances between modern and fossil specimens (O. A. Sherwood unpubl.), since diagenesis often leads to the synthesis of microbial biomass of different amino acid and stable isotope composition (Macko & Estep 1984). The degradation observed in fossil specimens appears to be the result of mechanical erosion, probably by the action of suspended sands, rather than organic diagenesis. This finding is consistent with gorgonin being one of the most chemically inert proteins known (Goldberg 1976). Over millennial timescales, therefore, isotope abundances in Primnoa spp. are preserved, making these corals durable archives of paleoceanographic information.

**Paleoceanographic applications**

Owing to little trophic level fractionation and excellent intra- and inter-colony reproducibility, δ^{13}C time-series from Primnoa spp. could reliably track variations in surface processes. From Fig. 8, it appears that the 3 colonies of *P. resedaeformis* from the NE Channel recorded a peak in δ^{13}C around the mid-twentieth century, with several decade-scale oscillations. The causes of these variations are not clear, but there may be an important link with known changes in plankton community composition since the 1960s in this region (Sameoto 2001). On the other hand, evidence of the oceanic Suess effect on the δ^{13}C composition of modern vs. fossil specimens (Fig. 9), and reconstruction of 20th century bomb radiocarbon (Sherwood et al. 2005) are examples of isotopic variability in source materials recorded in these corals. Therefore, there is evidence that both biological processes (i.e. correlation of δ^{13}C and δ^{15}N with AOU) and physical processes (changes in isotopic composition of bicarbonate substrate) are reflected in gorgonin isotope content. Smaller regional studies with multiple colonies and good chronological control will be required to deconvolute these different factors. Useful information may also be extracted from δ^{15}N profiles, provided that time-series variability is large relative to the intra- and inter-colony variability.

The annual nature of ring formation makes *Primnoa* spp. analogous to varved sediment cores, from which much useful paleoceanographic information has been retrieved (e.g. Tunnicliffe 2000). In similar fashion, annually resolved isotope time-series have been generated from preserved animal remains, such as fish scales (Wainright et al. 1993) and whale baleen (Schell 2001). Long lifespans (at least 400 yr, probably longer; Risk et al. 2002) lend to *Primnoa* spp. the advantages of both varved sediment cores (longer, *in situ* time-series) and preserved animal remains (widespread distribution, known TL). Isotopic reconstructions from *Primnoa* spp. could be useful in illustrating temporal variations in marine productivity, as well as tracking the relative importance of top-down vs. bottom-up influences over time (Schell 2001, Satterfield & Finney 2002, Rau et al. 2003). Oceanographic phenomena that may be recorded include changes in the position of water mass boundaries, upwelling strength, terrestrial nutrient inputs and atmospheric nutrient inputs.

Deep-sea corals could prove to be temporal and spatial recorders of the efficacy of the oceanic biological pump that transfers carbon dioxide from the atmosphere to the deep-ocean. By understanding natural processes that have affected the operation of this pump over century timescales, we can better predict the effects of global change and potential engineered approaches such as iron fertilization on oceanic carbon sequestration. The widespread occurrence and diversity of deep-sea corals is now only being fully appreciated. As more deep-sea corals are discovered in important oceanographic regions the applicability of this potential paleoceanographic archive is likely to increase.

**CONCLUSIONS**

Measurements of δ^{15}N indicate that *Primnoa resedaeformis* and *P. willeyi* have a TL of about 3.5. POM_{sink} and zooplankton appear to constitute the bulk of their diet, whereas DIC, DOC and POM_{susp} are not consumed. The TL for *P. pacifica* and *P. resedaeformis notialis* could not be determined but, based on similar sized polyps, it is likely that these species feed at a similar TL.

Average δ^{13}C and δ^{15}N compositions of the gorgonin fraction were strongly correlated with each other and with surface water AOU. This demonstrates strong coupling between surface biophysical processes and stable isotope compositions in *Primnoa* spp.

Isotope profiles from annual gorgonin rings showed excellent intra-colony and inter-colony reproducibility for δ^{13}C, while for δ^{15}N the reproducibility was not as good. The latter result may arise from differential feeding upon different sized prey items, depending on localized current regimes; however, more work is needed to address this issue.

Similarity in C:N, δ^{13}C and δ^{15}N between modern and fossil specimens demonstrates a lack of organic diagenesis in the tough gorgonin fraction. Isotopic signatures from the time of formation are therefore preserved over millennial timescales, making these corals excellent candidates for retrospective studies of the surface marine environment.
Acknowledgements. For providing coral samples, we are thankful to S. Atwood, L. Buhl Mortensen, S. Cairns, D. Gordon, D. Jones, D. Mackas, P. Mortensen, A. Muir, L. Talley, V. Tunncliffe, and R. Wylkins. J. McKay performed isotopic analyses at GEOTOP. We are grateful to B. Ghalay for providing radiometric dates. Stable isotopic analyses at UNM were performed by V. Atudorei, M. Hess, and Z. Sharp. We thank Jerry Preziosio of NOAA NMFS, Narragansett, Rhode Island, for kindly providing POM samples. Lawrence Plug offered valuable help with digital mapping. We also thank 3 anonymous reviewers for suggestions leading to improvement of this manuscript. Funding was provided by an NSERC Strategic Grant to M.J.R. and D.B.S., an Institute of Geophysics and Planetary Physics, Los Alamos National Laboratory, grant to J.M.H., and an NSERC postgraduate scholarship to O.A.S. Radiocarbon analyses were performed under the auspices of the US Department of Energy by the University of California Lawrence Livermore National Laboratory (contract W-7405-Eng-48).

LITERATURE CITED


Mousseau L, Legendre L, Fortier L (1996) Dynamics of size-fractionated phytoplankton and trophic pathways on the


Submissions: October 21, 2004; Accepted: April 21, 2005
Proofs received from author(s): September 5, 2005

Editorial responsibility: Otto Kinne (Editor-in-Chief), Oldendorf/Luhe, Germany