

TRADITIONAL AND EMERGING GEOCHEMICAL PROXIES IN FORAMINIFERA

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ABSTRACT

Geochemical analyses of the carbonate tests calcified by foraminifera have provided much of the foundation for reconstructions of past ocean and climate conditions, and for chemostratigraphy. In particular, reconstructions of climate history (including temperature, salinity, and ice volume), ocean paleocirculation patterns, the carbon cycle, paleoproductivity, marine carbonate chemistry, and chemostratigraphy have relied on measurements of isotopic and trace element composition of foraminiferal calcium carbonate, and variations in these geochemical records through time and space. Substantial work has been done on details of traditional proxies (e.g., $\delta^{18}\text{O}$, $\delta^{13}\text{C}$) and on emerging proxies (e.g., $\delta^{11}\text{B}$, ϵ_{Nd}) in recent years; hence, a new overview of these proxies provides a timely reference and educational tool. We review the geochemical proxies that utilize foraminiferal carbonate tests, including potential uses of the proxies for reconstructions through time: $\delta^{18}\text{O}$, $\delta^{13}\text{C}$, trace elements (Mg, Cd, Ba, Zn, B, U), $^{87}\text{Sr}/^{86}\text{Sr}$, $\delta^{26}\text{Mg}$, $\delta^{11}\text{B}$, and ϵ_{Nd} . Both planktic and benthic foraminifera are included; planktic foraminifera provide information on the upper few hundred meters of the surface ocean, whereas benthic foraminifera provide information on conditions at the seafloor and in shallow porewaters, from shallow seas to deep ocean basins.

INTRODUCTION

The development of myriad geochemical proxies and refinement of analytical techniques applied to calcareous foraminifera has exploded in recent decades, and has greatly enhanced our abilities to reconstruct a wide spectrum of climate and oceanographic parameters from the geologic record. Recent advances in the uses and understanding of both traditional proxies (e.g., $\delta^{18}\text{O}$, $\delta^{13}\text{C}$) and emerging proxies (e.g., $\delta^{11}\text{B}$, ϵ_{Nd}) warrant a new review that encompasses multiple geochemical proxies in foraminifera, from fundamentals to recent developments. Here, we review these proxies, including systematics and applications (Table 1). This paper provides a timely reference and educational tool, including overview, detail, and citations that can be pursued for further investigation. It is written for a wide audience, from students who need a broad-based background to specialists interested in exploring different proxies. We focus on open-ocean and deep-ocean applica-

tions, and note that these proxies can also be applied to more restricted environments such as coastlines and methane seeps.

A combination of parameters such as water chemistry, biological processes, and habitat preferences ultimately determine the geochemical signatures recorded in foraminiferal calcite. Understanding the impact of these factors on foraminiferal carbonate chemistry is critical to data interpretations. Therefore, we begin this review with a summary of these parameters.

BENTHIC FORAMINIFERA

Living benthic foraminifera are most abundant in the upper 1 cm of sediment, including the sediment surface, which is referred to as an epifaunal microhabitat. Benthic foraminifera may also live deeper in the sediment (most abundant in the upper few cm; occasionally as deep as ~20 cm), which is referred to as an infaunal microhabitat (e.g., Corliss, 1985; Gooday, 1986; Rathburn and Corliss, 1994; Jorissen and others, 1995; de Steiger and others, 1998). Benthic foraminifera may also reside in elevated habitats, such as attached to sea grass and coral, or within phytodetritus that accumulates on the seafloor. Benthic foraminifera may change microhabitat depth slightly (usually in response to food abundance or oxygen availability), or burrow up/down during a life cycle (Bender and others, 1985; Jorissen and others, 1992). Because of differences in microhabitats, the chemistry of epifaunal tests is thought to record water properties at the seafloor, whereas the chemistry of infaunal tests is thought to record porewater properties. Epifaunal genera most often used in geochemical analyses are *Cibicidoides*, *Nuttallides*, *Oridorsalis*, and *Planulina*. The most commonly used infaunal genus is *Uvigerina*.

Even before the widespread use of geochemical techniques, foraminiferal faunal distributions were used to reconstruct paleoenvironmental conditions and paleocirculation changes. This technique has been employed with varying degrees of success with benthic foraminifera, in large part because multiple environmental factors can influence the distributions of different species. The primary controls on these distributions are food availability and/or oxygen concentration, and hence, the abundances of certain benthic foraminifera have been used to reconstruct past food and oxygen availability (e.g. Lutze, 1978; Schnitker, 1979; Miller and Lohmann, 1982; Katz and Miller, 1993; Gooday, 1994; Rathburn and Corliss, 1994; Smart and others, 1994; Jorissen and others, 1995; Thomas and Gooday, 1996; Bernhard and others, 1997; Schmiedl and others, 1997; Cannariato and Kennett, 1999; Gooday and Rathburn, 1999; van der Zwaan and others, 1999; Kaiho, 1999.; Schmiedl and others, 2000; Morigi and others, 2001).

Streeter (1973) and Schnitker (1974) linked benthic foraminiferal species occurrences to physiochemical proper-

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TABLE 1. Geochemical proxies applied to calcareous foraminifera.

	$\delta^{18}\text{O}$	$\delta^{13}\text{C}$	$\delta^{11}\text{B}$	$^{87}\text{Sr}/^{86}\text{Sr}$	Mg/Ca	Cd/Ca	Ba/Ca	Zn/Ca	B/Ca	U/Ca	$\delta^{11}\text{Nd}$	$\delta^{26}\text{Mg}$
ice volume	x											
temperature	x				x							
circulation						x	x	x			x	
productivity		x										
carbonate saturation								x	x	x		
pH												
chemostratigraphy	x	x	x									
seawater [Mg]				x								x

ties of different water masses in the Atlantic; this type of association has been utilized to reconstruct changes in water-mass structure and circulation through time (e.g., Lohmann, 1978; Corliss, 1979b, a; Schnitker, 1979; Streeter and Shackleton, 1979; Murray, 1984; Boersma, 1985; Miller and Katz, 1987; Mackensen and others, 1993a; Murray, 1995; Schmiedl and Mackensen, 1997; Gooday, 2003; Katz and others, 2003b). The increasing use of isotope and trace element geochemical proxies has greatly enhanced the utility of benthic foraminifera in these types of studies.

PLANKTIC FORAMINIFERA

The distribution of planktic foraminiferal species in the modern ocean tends to correlate with temperature, with certain species more abundant in warm waters, while others are more abundant in cooler waters (Bradshaw, 1959; Bè and Tolderland, 1971; Imbrie and Kipp, 1971; Bè, 1977; Rutherford and others, 1999). As with other ecosystems, the greatest diversity in the planktic foraminiferal assemblages is in the tropics. Diversity decreases towards the poles with polar assemblages dominated by *Neogloboquadrina pachyderma* (Ehrenberg). The CLIMAP (Climate: Long-Range Investigation, Mapping, and Prediction) project exploited this relationship, producing the first comprehensive syntheses of sea surface temperatures (SSTs) for the last glacial maximum (CLIMAP, 1976, 1981). Detailed mathematical relationships between planktic foraminifera and SSTs were developed further in subsequent studies (e.g., Mix and others, 1986; Mix and others, 1999). Additional studies have used planktic foraminiferal assemblages to infer relative temperature changes during the Neogene (e.g., Thunell and Belyea, 1982; Kennett and others, 1985; Wright and Thunell, 1988). Only relative SSTs can be estimated for Miocene and older assemblages because many of the species comprising these assemblages are extinct.

The wide distribution of planktic foraminifera in surface waters and the calcitic mineralogy of their tests make them an attractive group for geochemical analysis. However, while planktic foraminiferal thanatocoenoses (fossils in an assemblage that may or may not have lived at the same time) can be related to the overlying surface-water conditions, large physical and chemical gradients exist in the upper water column, requiring knowledge of a species' preferred depth habitat and/or seasonal biases to make more informed interpretations of geochemical data (e.g., Fairbanks and Wiebe, 1980; Fairbanks and others, 1980). In many locations, this seasonal and/or vertical variability can be greater than the long-term climate signals to be ascertained.

Planktic foraminiferal assemblages reflect food availability in addition to water temperatures, which is a factor in determining different water depth habitats of different species (Berger and others, 1978; Fairbanks and others, 1982; Curry and others, 1983; Thunell and Reynolds, 1984; Ravelo and Fairbanks, 1992). Taxa are characterized as shallow, intermediate/thermocline, or deep dwellers, and use two feeding strategies to exploit the physiochemical differences in each niche. The surface mixed layer tends to have lower nutrient levels because most of the nutrients are quickly scavenged. As a result, shallow assemblages are dominated by symbiont-bearing species such as *Globigerinoides sacculifer* (Brady) and *Globigerinoides ruber* (d'Orbigny). These species adopt a dual feeding strategy such that when food is available (e.g., copepods), they capture prey (Anderson and Bè, 1976). The photosynthetic activity of the symbiotic algae within the foraminiferal protoplasm provides supplemental food for the host (Anderson and Bè, 1976). Planktic foraminifera with symbionts are predominantly spinose and maintain a relatively shallow position in the water column, presumably to maximize photosynthesis (Fairbanks and others, 1980; Bè and others, 1982; Fairbanks and others, 1982).

The majority of non-spinose planktic foraminifera rely on food within the water column. One of the most important hydrographic controls on the availability of food is the depth and intensity of the seasonal thermocline. This steep change in temperature in the upper water column creates a steep density gradient that concentrates food particles falling from the surface. When the thermocline occurs in the euphotic zone (i.e., the upper ~100 m of the water column), both high light intensity and high nutrient levels support high levels of primary production. The resulting phytoplankton biomass provides a food-rich environment that can support a large zooplankton biomass, including planktic foraminifera. The richest diversity and abundance of planktic foraminifera occurs when the thermocline is in the euphotic zone (Fairbanks and Wiebe, 1980; Ortner and others, 1980).

STABLE ISOTOPES ($\delta^{18}\text{O}$ AND $\delta^{13}\text{C}$)

OXYGEN ISOTOPES ($\delta^{18}\text{O}$)

The deviation from a standard of the ratio of isotopes of oxygen ($^{18}\text{O}/^{16}\text{O}$) can be measured in the CaCO_3 tests of planktic and benthic foraminifera, and is reported in “ δ ” notation, calculated as follows:

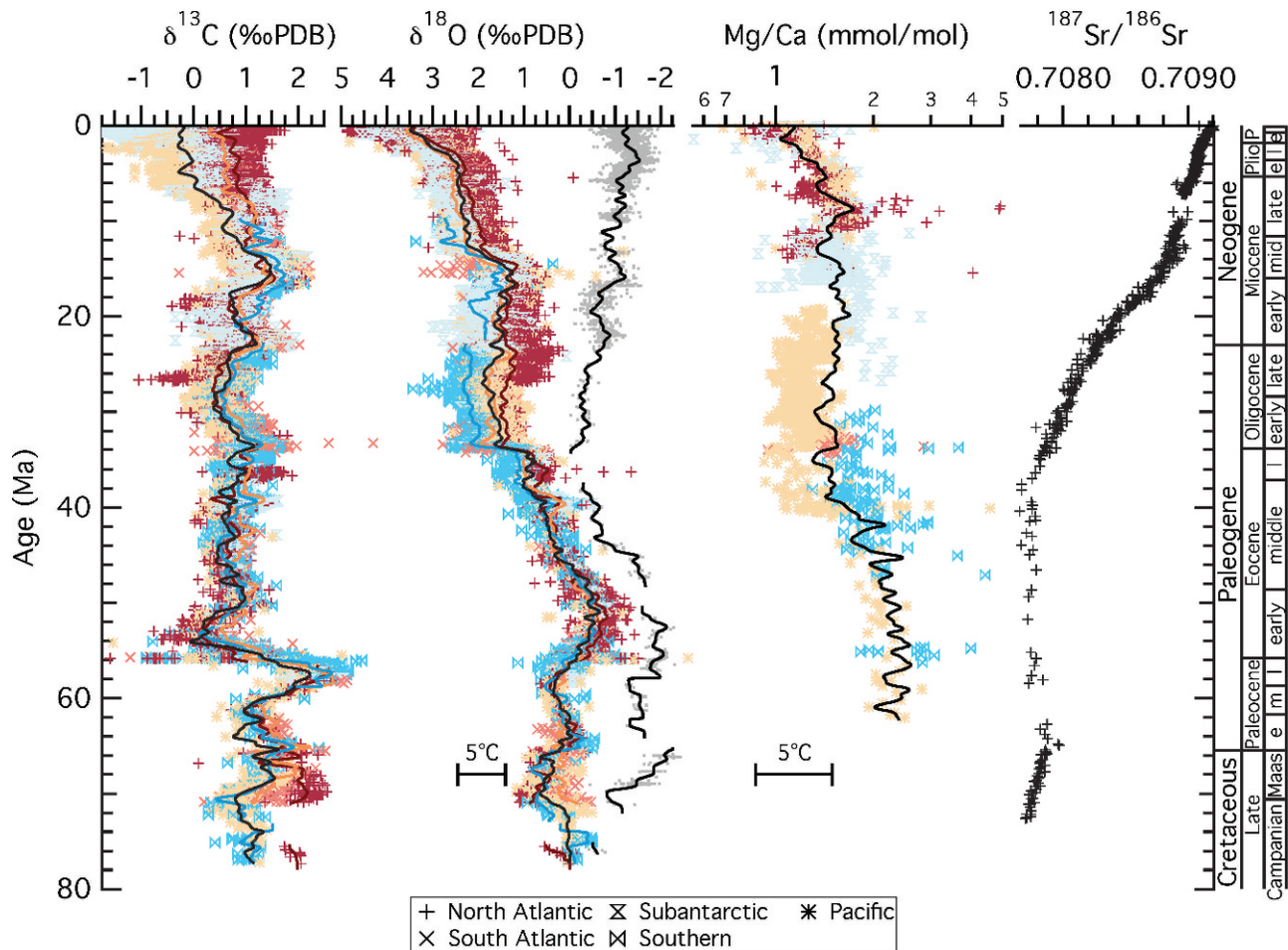


FIGURE 1. Compilations of geochemical proxies measured on foraminifera. (a) benthic foraminiferal $\delta^{13}\text{C}$; (b) benthic (colors) and planktic (gray with black trend line) foraminiferal $\delta^{18}\text{O}$; (c) benthic foraminiferal Mg/Ca, and (d) foraminiferal $^{87}\text{Sr}/^{86}\text{Sr}$. Separate trends are shown to highlight interbasinal gradients in the stable isotope records (black: tropical Pacific; dark red: North/Equatorial Atlantic; orange: South Atlantic/Subantarctic; blue: high latitude Southern Ocean). The equivalent in $\delta^{18}\text{O}$ and Mg/Ca of a 5°C temperature change is based on calibrations from Bemis and others (1998) and Lear and others (2002); note that the temperature signal in both records is complicated by variations in seawater chemistry, as discussed in the text. Benthic foraminiferal stable isotope compilations and trends are from Cramer and others (2009); planktic foraminiferal stable isotope compilation is from multiple sources (Wright 2001; Huber and others, 2002; MacLeod and others, 2005); Mg/Ca compilation includes data from multiple sources (Lear and others, 2000; Billups and Schrag, 2002, 2003; Lear and others, 2003; Lear and others, 2004; Dutton and others, 2005; Shevenell and others, 2008); $^{87}\text{Sr}/^{86}\text{Sr}$ compilation includes data from multiple sources (DePaolo and Ingram, 1985; Hess and others, 1986; McKenzie and others, 1988; Miller and others, 1988; Hodell and others, 1991; Miller and others, 1991a; Hodell and Woodruff, 1994; Oslick and others, 1994; Farrell and others, 1995; Reilly and others, 2002; Hodell and others, 2007). The procedure used to calculate the benthic foraminiferal $\delta^{18}\text{O}$ trend lines is described in Cramer and others (2009); the same procedure was used to calculate the planktic $\delta^{18}\text{O}$ and benthic Mg/Ca trend lines shown here.

$$\delta^{18}\text{O}_{\text{sample}} = 1000 \times \left[\frac{(^{18}\text{O}/^{16}\text{O})_{\text{sample}} - (^{18}\text{O}/^{16}\text{O})_{\text{standard}}}{(^{18}\text{O}/^{16}\text{O})_{\text{standard}}} \right]$$

The oxygen isotopic composition of the foraminiferal test reflects the $\delta^{18}\text{O}$ of the seawater (δ_w) in which the foraminifera calcify, as well as the temperature-dependent fractionation during the calcification process. In turn, δ_w is a function of: 1) global ice volume; and 2) local/regional river-water input and evaporation/precipitation patterns for shelf and surface waters, especially towards high latitudes. This is because the fractionation during evaporation (H_2^{16}O is more readily evaporated) produces ^{16}O -enriched precipitation over land, resulting in (1) ^{16}O -enriched glacial ice (and to a lesser extent, lakes and groundwater) sequestered on land; and (2) ^{16}O -enriched river runoff to the oceans.

$\delta^{18}\text{O}$, along with $\delta^{13}\text{C}$ measured in the same analysis, was the first geochemical proxy routinely measured in foraminifera (Fig. 1). It remains a central tool for paleoceanographic reconstructions, being widely used as a paleothermometer, ice-volume indicator, and sea-level indicator. While several paleotemperature equations have been developed for planktic (e.g., Bemis and others, 1998) and benthic (e.g., Shackleton and Kennett, 1975) foraminifera, they all are based on the original temperature equation proposed by Epstein and others (1953):

$$T = 16.5 - 4.3 \times (\delta^{18}\text{O}_{\text{calcite}} - \delta_w) + 0.14 \times (\delta^{18}\text{O}_{\text{calcite}} - \delta_w)^2$$

where T is the temperature ($^\circ\text{C}$), δ_w is the oxygen isotope value of the water in which the organism lived, and $\delta^{18}\text{O}_{\text{calcite}}$ is the oxygen isotope value of foraminiferal calcite

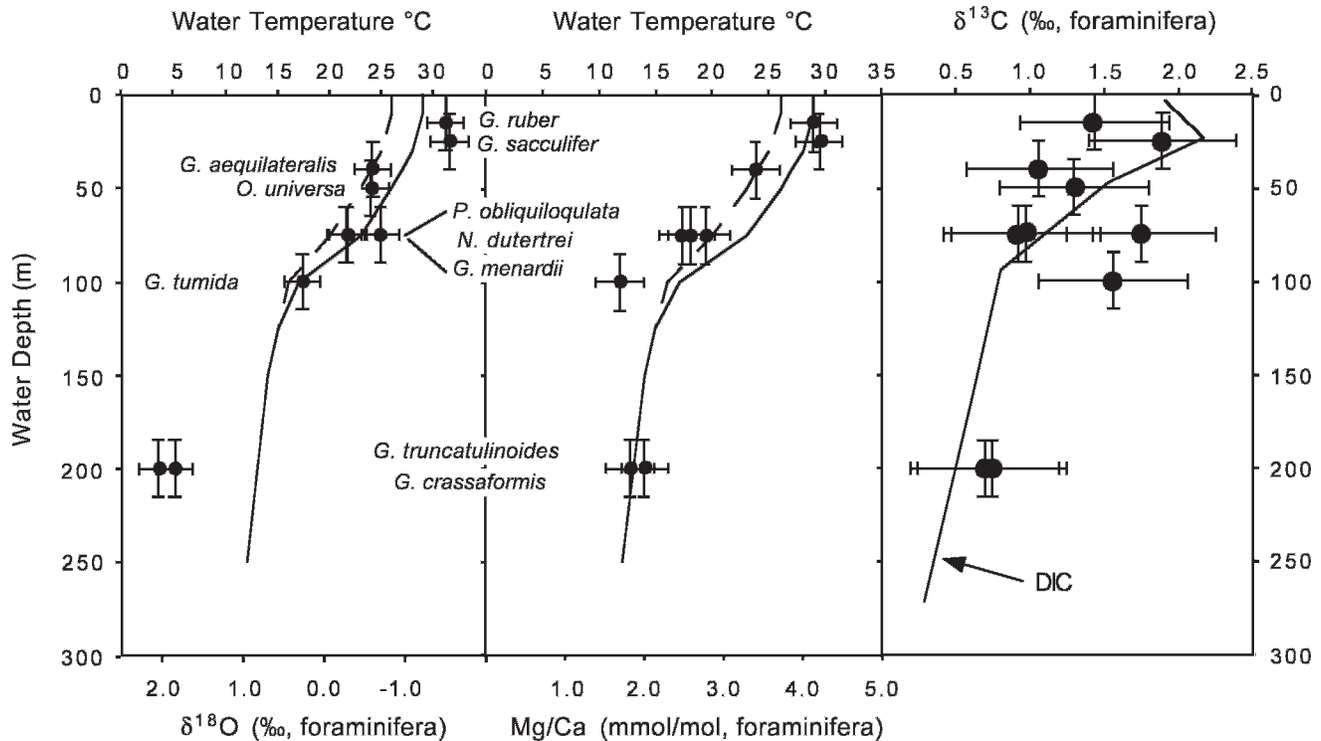


FIGURE 2. Planktic foraminiferal geochemistry reflects depth habitat, varying with changes in seawater properties. Planktic foraminiferal data are from core-top samples, Sierra Leone Rise EN66 38GGC, including $\delta^{18}\text{O}$ and Mg/Ca (Rosenthal and Boyle, 1993) and $\delta^{13}\text{C}$ (Rosenthal, unpublished data). Seawater seasonal temperature range (indicated by solid and dashed lines) and $\delta^{13}\text{C}$ of DIC are from nearby GEOSECS station 113. Error bars indicate estimated range in calcification depth and water properties. Species names are displayed at approximate calcification depth.

measured in a mass spectrometer. A change in the δ_w term will cause an equal change in the measured $\delta^{18}\text{O}_{\text{calcite}}$ value, whereas an inverse relationship exists between $\delta^{18}\text{O}_{\text{calcite}}$ and temperature such that for every 1°C increase, there is a $\sim 0.23\text{‰}$ decrease in the measured $\delta^{18}\text{O}_{\text{calcite}}$ value. These relationships allow us to interpret the $\delta^{18}\text{O}$ values in calcite-secreting organisms, including foraminifera.

Variations in the amount of water stored on land through time, usually in the form of ice, can have a significant effect on the mean ocean δ_w value, and hence, on the marine $\delta^{18}\text{O}_{\text{calcite}}$ record. During interglacial periods, most high-latitude precipitation returns to the oceans via summer ice and snow melt. During glacials, a larger fraction of ^{16}O -enriched snow and ice persists through cooler summers and accumulates into large continental ice sheets. The difference between ice sheet (-35 to -40‰) and mean ocean ($\sim 0\text{‰}$) $\delta^{18}\text{O}$ values is large, so fluctuations in ice-sheet size alter mean oceanic δ_w values.

For example, water stored in continental ice sheets during the last glacial maximum lowered global sea level by $\sim 120 \pm 5$ m, removed $\sim 3\%$ of the ocean's water volume, and resulted in a mean ocean δ_w value that was 1.2‰ higher than it is today. Therefore, there is a relationship between δ_w and sea level, with an empirical calibration of $0.10\text{--}0.12\text{‰}/10$ m, likely varying through time because of varying ice sheet $\delta^{18}\text{O}$ (Fairbanks and Matthews, 1978; Pekar and others, 2002; Miller and others, 2005; Katz and others, 2008; Miller and others, 2008).

Related to this, benthic foraminiferal $\delta^{18}\text{O}$ records can be used to reconstruct global ice-volume changes and past ocean temperature at the seafloor. Seafloor temperature reconstructions for times when large continental ice sheets existed must separate the temperature signal from the δ_w signal recorded in foraminiferal $\delta^{18}\text{O}$. The current approach to unravelling these two signals is to measure $\delta^{18}\text{O}$ and Mg/Ca in the same samples (see "Foraminiferal Mg/Ca paleothermometry" below). Glacial-interglacial deep-sea temperature changes are relatively small compared to SST changes at most latitudes; therefore, the amplitude in glacial-interglacial planktic foraminiferal $\delta^{18}\text{O}$ may be larger than the benthic foraminiferal $\delta^{18}\text{O}$ signal.

In addition to reconstructing past ocean temperature at the seafloor and global ice-volume changes, benthic foraminiferal $\delta^{18}\text{O}$ records can be used to reconstruct past deepwater source regions. The deep-ocean $\delta^{18}\text{O}$ of benthic foraminifera approximates the high-latitude surface-water conditions where the major deepwater masses originate. Interbasinal gradients in benthic foraminiferal $\delta^{18}\text{O}$ values (Fig. 1) can be a useful indicator of differentiation between deepwater source regions, and therefore of large-scale changes in the mode of ocean overturning (Cramer and others, 2009). Comparison of low-latitude planktic with high-latitude benthic foraminiferal $\delta^{18}\text{O}$ records provides the means to assess temperature gradients, both equator-to-pole and surface-to-deep (Fig. 1), providing the means to reconstruct the planetary temperature gradients and poleward heat transport that are a primary driver of climate

change (see “Foraminiferal Mg/Ca paleothermometry” below).

Planktic foraminiferal $\delta^{18}\text{O}$ can be a good proxy for SSTs if several issues are taken into account. Different species from the same sediment sample may show $\delta^{18}\text{O}$ offsets of as much as 1.5‰. This offset is attributed in part to species-specific vital effects (i.e., species-specific metabolic variation in isotopic fractionation), but some offset results from differences in seasonal temperatures or depth habitats of different species (Fig. 2), which may vary from region to region (e.g., Bè and others, 1966; Fairbanks and Wiebe, 1980; Fairbanks and others, 1980; Fairbanks and others, 1982; Curry and Crowley, 1987; Ravelo and Fairbanks, 1992; Spero and others, 2003). During the planktic foraminiferal life cycle, vertical migration through the water column with a surface-to-deep temperature gradient may result in $\delta^{18}\text{O}$ differences between inner and outer test chambers, depending on the depth of calcification. Because differences in seasonal surface-water temperatures or different depth habitats of various species can affect planktic foraminiferal $\delta^{18}\text{O}$, multi-species analyses may allow reconstructions of temperature gradients within the upper part of the surface ocean (e.g., Bè and others, 1966; Fairbanks and others, 1980; Fairbanks and others, 1982; Ravelo and Fairbanks, 1992; Spero and others, 2003).

Changes during the life cycle of some planktic foraminiferal species may affect specimens' $\delta^{18}\text{O}$ values. Some planktic foraminifera secrete a calcite layer towards the end of the life cycle, prior to gamete release (e.g., Bè and others, 1966; Fairbanks and others, 1980). As part of the gametogenesis cycle, many of the surface-dwelling planktic foraminifera will shed their spines and sink through the water column. Even while they are reproducing, some shell mass is added at depth; this can change the overall shell chemistry that was originally acquired in shallower water. This is a concern when making measurements in the tropics, where SSTs can be 10 to 15°C warmer than in the thermocline waters through which the foraminifera sink.

In the mid- to high-latitudes, the largest temperature variations occur at the surface and produce a seasonal succession of planktic foraminifera as SSTs warm and cool (e.g., Reynolds and Thunell, 1985, 1986; Deuser, 1987; Sautter and Thunell, 1989). Subsurface vertical temperature changes are small. During winter, nearly isothermal conditions extend downward from the surface through the depth range of the planktic foraminiferal habitat. Hence, the largest temperature variations are seasonal and are usually restricted to the upper 50 m. Phytoplankton, and hence zooplankton, bloom when light and nutrients exceed minimum threshold levels in the surface water. This event is diachronous regionally because it occurs progressively later at higher latitudes. Sediment traps provide a record of the planktic foraminiferal flux throughout the year (e.g., Reynolds and Thunell, 1985). In mid- to high-latitude regions, the flux of most planktic foraminifera to the seafloor is weighted towards a very short annual interval, and therefore may provide only seasonal information (e.g., Reynolds and Thunell, 1985, 1986; Donner and Wefer, 1994).

The $\delta^{18}\text{O}$ offsets among planktic foraminiferal species can provide the means to reconstruct surface-water structure

and seasonal temperature changes (Fig. 2). However, the offsets also may introduce error into paleoceanographic reconstructions through time. Nonetheless, careful correction for interspecies isotopic differences provides the opportunity to reconstruct water-column structure and paleoceanographic changes.

Diagenesis and $\delta^{18}\text{O}$

Post-depositional diagenesis can affect foraminiferal $\delta^{18}\text{O}$ values. Deep-burial diagenesis (>~500 m) tends to decrease $\delta^{18}\text{O}$ values because fossil foraminiferal tests may recalcify at equilibrium with sediment temperatures that are higher than at the seafloor, due to the geothermal gradient. Recent work indicates that shallow-burial diagenesis might alter planktic foraminiferal $\delta^{18}\text{O}$ values by overprinting a cool deepwater signal on warm surface-water planktic foraminiferal calcite (Pearson and others, 2001). This diagenesis may occur when calcium carbonate precipitates on the foraminiferal tests in the sediments in cold bottom waters. Benthic foraminifera would not be adversely affected by this process because they calcify in the same cold bottom waters in which they are buried. Perhaps the most effective strategy is to avoid diagenetic and recrystallization complications by analyzing foraminifera preserved in clay-rich sediments that are better insulated from potential pore-water diagenesis, making them more suitable for geochemical analyses (Pearson and others, 2001). Foraminifera preserved in clay-rich sediments appear to retain their original structures and geochemical signatures (Pearson and others, 2001; Sexton and others, 2006).

CARBON ISOTOPES ($\delta^{13}\text{C}$)

The analytical approach to measuring $\delta^{18}\text{O}$ results in the synchronous measurement of $\delta^{13}\text{C}$, the deviation from a standard of the carbon isotopic ratio ($^{13}\text{C}/^{12}\text{C}$), reported in the same “ δ ” notation as oxygen isotopes:

$$\delta^{13}\text{C}_{\text{sample}} = 1000 \times \left[\left(\frac{^{13}\text{C}/^{12}\text{C}_{\text{sample}}}{^{13}\text{C}/^{12}\text{C}_{\text{standard}}} \right) - 1 \right] \text{‰}$$

Whereas the ambient temperature and the $\delta^{18}\text{O}$ value of seawater (δ_w) determine the $\delta^{18}\text{O}$ value recorded in foraminiferal tests, the $\delta^{13}\text{C}$ value is primarily a function of the dissolved inorganic carbon (DIC) $\delta^{13}\text{C}$ value in seawater (e.g., Urey, 1947; Epstein and others, 1953; Emiliani, 1955; O'Neil and others, 1969; Graham and others, 1981; McCorkle and others, 1990).

The average $\delta^{13}\text{C}$ signature of the ocean's DIC reservoir on timescales longer than the residence time of carbon in the oceans (~180 kyr) is controlled by the $\delta^{13}\text{C}$ signatures and fluxes of carbon sources and sinks. Carbon is supplied to the ocean from hydrothermal/volcanic outgassing and from chemical weathering of continental rocks, while carbon is removed from the ocean through deposition of carbonate and organic carbon in marine sediments. Over millions of years, these sediments integrate large kinetic fractionations from the photosynthetic reduction of CO_2 to organic matter with small thermodynamic fractionations from ion exchange reactions during carbonate precipita-

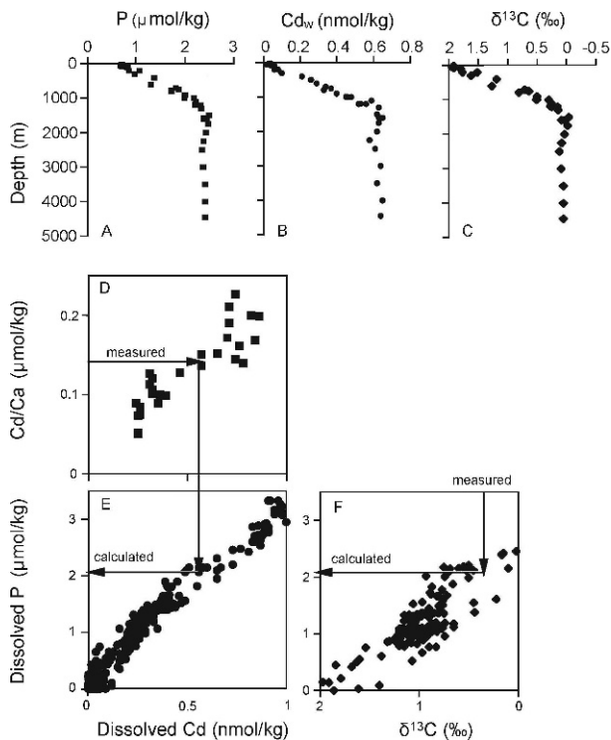


FIGURE 3. Vertical profiles of: A) phosphate; B) dissolved cadmium (Cd_w); and C) carbon isotopes ($\delta^{13}\text{C}$) of DIC from the Southern Ocean depicting the strong similarity among the major nutrient and the nutrient-like properties. Using measurements of Cd/Ca and $\delta^{13}\text{C}$ in benthic foraminifera and the modern relationships between these properties allows us to reconstruct both dissolved P concentration in the past and changes in deep ocean circulation. D) Core-top calibration between benthic foraminiferal Cd/Ca and bottom-water concentrations of dissolved Cd for deep cores. For intermediate depths, the partition coefficient varies with water depth, linearly increasing from 1.3 at depths ≤ 1.3 km to a value of 2.9 at depths > 3 km (Boyle, 1992). E) The modern $\text{Cd} = \text{P}$ relationships in deep waters is described using two equations: for $\text{P} < 1.3$, $\text{Cd}_w = 0.21 * \text{P}$; and for $\text{P} > 1.3$, $\text{Cd}_w = 0.40(\text{P} - 1.3) + 0.279$, where P is in $\mu\text{mol/kg}$ and Cd_w is in nmol/kg (Boyle, 1988). F) The modern $\delta^{13}\text{C}$ -P relationship in deep waters is described using the equation: $\delta^{13}\text{C} = 2.7 - 1.1 * \text{P}$.

tion. Changes in carbon isotopes of marine carbonates and organic matter through time serve as archives of changes in carbon sources and sinks. For detailed summaries of the geological carbon cycle, see Hayes and others (1999), Kump and Arthur (1999), Berner (2004), and Katz and others (2005).

Deepwater $\delta^{13}\text{C}$

On shorter time scales, there are considerable differences in $\delta^{13}\text{C}_{\text{DIC}}$ values throughout the oceans. For example, modern deep ocean values vary from $\sim 1.0\text{‰}$ in the North Atlantic to -0.5‰ in the North Pacific (Fig. 1). These $\delta^{13}\text{C}$ differences reflect (1) the basin-to-basin fractionation caused by different source regions and deepwater circulation patterns, and (2) surface-ocean productivity and related ^{12}C -enriched carbon export from the surface to the deep ocean (Kroopnick, 1985) (Fig. 3). In the modern ocean, deepwater source regions in the North Atlantic and

Southern Ocean are differentiated by temperature, salinity, and nutrient content. Interbasinal $\delta^{13}\text{C}$ fractionation occurs primarily because of differences in initial nutrient and $\delta^{13}\text{C}$ content of deepwater formed in different locations, and secondarily because the $\delta^{13}\text{C}$ value and nutrient content of deep/bottom waters changes as a function of the amount of time the water mass has been isolated from the surface. For example, whereas the Atlantic vs. Pacific $\delta^{13}\text{C}$ contrast in the modern ocean reflects aging, the $\delta^{13}\text{C}$ contrast in the deep Atlantic behaves conservatively and represents a mixing between North Atlantic deepwater and Antarctic bottom water end-member components. This is shown by the fact that the $\delta^{13}\text{C}$ distribution of the deep Atlantic mimics salinity (Wright and others, 1991).

The distinct $\delta^{13}\text{C}$ characteristics of different deepwater masses have been exploited to reconstruct deepwater circulation patterns (Figs. 1, 3). The deepwater $\delta^{13}\text{C}_{\text{DIC}}$ is recorded in the CaCO_3 tests of benthic foraminifera, providing a means to reconstruct DIC through time and space. Thus, for times when the nutrient/ $\delta^{13}\text{C}$ characteristics of each deepwater source region are very different (e.g., Neogene), carbon isotope records from deep-ocean benthic foraminifera are among the best tools available for inferring past deepwater circulation patterns. These reconstructions are done by comparing benthic foraminiferal $\delta^{13}\text{C}$ values at various locations in the deep ocean and assuming that the differences reflect mixing between end-member deepwater source regions (e.g., Belanger and others, 1981; Graham and others, 1981; Curry and Lohmann, 1982; Oppo and Fairbanks, 1987; Wright and others, 1991; Wright and Miller, 1996; Poore and others, 2006; Cramer and others, 2009).

As water masses sink and move away from their source regions, ^{12}C -enriched organic matter sinking from the surface ocean collects at depth, either in the water column or on the seafloor. Oxidation of this organic matter releases nutrients and CO_2 with low $\delta^{13}\text{C}$ values (approximately -25‰ today), lowering the $\delta^{13}\text{C}_{\text{DIC}}$ value and increasing nutrient levels (Fig. 3) as the deepwater mass "ages" (i.e., flows farther from its source region). Because of these processes, a deepwater mass proximal to its source region is likely to have relatively high $\delta^{13}\text{C}_{\text{DIC}}$ values and low nutrient levels; in contrast, the more distal (older) deepwater mass is likely to have lower $\delta^{13}\text{C}_{\text{DIC}}$ values and higher nutrient levels than near the source region (Fig. 1).

Surface Ocean $\delta^{13}\text{C}$

The $\delta^{13}\text{C}$ recorded in planktic foraminiferal tests primarily reflects $\delta^{13}\text{C}_{\text{DIC}}$ in the upper few hundred meters of the surface ocean (Fig. 2), along with the influence of foraminiferal respiration and symbiotic photosynthesis (e.g., Spero and Lea, 1993; Spero and Lea, 1996; Bemis and others, 1998) (Fig. 2). The abundance of phytoplankton, both in surface waters and as symbionts in some species of planktic foraminifera, can alter the $\delta^{13}\text{C}_{\text{DIC}}$ on a scale that ranges from regional to the microhabitat of an individual. Additional factors may influence planktic foraminiferal $\delta^{13}\text{C}$, such as test size, pH, and light intensity (e.g., Oppo and Fairbanks, 1989; Spero and others, 1991).

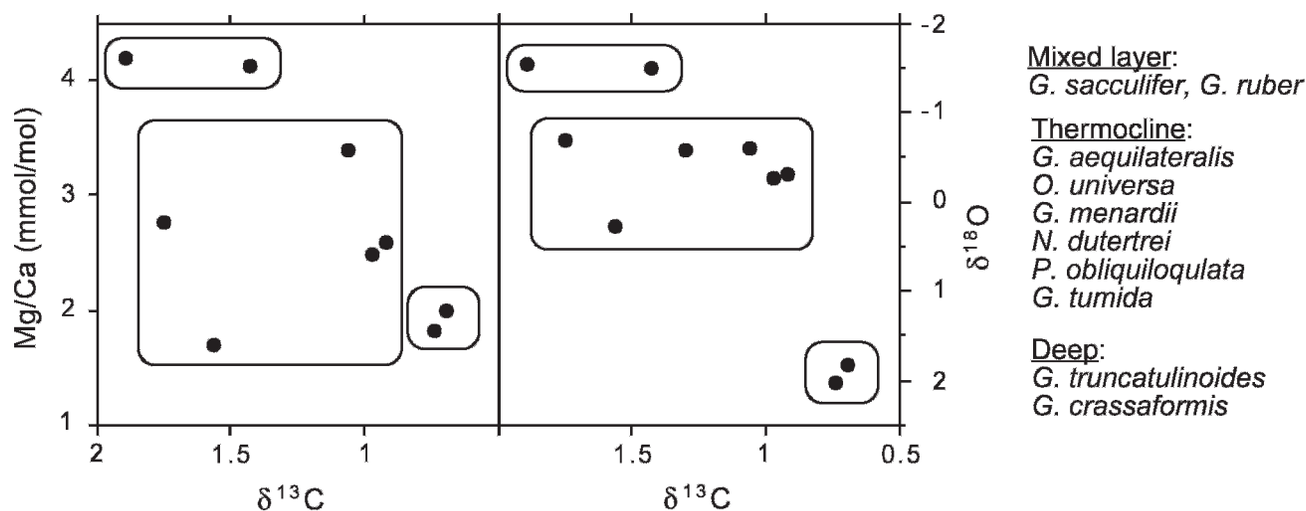


FIGURE 4. Comparison of $\delta^{18}\text{O}$ and Mg/Ca (Rosenthal and Boyle, 1993) with $\delta^{13}\text{C}$ (Rosenthal, unpublished data) of planktic foraminifera from core-top samples (Sierra Leone Rise EN66 38GGC) illustrates the correspondence between depth habitat and geochemistry of foraminiferal tests. Species are grouped into mixed layer, thermocline, and deep dwellers.

Phytoplankton preferentially take up ^{12}C during photosynthesis, leaving the surrounding waters enriched in ^{13}C and yielding higher $\delta^{13}\text{C}_{\text{DIC}}$. In areas of high primary and export production, phytoplankton can deplete the shallowest surface waters of ^{12}C , and enrich the deeper surface waters with ^{12}C . As a result, planktic foraminiferal $\delta^{13}\text{C}$ decreases with depth from the sea surface to the nutricline (Figs. 2, 3) (e.g., Fairbanks and others, 1980; Curry and Crowley, 1987).

Similarly, photosynthetic symbionts enrich the micro-environment of the host foraminifera in ^{13}C , resulting in $\delta^{13}\text{C}_{\text{calcite}}$ that is higher than surrounding seawater (e.g., Oppo and Fairbanks, 1989; Spero and Lea, 1993). At the same time, foraminiferal respiration reintroduces ^{12}C -enriched CO_2 back into the DIC pool (Kroopnick, 1974). Studies have shown that there is a positive relationship between test size of symbiotic-bearing planktic foraminifera and $\delta^{13}\text{C}$ (e.g., Berger, 1978; Curry and Matthews, 1981; Fairbanks and others, 1982; Oppo and Fairbanks, 1989; Spero and Lea, 1993; Billups and Spero, 1995; Faul and others, 2000).

The trend towards lower seawater temperature and ^{12}C -enrichment in DIC with increasing depth in the surface ocean produces a trend towards increasing $\delta^{18}\text{O}$ and decreasing $\delta^{13}\text{C}$ in planktic foraminifera from the shallow mixed layer down to the base of the thermocline (Fig. 2) (e.g., Fairbanks and others, 1980; Curry and Crowley, 1987; Ravelo and Fairbanks, 1992; Faul and others, 2000). This relationship tends to produce a correlation between $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$ in planktic foraminifera that has been documented in plankton tows (Fig. 4), and that has been utilized in fossil assemblages to reconstruct changes in the surface ocean structure and productivity (e.g., Faul and others, 2000). It has also provided the basis for reconstructions of surface ocean hydrography and productivity using extinct planktic foraminiferal species, under the assumption that the general $\delta^{18}\text{O}$ - $\delta^{13}\text{C}$ relationship is valid through time (e.g., Pearson and others, 2001).

Paleoproductivity: Integrated Geochemical ($\delta^{13}\text{C}$) and Paleontological Proxies

Numerous studies document that organic matter sinking from the surface ocean to the seafloor is the primary food supply for benthic foraminifera, and hence that surface-water productivity (= food supply) is a primary control on the species composition, accumulation rate, and geographic distribution of benthic foraminiferal faunas (e.g., Miller and Lohmann, 1982; Thomas and Vincent, 1987; Loubere, 1994; Rathburn and Corliss, 1994; Jorissen and others, 1995; Thomas and Gooday, 1996; Schmiedl and Mackensen, 1997; Schmiedl and others, 2000). This relationship between surface and deep water organisms is recorded in several types of geochemical and paleontological data, providing the potential to use a multi-proxy approach to constrain and monitor productivity changes through time.

A classic method used to reconstruct surface productivity and export production is to compare planktic foraminiferal $\delta^{13}\text{C}$ from phosphate-free surface waters with benthic foraminiferal $\delta^{13}\text{C}$ (mean deepwater proxy). As discussed above, ^{12}C -enriched organic matter sinks from the surface ocean to the seafloor, driving $\delta^{13}\text{C}$ of DIC higher at the surface and lower at the seafloor. Therefore, high export production drives $\delta^{13}\text{C}$ higher in planktics and lower in benthics, resulting in an offset between the two, referred to as $\Delta\delta^{13}\text{C}$. Whereas a paleoproductivity signal can be extracted from $\Delta\delta^{13}\text{C}$, it is actually a proxy for the carbon to phosphorous ratio (C/P) (Broecker and Peng, 1982), which is proportional to productivity only as a function of stability/mixing.

Another means of estimating paleoproductivity is to compare $\delta^{13}\text{C}$ of epifaunal vs. infaunal benthic foraminifera in the same sample (within the constraints of the amount of time represented in a sample by sedimentation rate). Decay of organic matter within sediments enriches porewaters in ^{12}C , which can be incorporated into the tests of infaunal benthic foraminifera as they calcify. Therefore, an offset between coeval epifaunal and infaunal $\delta^{13}\text{C}$ may reflect

greater organic matter preservation within sediments than at the sediment surface, where organic matter may be oxidized before it is consumed (Mackensen and others, 1993b). When $\Delta\delta^{13}\text{C}$ between epifaunal and infaunal species approaches 0‰ and the $\delta^{13}\text{C}$ values of both are low, there was abundant food supply at the surface and within sediments (i.e., surface-water productivity was high). The higher the $\delta^{13}\text{C}$, the lower the food supply, and hence, paleoproductivity.

Support for conclusions drawn from geochemical paleoproductivity proxies can be found in paleontological proxies for food availability and by extension, surface-water productivity. The paleontological proxies must be studied using specimens from the same samples that were used to produce the geochemical records. For instance, a decrease in planktic foraminiferal flux and concomitant increase in benthic foraminiferal flux may indicate higher surface ocean primary productivity, under the assumption that a higher food supply will support more benthic foraminifera. Similarly, increases in benthic foraminiferal accumulation rates (as opposed to relative abundances) may indicate that food supply increased, thereby sustaining larger populations (e.g., Herguera, 1992). Epifaunal vs. infaunal benthic abundances are also used as a paleoproductivity indicator because epifaunal taxa tend to dominate assemblages when oligotrophic conditions prevail, whereas infaunal taxa dominate as food supply increases (e.g., Rathburn and Corliss, 1994; Jorissen and others, 1995; van der Zwaan and others, 1999). There are several trochospiral epifaunal benthic foraminiferal species that flourish in seasonal phytodetritus (e.g., Gooday, 1988; Smart and others, 1994; Loubere and Fariddudin, 1999) that are used as indicators of high export production. Thomas and Gooday (1996) identified the “phytodetritus species” *Epistominella exigua* (Brady) and *Alabaminella weddellensis* (Earland) in the Paleogene, and noted that species richness declined at southern high latitudes, but not in the Pacific tropics. This led them to speculate that expansion of these taxa in the Oligocene was the result of increased seasonal food supply at high latitudes. Finally, comprehensive benthic assemblage data have been used to estimate paleoproductivity (e.g., Loubere and Fariddudin, 1999).

BENTHIC FORAMINIFERAL INTERSPECIES $\delta^{18}\text{O}$ AND $\delta^{13}\text{C}$ OFFSETS

Studies that utilize geochemical analyses of foraminifera rely on the assumption that variations in foraminiferal test chemistry primarily reflect environmental changes through time or contemporaneous environmental differences related to geography or bathymetry. A potential complicating factor is that measurements of different foraminiferal species in the same sample are often consistently offset from each other and from calcite precipitated in equilibrium with the surrounding water, both in benthic (e.g., Duplessy and others, 1970; Shackleton, 1974; Belanger and others, 1981; Graham and others, 1981; McCorkle and others, 1990; Katz and others, 2003a) and planktic foraminifera (e.g., Bè and others, 1966; Fairbanks and others, 1980; Fairbanks and others, 1982; Ravelo and Fairbanks, 1992;

Spero and others, 2003). Planktic foraminiferal species isotopic offsets are discussed in the preceding sections.

Benthic foraminiferal interspecies isotopic offsets have been attributed to microhabitat differences, in which tests grow in geochemically distinct environments (e.g., Belanger and others, 1981; Corliss, 1985; McCorkle and others, 1990), presence of symbionts (e.g., Spero and Lea, 1993; Saraswati and others, 2004), and vital effects (e.g., Duplessy and others, 1970; McCorkle and others, 1990). Benthic foraminifera have different microhabitat preferences, ranging from deep infaunal to epifaunal to elevated substrates (e.g., Corliss, 1985; Gooday, 1986; Bernhard, 1989) (see Introduction).

Porewater chemistry differs from overlying seawater chemistry because organic matter decomposition within the sediments alters the porewater chemistry (e.g., Bender and others, 1977; Froelich and others, 1979; Emerson and others, 1980). In addition, porewaters may migrate within sediments. Carbon isotopes may be especially sensitive because the oxidation of organic matter within sediments lowers the $\delta^{13}\text{C}_{\text{DIC}}$ value of the porewater; therefore, porewater $\delta^{13}\text{C}_{\text{DIC}}$ profiles tend to decrease with increasing sub-seafloor depth. Isotopic analyses of modern benthic foraminifera show that epifaunal forms tend to approximate seafloor seawater values, whereas infaunal taxa are influenced by porewater values (e.g., McCorkle and others, 1990). Microhabitat effects may play a larger role than seawater $\delta^{13}\text{C}$ in some settings, as concluded from a study of living benthic foraminifera in the Bay of Biscay (Fontanier and others, 2006).

Culture studies of $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$ in *Bulimina aculeata* d'Orbigny and *Rosalina vilardeboana* d'Orbigny yielded similar results to field studies, even though maintaining culture system stability can be challenging (McCorkle and others, 2008). Systematic offsets in isotopic values of these two species, grown in the same conditions, supports species-specific, non-environmental controls on benthic foraminiferal test chemistry. The authors also found that very small specimens tend to be depleted in ^{18}O and ^{13}C , although these ontogenetic effects are not likely to skew paleoceanographic studies that typically use substantially larger specimens (McCorkle and others, 2008). We note that the two species used in these culture studies are typically not used in paleoceanographic studies.

To avoid potential bias from vital effect and microhabitat differences, the best strategy for reconstructing past oceanographic changes is to use monospecific (specimens of the same species) or monogeneric (specimens of multiple species in the same genus) records. Many benthic foraminiferal taxa are long-lived, and therefore one species/genus may be used to construct records spanning millions to tens of millions of years. In contrast, planktic taxa are shorter-lived in general (a few million years), necessitating spliced records from several species, corrected for interspecies offsets, for long-term planktic isotopic records. It is interesting to note that the first $\delta^{18}\text{O}$ Cenozoic syntheses were based on mixed species analyses (Savin and others, 1975; Shackleton and Kennett, 1975), yet the basic features captured in these older curves are clear in more recent compilations (Miller and others, 1987; Zachos and others, 2001b; Cramer and others, 2009). This attests to the

robustness of these records in spite of offsets due to vital effects.

EFFECTS OF pH AND CARBONATE ION CONCENTRATION ([CO₃²⁻]) ON δ¹⁸O AND δ¹³C

Culture studies with several symbiotic and non-symbiotic planktic foraminiferal species have shown that δ¹⁸O and δ¹³C of the foraminiferal calcite decreases with increasing pH (Spero and others, 1997; Spero and others, 1999). This finding has been explained by the fact that as pH increases, the dominant form of DIC shifts from HCO₃⁻ to CO₃²⁻, and because the fractionation of oxygen isotopes with respect to H₂O is greater (i.e., higher δ¹⁸O) for HCO₃⁻ than for CO₃²⁻, the isotopic composition of calcite decreases as CO₃²⁻ is increasingly used for calcification (Zeebe and others, 1999). This interpretation is supported by theoretical study of inorganically precipitated calcite (Zeebe and others, 1999), and a culture experiment with constant pH and variable [CO₃²⁻], which indicated that it is the CO₃²⁻:HCO₃⁻ ratio, rather than the [CO₃²⁻], that affects δ¹⁸O of *Orbulina universa* d'Orbigny (Bijma and others, 1999). In contrast, the carbonate ion effect on δ¹³C is mechanistically much less well understood (Zeebe and others, 1999), although the different response (i.e., δ¹³C/CO₃²⁻ slope) of various foraminiferal species has been used to quantify past changes in the carbonate ion (Spero and others, 1997).

The impact of changes in pH through time could affect interpretations of glacial-interglacial changes and longer-term trends in isotopic records. Correcting for this effect would decrease glacial tropical SST estimates by as much as 1°C, and attribute a portion of the glacial δ¹³C decrease to increasing [CO₃²⁻] (Spero and others, 1997). In the older record, Cretaceous SSTs may have been 2–3.5°C warmer than initially estimated (Zeebe and Wolf-Gladrow, 2001), although the difference may be at least partially offset by lower Cretaceous δ_w due to hydrothermal cycling of water through oceanic crust (Wallmann, 2004).

CHEMOSTRATIGRAPHY

Geochemical proxies can be used for stratigraphic correlation in one of two ways. First, for proxies (e.g., Sr isotopes) with residence times in the ocean much longer than the ocean mixing time, and substantial secular variability in the measured values, the proxy data can be converted to an age using a look-up table of age-calibrated values (Howarth and McArthur, 1997). Second, rapid changes in a geochemical proxy (typically δ¹⁸O or δ¹³C) can be correlated between geographically distant sites, based on the assumption that mixing of the ocean and atmosphere requires that such changes are synchronous on 10³-yr timescales. Chronostratigraphic use of geochemical proxies includes radiometric dating (¹⁴C) and identification of predictable orbitally forced cyclicity that can be directly related to temporal duration. Milankovitch orbital cycles result from gravitational interactions among Earth and the other planets, Sun, and Moon that alter Earth's orbit at regular cycles (averaging ~23, 41, 100, and 400 kyr, and 2.5 Myr). Milankovitch-scale climate variability results from amplification of orbitally driven variations in the geographic and intra-annual distribution of

solar energy on Earth, which in turn affects a range of sedimentary and geochemical parameters, including the chemistry of foraminiferal tests.

STABLE OXYGEN AND CARBON ISOTOPES

The value of δ¹⁸O in foraminifera for stratigraphic correlation was recognized even from the earliest measurements (Emiliani and Shackleton, 1974). Reconstructing Pleistocene glacial cycles was an early impetus for down-core stable isotopic measurements, and the implicit strategy of correlating glacial and interglacial maxima between widely separated sites was exploited as soon as a sufficient number of downcore records was available (Emiliani and Shackleton, 1974; Imbrie and others, 1984). This has been refined into a series of over 200 marine isotope stages, numbered within successive magnetic polarity chrons, extending into the late Miocene (~5.3 Ma) (Lisiecki and Raymo, 2005). Correlation to this series of glacials and interglacials can be used to refine an age model derived using other methods (e.g., biostratigraphy, magnetostratigraphy), although the cyclic nature of the glacial-interglacial signal prevents using it as a primary stratigraphic tool. Recognition of the glacial imprint on older δ¹⁸O records has led to the designation of a series of Myr-scale Oligocene and Miocene isotope events that are Myr composites of higher-frequency Milankovitch changes (Oi-1 through Oi-2b and Mi-1 through Mi-7 series (Miller and others, 1991b; Pekar and Miller, 1996).

Over longer time spans, variability in the marine δ¹³C record has proven useful as a stratigraphic correlation tool. With a few exceptions (Monterey hypothesis, Vincent and Berger, 1985), the causal mechanisms for δ¹³C changes are not as clearly defined as for δ¹⁸O. Nonetheless, δ¹³C stratigraphy has been critical in constructing robust core-to-core correlations to allow reconstruction of variations in deepwater circulation (e.g., Wright and Miller, 1996) and to test the synchrony of biostratigraphic datums (Cramer and others, 2003).

In conjunction with the Milankovitch theory of glacial-interglacial cycles forced by cyclic variations in Earth's orbital geometry, the recognition of glacial-interglacial cyclicity led to early use of δ¹⁸O cyclicity as a chronostratigraphic tool. The tuning of δ¹⁸O records to an ice-volume model driven by summer insolation at 65°N, which varies with the orbital geometry, was the foundation of the SPECMAP timescale (Imbrie and others, 1984; Martinson and others, 1987) and culminated in the recalibration of the age of Brunhes/Matuyama magnetic polarity reversal (Shackleton and others, 1990). This strategy of tuning the δ¹⁸O record is still used (e.g., Lisiecki and Raymo, 2005), although it may create the impression that ice volume is more linearly related to insolation than is actually the case (Huybers and Wunsch, 2004). The orbital theory for climate change received a boost from the correlation of the ~100 kyr variability with periodic variability in the eccentricity of Earth's orbit around the Sun (Hays and others, 1976). However, the δ¹⁸O spectrum in the 10 cycles/Myr range has never fit the spectrum of eccentricity very well (e.g., Kominz and Pisias, 1979), precluding a simple

model for orbital forcing of the Pleistocene glacial-interglacial cycles (e.g., Imbrie and others, 1993). It has been suggested that the large Pleistocene ~ 100 -kyr cycles in $\delta^{18}\text{O}$ result from temperature changes due to atmospheric CO_2 responding to oceanographic changes in the Southern Ocean, and largely independent of Northern Hemisphere ice sheets (Shackleton, 2000; Toggweiler, 2008). Most orbital stratigraphies in the pre-Pliocene instead rely on tuning of sedimentary parameters (e.g., % CaCO_3 , magnetic susceptibility), with the resulting chronology used to test for orbital control on geochemical proxies (e.g., Zachos and others, 2001a; Pälike and others, 2006).

The presence of orbital cyclicity in $\delta^{13}\text{C}$ records has frequently been noted, but typically not emphasized (e.g., see Zachos and others, 2001a). In the pre-Pleistocene, the eccentricity (~ 100 kyr, ~ 400 kyr, and ~ 2.5 Myr) cyclicity is especially pronounced (Woodruff and Savin, 1991; Zachos and others, 2001a; Pälike and others, 2006). Cramer and others (2003) explained this as due to the low-pass filter effect of the carbon residence time (~ 100 kyr), and proposed that the eccentricity cyclicity should be used as a primary tool in constructing orbital stratigraphies especially for intervals older than 50 Ma when the uncertainty in the orbital solution is too large to be of use as a tuning target curve.

The investigation of the orbital imprint on foraminiferal geochemical proxies is based on $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$ records, primarily because of the relative simplicity of these measurements. At present, measurement of trace element ratios requires a large sample size that makes high-resolution studies difficult. Further investigation of orbital cyclicity in other geochemical proxies should be of interest, both in constraining the climate response to orbital forcing and in establishing the robustness of the proxies.

STRONTIUM ISOTOPES

Two isotopes of strontium (^{87}Sr , ^{86}Sr) are introduced into the oceans through: 1) hydrothermal exchange, primarily at mid-ocean ridges (low $^{87}\text{Sr}/^{86}\text{Sr}$ values, ~ 0.703); and 2) continental erosion (high $^{87}\text{Sr}/^{86}\text{Sr}$ values, ~ 0.718). However, the average $^{87}\text{Sr}/^{86}\text{Sr}$ in seawater may be complicated by changes in dominant continental source rock type (e.g., Ravizza, 1993) and variable riverine fluxes (e.g., Lear and others, 2003). The $^{87}\text{Sr}/^{86}\text{Sr}$ ratio in seawater is essentially uniform at any given time because the residence time (2–4 Myr) is much longer than oceanic mixing times (Broecker and Peng, 1982). Note the much smaller short-term variability relative to long-term variability in $^{87}\text{Sr}/^{86}\text{Sr}$, in contrast with large short-term variability in $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$ (Fig. 1).

Strontium can substitute for calcium during calcification of foraminiferal tests. The $^{87}\text{Sr}/^{86}\text{Sr}$ ratio of the seawater is preserved in foraminiferal tests, provided diagenetic alteration does not occur (Elderfield, 1986). Strontium isotope ratios provide an important correlation tool in stratigraphic studies of marine sediments with a precision as good as ± 0.3 Myr during times of high rates of change or inflection points in the $^{87}\text{Sr}/^{86}\text{Sr}$ record. For example, $^{87}\text{Sr}/^{86}\text{Sr}$ can provide good stratigraphic control for much of the late Eocene to Recent because of changing $^{87}\text{Sr}/^{86}\text{Sr}$ values (Fig. 1; Burke, 1982; DePaolo, 1985; Hess, 1986).

Sr-isotope stratigraphy using foraminifera has the potential to significantly enhance stratigraphic resolution in sections where biomagnetostratigraphy is poor or lacking, such as in the Arctic (McNeil and Miller, 1990), in shallow-water sections (Miller and others, 1988; Sugarman and others, 1993), and in the Oligocene (Hess and others, 1986). Because of the long residence time and uniform distribution of Sr isotopes in the oceans, mixed planktic and benthic foraminifera from a single sample can be analyzed without concern for potential surface vs. deep ocean signal bias.

Reference sections for $^{87}\text{Sr}/^{86}\text{Sr}$ analyzed in foraminiferal samples have been developed using magnetobiostratigraphic correlations (Fig. 1). Composite records from different sites have been developed because no single site contains a complete, unaltered Cenozoic section (DePaolo and Ingram, 1985; Hess and others, 1986; McKenzie and others, 1988; Hodell and others, 1991; Hodell and Woodruff, 1994). Farrell and others (1995) compiled a record for the late Neogene (0–7 Ma) from Ocean Drilling Program (ODP) Site 758. Hess and others (1986) produced a record from 137 samples for the entire Cenozoic. Miller and others (1991a) and Olsick and others (1994) developed a lower to middle Miocene reference section at North Atlantic Deep Sea Drilling Project (DSDP) Site 608 that has a very good magnetostratigraphic record, and compares well with DSDP Site 588 (Hodell and others, 1991). Miller and others (1988) and Reilly and others (2002) compiled an Oligocene Sr-isotope reference section from DSDP Site 522 that has an excellent magnetostratigraphy. Sr-isotope variations were very small during the Paleocene–Eocene, limiting stratigraphic resolution (Hodell and others, 2007). Rates of change in the Maestrichtian were sufficiently high to yield ± 1 Myr resolution when calibrated to paleomagnetism at South Atlantic DSDP Site 525 (Sugarman and others, 1993).

In this paper, we include a new Maestrichtian through Cenozoic compilation that uses only foraminiferal analyses (Fig. 1). The most comprehensive compilation is the “look-up table” of Howarth and McArthur (1997), which provides a record for the Jurassic to present using a variety of fossil types. However, the error analysis in their look-up tables does not reflect the full errors. Regressions provided by Olsick and others (1994) and Reilly and others (2002) for the interval 34–8 Ma provide a rigorous evaluation of age errors.

MINOR AND TRACE ELEMENTS

Minor and trace elements, which substitute for calcium (Ca) or carbonate ion (CO_3^{2-}) in the crystal latticework of calcareous foraminiferal tests, have been used to reconstruct past oceanic conditions. Because the uptake of these elements is often a function of ambient environmental conditions at the time of test formation, fossil tests can provide powerful tools for a range of environmental reconstructions. Elemental ratios reported relative to Ca (e.g., Mg/Ca, Cd/Ca) in biogenic carbonates are especially useful because they are measured on the same phase as $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$, reducing the uncertainties associated with using multiple proxies in reconstructions.

FORAMINIFERAL Mg/Ca PALEOTHERMOMETRY

As the ocean plays a key role in climate dynamics, a full understanding of the complexity of past climate change requires thorough knowledge of variations in ocean hydrography and circulation through time. Therefore, obtaining proxies for reconstructing the oceanic distribution of temperature and salinity has been a primary objective in paleoceanography. The oceanic residence time of minor elements is relatively long (10 Myr for Mg and 1 Myr for Ca), resulting in constant seawater Mg/Ca on timescales of $<10^6$ years (Broecker and Peng, 1982). In view of this long residence time, the possibility that temperature controls the Mg content of marine carbonates has been explored for decades. However, it was not until more recently that the prospects of using foraminiferal Mg/Ca for paleothermometry have been demonstrated under both controlled laboratory and field calibrations (Nürnberg, 1995; Nürnberg and others, 1996; Rosenthal and others, 1997a; Lea and others, 1999; Elderfield and Ganssen, 2000). Following these initial studies, there has been an explosion in studies exploring both the reliability of the new proxy, and its application for reconstructing past ocean temperatures (see below).

Planktic Foraminiferal Mg/Ca Paleothermometry

Mg/Ca-paleothermometry, measured in conjunction with $\delta^{18}\text{O}$, offers a distinct advantage over other SST proxies (e.g., faunal assemblage analysis and alkenone unsaturation index). Because Mg/Ca indicates the temperature at which the foraminiferal test precipitated, it allows the estimation of the $\delta^{18}\text{O}$ composition of seawater (δ_w) (Mashiotta and others, 1999), which in turn can provide the means to reconstruct changes in ice volume and/or salinity (e.g., Lea and others, 2000; Lear and others, 2000).

The temperature dependence of Mg uptake in planktic foraminiferal tests has been determined using three different calibration approaches: 1) controlled culture experiments where changes in growth parameters (e.g., light, temperature, salinity, and pH) are set independently of each other (e.g., Nürnberg and others, 1996; Lea and others, 1999; Russell and others, 2004; Langen and others, 2005); 2) measurements of planktic foraminiferal tests collected in sediment trap time-series from sites that are characterized by significant seasonality in SSTs (Anand and others, 2003; McConnell and Thunell, 2005; Huang and others, 2008; Mohtadi and others, 2009); and 3) analysis of fossil planktic foraminifera from core-top sediments (e.g., Elderfield and Ganssen, 2000). These calibrations suggest an exponential dependence of the Mg/Ca content of planktic foraminifera on temperature in the form:

$$\text{Mg/Ca} (\text{mmol mol}^{-1}) = B e^{AT}$$

where T is the calcification temperature in $^{\circ}\text{K}$. In this relationship, the exponential constant (A) reflects the Mg/Ca response to a given temperature change (in mmol mol^{-1} per $^{\circ}\text{K}$), implying increased sensitivity with temperature. An examination of available data obtained by these three independent calibration methods suggests a similar temperature sensitivity for all planktic species within the range

$0.9 \pm 0.2 \text{ mmol mol}^{-1} \text{ }^{\circ}\text{K}^{-1}$ (1 SD). The pre-exponential constant (B) differs among the species and depends on the water depth at which each of the species calcifies (Anand and others, 2003).

The good agreement among these three calibration methods provides robust evidence that the temperature signal imprinted during test formation is generally preserved after burial of foraminiferal tests, thereby supporting the use of planktic foraminiferal Mg/Ca for paleotemperature reconstructions. However, intra-species variability (Elderfield and Ganssen, 2000; Anand and others, 2003), diagenetic overprints including post-depositional dissolution (Rosenthal and others, 2000a; Rosenthal and Lohmann, 2002), variable contaminants (Lea and others, 2005; Pena and others, 2005; Pena and others, 2008), and methodological biases among laboratories (Rosenthal and others, 2004; Greaves and others, 2008) can introduce errors in absolute temperature reconstructions. Therefore, Mg/Ca paleothermometry is best used to estimate relative changes in seawater temperatures.

In general, inter-species variability in Mg/Ca is correlated with calcification depth of the planktic foraminiferal tests (Figs. 2, 4). Shallow mixed-layer dwellers (e.g., *Globigerinoides ruber* and *Gln. sacculifer*) have high Mg/Ca, whereas deep dwellers (e.g., *Globorotalia tumida* (Brady) and *Neogloboquadrina dutertrei* (d'Orbigny)) are characterized by relatively low Mg/Ca (Bender and others, 1975; Rosenthal and Boyle, 1993). This trend is consistent with the decrease in calcification temperature with depth, as recorded by the increase in $\delta^{18}\text{O}$ from mixed-layer to deep species (Fairbanks and others, 1980; Fairbanks and others, 1982). Accurate species-specific calibrations could thus allow for the reconstruction of the thermocline structure of past upper oceans (Regenberg and others, 2009). Even though this general trend is valid, it can be complicated by (1) species that calcify while migrating vertically through the water column (Hemleben and others, 1989), or (2) species that add an outer calcite crust prior to reproduction (called "gametogenic crust") at depths significantly deeper (and colder) than their principal habitat depth (e.g., *Gln. sacculifer*). These processes can cause significant variability in the Mg distribution within and among foraminiferal tests. Indeed, the "whole test" Mg/Ca composition that is most often used for temperature reconstructions is arguably a weighted average of calcite layers formed at different depths/temperatures (Lohmann, 1995; Rosenthal and others, 2000b; Benway and others, 2003). This effect is minimal in the shallow, mixed-layer species *Gln. ruber*, which does not produce a gametogenic crust. In contrast, it is expressed strongly in deep-dwelling species (e.g., *Neogloboquadrina dutertrei*, *Globorotalia tumida*, and *Gln. truncatulinoides* (d'Orbigny)).

There is also significant test size-dependent variability in Mg/Ca ratios among specimens of the same species. Mg/Ca increases with increasing test size in most species, possibly because small individuals calcify faster than large individuals or because larger individuals spend relatively more time near the sea surface, where the water is warmer and the higher light intensity increases symbiont photosynthetic activity, which in turn allows growth to larger sizes (Spero and Lea, 1993; Hönisch and Hemming, 2004). In addition,

Mg/Ca can vary among different morphotypes of the same species, reflecting differences in growth season (e.g., the white vs. pink variety of *Gln. ruber*; Anand and others, 2003) or depth habitat (e.g., *Gln. ruber* s.s. vs. *Gln. ruber* s.l.; Steinke and others, 2005). The intraspecific variability implies that Mg/Ca analyses and calibration should be performed on the same morphotypes and within a narrow size range. Also, when possible, the same size fraction should be used for both Mg/Ca and isotopic analyses.

Interpretations of planktic foraminiferal Mg/Ca in terms of temperature might be complicated by the fact that the uptake of Mg into the calcifying tests appears to depend on additional factors, including salinity and pH. Furthermore, dissolution on the seafloor may alter the primary Mg/Ca signal and bias paleotemperature estimates. Culture studies show a direct relationship between planktic foraminiferal Mg/Ca and seawater salinity on the order of $5 \pm 3\%$ per salinity unit (Lea and others, 1999; Kisakurek and others, 2008). In contrast, evidence from a surface sediment calibration in the Mediterranean Sea suggesting significantly higher salinity effect (e.g., $>16\%$ per psu for *Gln. ruber* (Ferguson and others, 2008) is still controversial and at odds with the culture studies, which cover a similar salinity range. Given a temperature sensitivity of $\sim 9\%$ per $^{\circ}\text{C}$, and the conservative salinity dependence, these non-temperature related effects translate to an uncertainty of $0.5 \pm 0.3^{\circ}\text{C}$ per 1 salinity unit. pH changes may exert an additional influence; culture data indicate an inverse Mg/Ca dependence of about $7 \pm 5\%$ per 0.1 pH unit, equivalent to $-0.8 \pm 0.5^{\circ}\text{C}$ per 0.1 pH unit (Lea and others, 1999). The experimental data indicate only minimal pH dependence under higher-than-ambient pH conditions (Russell and others, 2004). While significant, these uncertainties are most likely smaller than the glacial–interglacial variability during the Pleistocene. On longer time scales, however, non-temperature related effects might have been large enough to mask SST variability when surface salinity variations were substantial. For example, the large increase in Mg/Ca of *Gln. sacculifer* observed at ODP Site 1000 from 4.5–3.9 Ma has been attributed to a substantial increase in Caribbean Sea surface salinity as a result of the closure of the Panamanian Gateway (Groeneveld and others, 2008). On long time scales, variations in seawater Mg/Ca also needs to be accounted for when estimating secular changes in temperature. One promising method for constraining changes in the Mg content of the oceans is $\delta^{26}\text{Mg}$ (see below).

Interpretations of SST estimates based on planktic foraminiferal Mg/Ca are also confounded by the fact that the primary Mg/Ca ratio of planktic foraminiferal tests might be altered by post-depositional dissolution (Savin and Douglas, 1973; Lorens and others, 1977). The extent of diagenetic alteration increases with water depth and the accompanying decrease in the degree of calcite saturation (Rosenthal and Boyle, 1993; Russell and others, 1994; Brown and Elderfield, 1996). Preferential dissolution of Mg-rich calcite (which is more susceptible to dissolution) results in a systematic decrease of Mg/Ca ratios in planktic foraminiferal tests (Bè and others, 1975; Hecht and others, 1975; Brown and Elderfield, 1996), thereby shifting temperature estimates toward apparently colder

values (Rosenthal and others, 2000a). The decrease in Mg/Ca with depositional water depth is due to changes in each individual test, rather than due to the progressive loss of more dissolution-prone tests (Rosenthal and others, 2000a). This effect is higher in non-spinose, thermocline-dwelling species (e.g., *Glr. tumida* and *Nq. dutertrei*) than in spinose, shallow-dwelling species (e.g., *Gln. ruber* and *Gln. sacculifer*), reflecting the larger range of water temperatures over which the deep-dwellers calcify (Brown and Elderfield, 1996; Dekens and others, 2002; Huang and others, 2008). Based on core-top calibrations, various approaches have been proposed to test and correct for dissolution effects on Mg/Ca (Dekens and others, 2002; Rosenthal and Lohmann, 2002; Bassinot and others, 2004; Regenberg and others, 2006), but none are considered to completely compensate for the effect. However, with careful selection of samples and novel approaches to assess downcore dissolution effects on foraminiferal tests, errors associated with dissolution effects might be greatly minimized.

Benthic Foraminiferal Mg/Ca Paleothermometry

Whereas several additional proxies are available for estimating SST (e.g., faunal assemblage analysis, alkenone unsaturation index), Mg/Ca measurements in benthic foraminifera and ostracodes are the only alternative to $\delta^{18}\text{O}$ for reconstructing bottom-water temperature changes during the late Paleogene–Neogene, when large ice sheets affected the $\delta^{18}\text{O}$ of seawater, and hence, of foraminiferal calcite (Fig. 1). Because benthic foraminifera calcify on (or just below) the seafloor and are preserved in the same environment in which they lived, errors related to the exact calcification water depth and *post-mortem* alteration are minimized in comparison to planktic foraminifera.

The first benthic foraminiferal Mg/Ca calibration was based on the species *Cibicidoides pachyderma* (Rzehak) from a shallow bathymetric transect in Little Bahama Banks (LBB) that spanned a temperature range of $5\text{--}18^{\circ}\text{C}$ (Rosenthal and others, 1997b). Two subsequent studies that expanded on the LBB calibration supported the initial results suggesting that the temperature sensitivity of combined *Cibicidoides pachyderma* and *Planulina wuellerstorfi* (Schwager) Mg/Ca data is apparently identical with the $\sim 10\%$ increase in Mg/Ca per $^{\circ}\text{K}$ observed in planktic foraminifera (Lear and others, 2002; Martin and others, 2002). Similar temperature sensitivity has been observed in other benthic taxa, such as *Planulina* spp., *Oridorsalis umbonatus* (Reuss), and *Melonis* spp., whereas *Uvigerina* spp. and *Planulina arimenensis* d'Orbigny exhibited significantly lower sensitivity of about 6% per $^{\circ}\text{C}$ (Lear and others, 2002).

Using the above calibrations, paired $\delta^{18}\text{O}$ and Mg/Ca measurements in benthic foraminifera have been used to infer changes in seafloor temperature and whole-ocean seawater $\delta^{18}\text{O}$ (δ_w) composition. Combined, these records provided new insights on Cenozoic climate history and the evolution of the cryosphere. A low-resolution benthic foraminiferal Mg/Ca record showed a general global cooling through the Cenozoic, with the greatest cooling occurring during the Eocene and the Plio-Pleistocene (Lear

and others, 2000; Billups, 2002; Billups and Schrag, 2003). The inferred δ_w record suggested that the Cenozoic expansion of continental ice sheets occurred primarily in three steps, during the Eocene–Oligocene transition, the middle Miocene, and Plio–Pleistocene (Lear and others, 2000), consistent with the $\delta^{18}\text{O}$ record of Cenozoic temperature and ice-volume changes (Miller and others, 1987; Zachos and others, 2001b; Cramer and others, 2009).

While initially very promising, recent studies have raised concerns about the accuracy of the above calibrations. First, it has been argued that the shallow LBB samples are compromised by diagenetic calcite overgrowths enriched in Mg, yielding an apparent large exponential dependence on temperature. A new calibration study of *C. pachyderma* from depth transects in the Florida Straits and Great Bahama Banks, where the problem of diagenetic calcite overgrowth is minimal, suggests a linear relationship between Mg/Ca and BWT of $\text{Mg/Ca} = 0.116T + 1.2$ (i.e., a sensitivity of $\sim 0.12 \text{ mmol mol}^{-1} \text{ per } ^\circ\text{C}$) for the temperature range from 5.8 to 18.6°C (Marchitto and others, 2007). Using a micro-analytical technique (SIMS) to avoid contamination by diagenetic overgrowth, Curry and Marchitto (2008) obtained an identical calibration on individual *C. pachyderma* samples from Bahama Banks core tops.

At the colder end of the calibration, there is growing evidence for CO_3^{2-} control, which affects the incorporation of trace metals into benthic foraminiferal tests, thus significantly altering the Mg/Ca-temperature relationships in cold waters at low or under-saturated conditions with respect to carbonate minerals (Elderfield and others, 2006; Rosenthal and others, 2006). Using a global dataset, Elderfield and others (2006) proposed a carbonate ion-Mg/Ca sensitivity of $0.0086 \text{ mmol mol}^{-1} \text{ per } \mu\text{mol kg}^{-1}$ in calcitic benthic tests that becomes significant below 3°C. Using a similar deep-ocean calibration, Healey and others (2008) found identical sensitivity to CO_3^{2-} and a similar threshold (i.e., $\sim 3^\circ\text{C}$). Interestingly, their data from the warm Sulu Sea (10°C) fall right on the Marchitto and others (2007) calibration, thus lending support for a dominant temperature effect at $>3\text{--}4^\circ\text{C}$. At colder temperatures, carbonate saturation effects may dominate, thus altering the Mg/Ca-temperature relationship (Yu and Elderfield, 2008).

The aragonitic *Hoeghundina elegans* (d'Orbigny) may be particularly useful for reconstructing bottom-water temperatures. This species has a widespread distribution (Atlantic, Pacific, and Indian Oceans), but because of its high dissolution susceptibility, its occurrence in the geological record is sporadic in deepwater sections. Recently, core-top calibrations from Atlantic and Pacific locations showed that Mg/Ca in *Hoeghundina elegans* may be used to reconstruct thermocline temperatures (Rosenthal and others, 2006). However, there is also evidence for additional significant CO_3^{2-} effect on the test Mg/Ca at low carbonate saturation levels.

Culture studies of trace elements in *Bulimina aculeata* and *Rosalina vilardeboana* yielded similar results to field studies, and show that while water chemistry is the major environmental influence on trace elemental ratios in benthic foraminifera, there are interspecies and size offsets in these

ratios that can be attributed to biological processes (Hintz and others, 2006a; Hintz and others, 2006b). As with $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$, interspecies offsets in Mg/Ca are typically taken into account in paleoceanographic studies (e.g., Lear and others, 2000). Size-related variability in Mg/Ca can be minimized by analyzing a narrow test size range (Hintz and others, 2006b).

While the exact magnitude of the various effects (especially temperature and $[\text{CO}_3^{2-}]$) is still under investigation, it is becoming clear that benthic foraminiferal Mg/Ca thermometry is not as straightforward as initially thought. Although it is still likely that in warm, saturated waters (e.g., upper thermocline), secondary effects may have only a small influence on seafloor temperature estimates from Mg/Ca, they may lead to a large bias in deep waters where changes in temperature and CO_3^{2-} saturation occur concomitantly.

In conclusion, at this time there is a large uncertainty about the application of benthic foraminiferal Mg/Ca for estimating deep-ocean temperatures in cold and unsaturated conditions. There is also some uncertainty about the nature of the calibration under warm temperatures and over-saturated conditions (Bryan and Marchitto, 2008). Interestingly, estimation of Cenozoic bottom-water temperatures remain largely consistent using the calibration of Lear and others (2002) (Katz and others, 2008; Cramer and others, 2009).

Mg Isotopes ($\delta^{26}\text{Mg}$)

As discussed above, the Mg/Ca ratio in foraminiferal calcite is strongly influenced by environmental parameters, including temperature and seawater Mg content. An emerging proxy that holds promise for providing constraints on Mg/Ca changes through time is $\delta^{26}\text{Mg}$. In contrast to Mg/Ca, the Mg-isotope uptake ($\delta^{26}\text{Mg}$) shows no significant correspondance to environmental changes, such as temperature or pH (Pogge von Strandmann, 2008). Rather, mechanisms that dominate Mg isotopic fractionation are continental weathering (Mg source) and dolomite formation (Mg sink). Very little fractionation occurs during hydrothermal exchange, and ion-exchange reaction with clays is a very small sink. Hence, changes in foraminiferal $\delta^{26}\text{Mg}$ should reflect changes in the Mg sources and sinks through time. The 10 Myr residence time of Mg in the oceans (Berner and Berner, 1996) is ideally suited to constrain long-term global variations in source and sink Mg fluxes (Pogge von Strandmann, 2008).

TRACE ELEMENT NUTRIENT PROXIES

Unlike Mg and Ca, which exhibit conservative behavior in the ocean (i.e., maintain constant proportion with salinity), the distribution of the trace elements cadmium (Cd), zinc (Zn), and barium (Ba) in the ocean varies spatially and temporally on much shorter time scales. These trace elements exhibit a nutrient-like behavior, being depleted in surface waters and enriched at depth, following the cycling of organic matter and its redistribution by deepwater circulation (Bruland, 1983). These trace elements have initially been explored as potential proxies for nutrient paleochemistry, as it relates to changes in productivity.

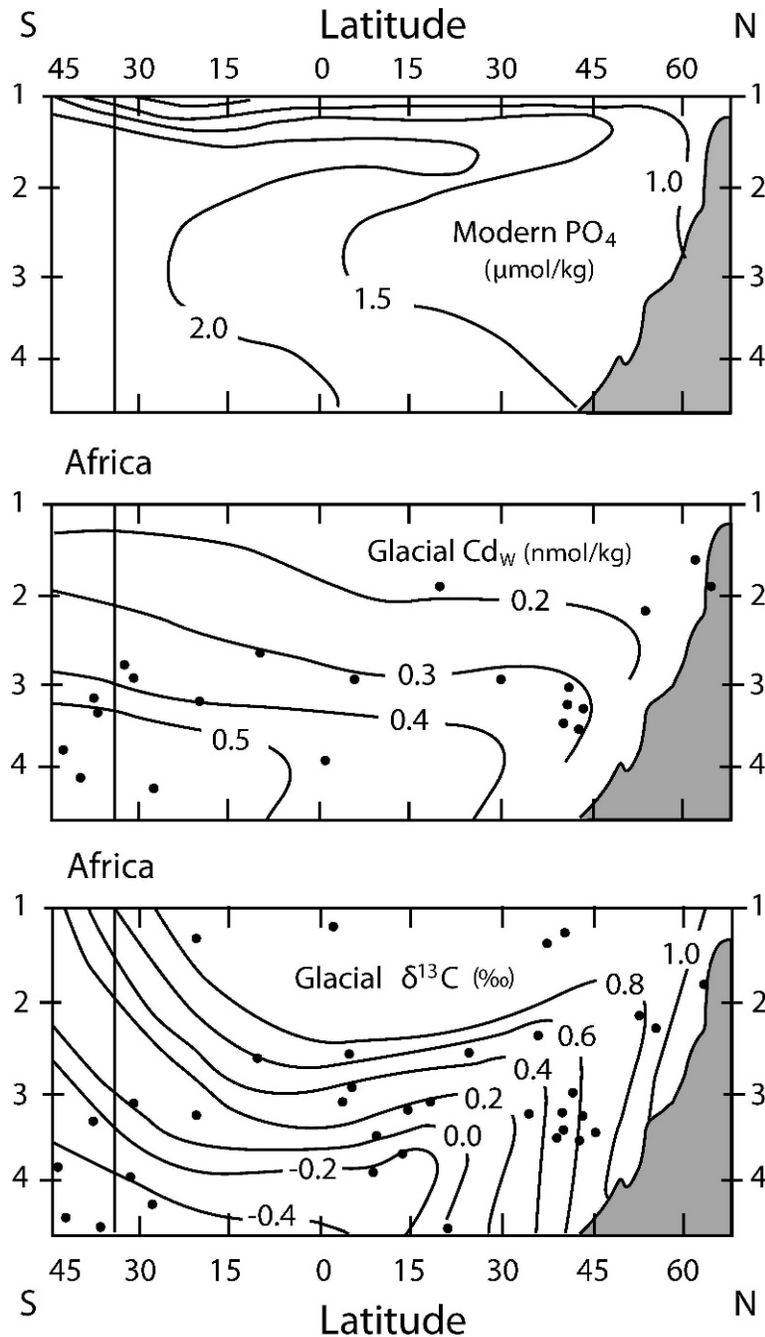


FIGURE 5. Geochemical sections through the western Atlantic Ocean depicting: the modern distribution of labile nutrient phosphate (A), and the glacial distributions of Cd (B) and $\delta^{13}\text{C}$ (C) in seawater, reconstructed from analyses of benthic foraminifera in cores at different water depths. In the same way that phosphate is used in the modern ocean to trace the different deep ocean masses, the seawater distribution of Cd (Cd_w) and $\delta^{13}\text{C}$ are used as nutrient (phosphate) analogs to reconstruct variations in past ocean circulation. Although similar, Cd_w , unlike $\delta^{13}\text{C}$, is not affected by air-sea CO_2 exchange. Dots indicate depth and latitude of each sample analyzed. The vertical line marked as AFRICA delineates the transition into the Antarctic Circumpolar Current at the southern extent of the Atlantic Ocean (modified from Rosenthal [1994]).

However, subsequent studies have shown that these proxies may be better used to reconstruct deepwater circulation. This approach takes advantage of the fact that the distribution of Cd, Ba, and Zn in the ocean “tags” the different deepwater masses, increasing substantially with water “aging” from the North Atlantic to the North Pacific. Hence, a downcore record from sites with strong mixing gradients primarily will reflect changes in deepwater

circulation and, to a lesser extent, global ocean chemistry (e.g., Boyle, 1988, 1990).

The application of trace-element proxies has played a key role in understanding recent glacial–interglacial variability. However, there has been very limited work on longer timescales (e.g., Delaney and Boyle, 1987). This is largely because of the relatively short oceanic residence time of trace elements (on the order of 10^4 – 10^5 years) and the

uncertainty about the long-term controls on their oceanic distributions. For example, on these time scales, changes in the extent of reducing (suboxic and anoxic) sediments may significantly alter the oceanic inventory of Cd and Zn, independent of variations in ocean circulation. This is because precipitation of sulphidic minerals (e.g., CdS and ZnS) in reducing sediments is the major oceanic sink for these elements (Rosenthal and others, 1995a; Rosenthal and others, 1995b).

Cadmium (Cd/Ca)

The oceanic distribution of dissolved Cd is highly correlated with that of phosphate (PO_4) (Fig. 5). Like PO_4 , the vertical distribution of Cd is characterized by a concentration maximum at ~ 1 km water depth, reflecting its association with the degradation of labile organic matter. Although evidence suggests that Cd plays an important enzymatic role, especially at low Zn conditions (Cullen and others, 1999), it is not clear that this is sufficient to explain its ocean-wide nutrient-like distribution. The global Cd- PO_4 correlation is significant, but follows a slight curve that has been attributed to differential uptake rates of Cd and PO_4 during photosynthesis that is likely related to Fe bioavailability (Cullen, 2006; Hendry and others, 2008). Therefore, the modern Cd-P relationships in deep waters is described using two equations: 1) for $P < 1.3$, $\text{Cd}_w = 0.21 \times P$; and 2) for $P > 1.3$, $\text{Cd}_w = 0.40 \times (P - 1.3) + 0.279$, where P is in $\mu\text{mol kg}^{-1}$ and Cd_w is in nmol kg^{-1} (Boyle, 1988).

The high variability of, and uncertainties about, the controls on Cd/ PO_4 ratios in surface waters and evidence for possible strong temperature control on the incorporation of Cd into planktic tests (Rickaby and Elderfield, 1999) have limited the application of planktic foraminiferal Cd/Ca for reconstructing biological productivity. Furthermore, there is also a suggestion of possible diagenetic alteration of the primary Cd/Ca signal in planktic tests that may add another complication (Ripperger and others, 2008). Instead, the real strength of this proxy has been in reconstructing deepwater circulation by taking advantage of the fact that like phosphorous, Cd distribution in the deep ocean reflects the flow-related mixing of different deepwater masses (Fig. 5), increasing about five-fold between the deep North Atlantic and North Pacific Oceans (e.g., Boyle and Keigwin, 1985, 1987; Boyle, 1988, 1990).

Benthic foraminiferal Cd/Ca ratios vary proportionally to seawater Cd concentrations, a relationship that can be expressed in terms of the partition coefficient:

$$D_{\text{Cd}} = (\text{Cd/Ca})_{\text{foram}} / (\text{Cd/Ca})_{\text{seawater}}$$

The D_{Cd} of calcareous benthic foraminifera typically used for geochemical reconstructions (e.g., *Cibicides mundulus* (Brady, Parker, and Jones), *Oridorsalis umbonatus*, *Planulina wuellerstorfi*, and *Uvigerina* spp.) varies with water depth, linearly increasing from 1.3 at depths ≤ 1.3 km to 2.9 at depths > 3 km (Boyle, 1992). D_{Cd} of cultured *Bulimina aculeata* is consistent with these taxa from intermediate water depths (Hintz and others, 2006a). Studies showing that temperature does not have a significant influence on the uptake of Cd into foraminiferal calcite (Rosenthal and

others, 1997a; Marchitto, 2004) suggested that the depth-dependent change in D_{Cd} might be related to changes in the degree of calcite saturation, which is evident in the highly undersaturated water of the deep Pacific (McCorkle and others, 1995; Marchitto and others, 2000). In contrast, the aragonitic benthic foraminifera *Hoeglundina elegans* has a D_{Cd} of 1.0, which is invariant with depth (Boyle and others, 1995) or temperature (Rosenthal and others, 1997a).

Benthic foraminiferal Cd/Ca reconstructions have provided important insights on glacial-interglacial variability in deep-ocean circulation (Fig. 5). In particular, the last glacial maximum (LGM) reconstruction indicates reduced extent of low-nutrient North Atlantic Deep Water (NADW) in the deep Atlantic Ocean and its presence as a shallower glacial North Atlantic Intermediate Water (GNAIW) (Boyle and Keigwin, 1987; see review by Marchitto and Broecker, 2006). Additional important information might be gained from coupled $\delta^{13}\text{C}$ and Cd measurements, because the two tracers have somewhat different controls. Specifically, air-sea exchange affects $\delta^{13}\text{C}$ but not Cd, so combining the two can potentially reveal past air-sea processes. Applying this approach to various regions of the glacial ocean have shown that different deepwater masses had different air-sea signatures, allowing for the separation of biogeochemical aging and water-mass mixing (Lynch-Stieglitz and Fairbanks, 1994).

Zinc (Zn/Ca)

The distribution of dissolved Zn in the ocean mimics that of the nutrient silica; both are depleted in surface waters and gradually increase with depth. Unlike labile constituents (PO_4 , Cd), Zn behaves like a refractory nutrient, with no distinct intermediate-depth concentration maximum. Zinc is an essential constituent for algal metalloenzymes, especially in diatoms, and this may explain its oceanic association with silica. Zn concentrations increase more than ten-fold between the deep North Atlantic and North Pacific, and therefore, it was initially thought that foraminiferal Zn/Ca may be a very sensitive tracer of past deepwater circulation. However, further research has shown that the partition coefficient of Zn in calcareous benthic foraminifera strongly depends on saturation state. In saturated waters, D_{Zn} is ~ 9 and decreases to about ~ 4 in undersaturated conditions, such as found in the modern deep Pacific (Marchitto and others, 2000). Consequently, downcore benthic foraminiferal records may reflect not only changes in the mixing between the different deepwater masses, but also in situ variations in carbonate saturation state. For example, the glacial increase in deep North Atlantic Zn due to the increased presence of Antarctic Bottom Water (AABW) (high Si, high Zn, low $[\text{CO}_3^{2-}]$) was partially muted in Zn/Ca because of AABW's lower saturation state.

The strong relationship between D_{Zn} and saturation state allows for reconstruction of past deep ocean carbonate chemistry. Using this approach, Marchitto and others (2005) have reconstructed deep equatorial Pacific CO_3^{2-} over the last glacial-interglacial cycle. Their data support the "CaCO₃ compensation hypothesis" (Broecker and Peng, 1987), showing a transient drop in deep-ocean

CO_3^{2-} at the beginning of the LGM, associated with the transfer of excess metabolic CO_3^{2-} from the upper ocean to the deep ocean, and a subsequent reversal due to seafloor dissolution of CaCO_3 (Marchitto and others, 2005).

Barium (Ba/Ca)

Dissolved Ba displays nutrient-like vertical distribution, with low concentrations at the surface that gradually increase with depth, similar to that of biogenic “hard parts” (CaCO_3 and SiO_2) (Chan and others, 1977). The modern distribution of dissolved Ba resembles that of alkalinity, but the association might be fortuitous; whereas biogenic CaCO_3 skeletons are the main carrier of alkalinity to the deep sea, the Ba export to the deep ocean is primarily through marine barite formed in decaying organic matter (Lea and Boyle, 1989; Lea, 1993, 1995). Furthermore, a recent study suggests that unlike CaCO_3 dissolution that occurs largely on the seafloor, Ba is preferentially regenerated in the water column (Rubin and others, 2003). Consequently, despite the strong correlation between Ba and alkalinity in the modern ocean, its application for reconstructing past ocean alkalinity might be limited. However, as for Cd, the large Atlantic–Pacific difference in Ba offers the possibility to reconstruct deepwater circulation from benthic foraminiferal Ba/Ca (Lea and Boyle, 1989; Lea, 1990).

For the LGM, benthic foraminiferal Ba/Ca supports the view inferred from Cd/Ca and $\delta^{13}\text{C}$ that the low-nutrient NADW was replaced by the shallower GNAIW, while the nutrient-rich AABW expanded much further northward than at present (Lea and Boyle, 1990). In contrast to $\delta^{13}\text{C}$ and Cd/Ca, however, the benthic foraminiferal Ba/Ca shows no apparent interbasinal gradient between the deep Atlantic and Pacific oceans. This indicates that the Ba-alkalinity relationship in the modern ocean might be different than it was in the glacial ocean, or that there are other, hitherto undiscovered, factors controlling the partitioning of Ba into benthic foraminiferal tests.

A potentially more promising application is using planktic foraminiferal Ba/Ca to reconstruct past variability in riverine and glacial meltwater inputs to the ocean. River waters are concentrated in barium compared to the depleted surface ocean. A further enrichment occurs in the river–sea mixing zone as Ba desorbs from suspended particles in intermediate salinity waters (e.g., Hanor and Chan, 1977). Taking advantage of the fact that Ba incorporates into planktic foraminiferal tests in direct proportion to its seawater concentration and does not vary with temperature or salinity (Lea and Boyle, 1991; Lea and Spero, 1992, 1994), Ba/Ca has been successfully used to reconstruct variations in fluvial inputs to the ocean in the past (Weldeab and others, 2007). This relatively new application shows the greatest promise for the use of Ba as a proxy in paleoceanography.

PROXIES FOR pH AND CARBONATE ION CONCENTRATION

Paleo-pH and carbonate ion concentration [CO_3^{2-}] have been estimated from the variation in size-normalized test

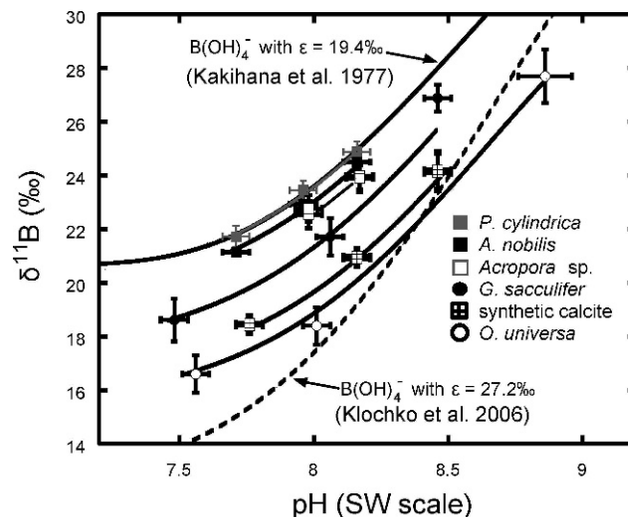


FIGURE 6. Empirical and theoretical calibration curves of $\delta^{11}\text{B}$ of aqueous borate (Kakihana and others, 1977; Klochko and others, 2006) and marine carbonates (Sanyal and others, 1996; Sanyal and Nugent, 2000; Sanyal and others, 2001; Hönisch and Hemming, 2004; Reynaud and others, 2004) versus seawater pH. Note that marine carbonates are $\sim 17\%$ lighter than $\delta^{11}\text{B}$ of seawater ($\sim 39.8\%$), suggesting borate must be the dominant species incorporated into marine carbonates (Hemming and Hanson, 1992). Although offset from each other, empirical carbonate calibrations parallel $\delta^{11}\text{B}$ of borate after Kakihana and others (1977) rather than the experimental curve of Klochko and others (2006), which demonstrates the need to use species-specific empirical carbonate curves for paleo-pH reconstructions.

weights of planktic foraminifera, with mixed results (e.g., Lohmann, 1995; Broecker and Clark, 2004). This method states that fossil test weight is a function of: 1) the test weight while the organism was alive, assuming it is determined solely by surface water [CO_3^{2-}]; and 2) carbonate loss after deposition, resulting from dissolution. The greater the dissolution, the lower the [CO_3^{2-}] at the seafloor. However, there are additional factors that may affect fossil test weight, which can complicate interpretation solely in terms of [CO_3^{2-}]. Dissolution that occurs upon initial deposition cannot be distinguished from post-burial porewater dissolution, the initial test wall thickness depends on growth conditions, and environmental parameters (e.g., temperature, nutrient availability) can make it difficult to link test weight directly and exclusively to [CO_3^{2-}] (Broecker and Clark, 2004). Nonetheless, test weights of planktic foraminifera have been linked to surface ocean [CO_3^{2-}] (Barker and Elderfield, 2002; Moy and others, 2009).

There are three emerging geochemical proxies for seawater pH and [CO_3^{2-}] that offer alternatives to the test weight method. Boron isotope ($\delta^{11}\text{B}$) analyses of planktic foraminifera are the most widely studied so far. $\delta^{11}\text{B}$ in benthic foraminifera, B/Ca, and U/Ca also show promise. In addition, the nutrient tracer Zn/Ca can also be used as a [CO_3^{2-}] proxy (see above).

BORON ISOTOPES ($\delta^{11}\text{B}$)

Boron isotopes provide a proxy for past seawater pH, based on the equilibrium reaction between the two dominant species of B in seawater, boric acid ($\text{B}(\text{OH})_3$) and borate

($B(OH)_4^-$). At low pH (<7), almost all B is present in the form of $B(OH)_3$, which is enriched in ^{11}B . At high pH (>10), it is present in the form of $B(OH)_4^-$, which is enriched in ^{10}B (Fig. 6). Therefore, $\delta^{11}B$ increases with increasing pH.

Boron isotope ratios are traditionally measured by thermal ionization mass spectrometry (TIMS), although recent developments in the field of MC-ICP-MS methods now also allow analysis of small samples (Foster, 2008). Analogous to $\delta^{18}O$ and $\delta^{13}C$, the boron isotopic composition is reported relative to a standard, in δ notation:

$$\delta^{11}B_{\text{sample}} = 1000 \times (R_{\text{sample}}/R_{\text{standard}} - 1)$$

where R is the isotopic ratio measured. Because marine carbonates record a boron isotopic composition that falls close to the $\delta^{11}B$ of borate, Hemming and Hemming (1992) suggested that borate must be the boron species adsorbed and incorporated into the $CaCO_3$ lattice. This has been supported in a number of empirical calibration studies (Sanyal and others, 1996; Sanyal and Nugent, 2000; Sanyal and others, 2001; Hönisch and Hemming, 2004; Reynaud and others, 2004), where biogenic and inorganic carbonates have been grown over a wide range of seawater pH (Fig. 6). Based on the boron speciation in seawater and the respective isotopic composition, pH can be inferred from the measured $\delta^{11}B_{\text{carbonate}}$ by:

$$pH = pK_B - \log\left(-(\delta^{11}B_{\text{seawater}} - \delta^{11}B_{\text{carbonate}} + a) / (\delta^{11}B_{\text{seawater}} - \alpha \times (\delta^{11}B_{\text{carbonate}} - a) - \varepsilon)\right)$$

where pK_B is the equilibrium constant of the boric acid/borate system (after Dickson, 1990), $\delta^{11}B_{\text{seawater}} = 39.5\%$, fractionation $\varepsilon = (\text{fractionation factor } \alpha - 1) \times 1000$, and 'a' is a constant offset between the theoretical borate curve and empirical calibration data for a specific sample type.

A complication in the interpretation of this system arises from the mismatch between the measured boron isotope fractionation in synthetic seawater (Klochko and others, 2006) and the shallower $\delta^{11}B/pH$ -curves described by the marine carbonates, which fall close to and follow the shape of the theoretical borate curve calculated from the fractionation factor determined by Kakihana and others (1977) (Fig. 6). A mechanistic explanation for this discrepancy has not yet been found, but may be related to analytical matrix effects. Because it is unacceptable to use a fractionation factor that does not fit the data, translating $\delta^{11}B$ to seawater pH has to rely on the species-specific empirical calibrations. Using the above $\delta^{11}B/pH$ equation requires the application of the theoretical fractionation factor by Kakihana and others (1977), which is statistically the factor that best describes the shape and inflection of all empirical carbonate calibrated to date (Hönisch and others, 2007).

Boron Isotopes in Planktic Foraminifera—Vital Effects and Dissolution

Of all marine carbonates, B isotopes have been studied most intensively in planktic foraminifera. The $\delta^{11}B/pH$ relationship of cultured *Gln. sacculifer* and *O. universa* has been calibrated (Fig. 6; Sanyal and others, 1996; Sanyal and others, 2001), and light experiments have quantified the effect of symbiont photosynthesis on the integrated pH

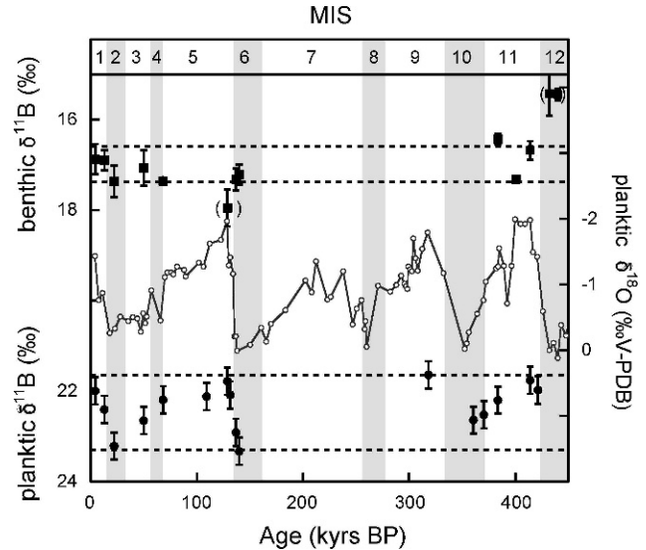


FIGURE 7. Glacial-interglacial variations in planktic and benthic foraminiferal $\delta^{11}B$, with planktic $\delta^{18}O$ for reference. The range of benthic $\delta^{11}B$ is approximately half the range of planktic $\delta^{11}B$, and indicates the small glacial pH increase of ~ 0.1 unit in the deep sea, compared to ~ 0.2 units in the surface ocean (from Hönisch and others, 2008).

recorded by $\delta^{11}B$ in *O. universa*, which is elevated relative to the pH of the surrounding seawater (Hönisch and others, 2003; Zeebe and others, 2003).

This vital effect can also be observed in different test sizes of *G. sacculifer*, where $\delta^{11}B$ increases with test size (Hönisch and Hemming, 2004; Ni and others, 2007). Hönisch and Hemming (2004) proposed that larger tests with higher $\delta^{11}B$ reflect growth in shallower habitat, resulting in higher symbiont photosynthetic activity and therefore higher microenvironment pH. Their depth transect studies showed that $\delta^{11}B$ decreases in deeper cores, and that the effect is stronger in the smaller sizes. This is consistent with the higher surface area to volume ratio of smaller tests, exposing more surface area to the corrosive bottom waters. This finding is supported by test weights and SEM imaging at one Atlantic site, which show that gametogenic calcite was removed during dissolution. In contrast, Ni and others (2007) found no $\delta^{11}B$ -variation with *G. sacculifer* size, and proposed that $\delta^{11}B$ is originally uniform in all test sizes, and that dissolution causes the size effect. However, the bottom waters of all core sites studied by Ni and others (2007) are supersaturated with respect to calcite, their scanning electron microscope (SEM) investigation revealed that gametogenic calcite was preserved, and there is no test weight data to support their dissolution hypothesis.

The actual mechanisms for lowering $\delta^{11}B$ in partially dissolved foraminifera remains a matter of debate. Nonetheless, an important observation is that $\delta^{11}B$ of *G. sacculifer* tests $>425 \mu m$ is the same for all sites (Hönisch and Hemming, 2004; Ni and others, 2007), indicating that large tests reflect surface seawater pH. Because these are also the tests that are the least affected by dissolution, the use of large tests is recommended for paleoreconstructions (Hönisch and Hemming, 2004). In contrast to *G. sacculifer*, Ni and others (2007) demonstrated that the symbiont-

bearing *Gln. ruber* does not show a test size effect. The effect of dissolution on this and other species remains to be tested further.

Paleoreconstructions Based on Boron Isotopes in Planktic Foraminifera

Surface ocean pH-reconstructions have focused primarily on the late Pleistocene. Ocean areas where aqueous $p\text{CO}_2$ is in equilibrium with the atmosphere can be used for reconstructing past atmospheric $p\text{CO}_2$ variations. Such studies found glacial surface ocean pH consistently ~ 0.15 pH units higher than during interglacials (Sanyal and others, 1995; Sanyal and others, 1997; Hönisch and Hemming, 2005), and estimated aqueous $p\text{CO}_2$ by pairing the pH estimates with paleoalkalinity, which is assumed to have changed only in response to salinity variations in the late Pleistocene (Fig. 7). The aqueous $p\text{CO}_2$ estimates match atmospheric $p\text{CO}_2$, as measured in Antarctic ice cores within ± 30 μatm , demonstrating the high quality and potential of B isotope reconstructions.

These initial validation studies led to attempts to use $\delta^{11}\text{B}$ to reconstruct the evolution of Cenozoic atmospheric $p\text{CO}_2$, a time when benthic foraminiferal $\delta^{18}\text{O}$ and Mg/Ca reconstructions indicate large variations in Earth's climate (Miller and others, 1987; Lear and others, 2000; Zachos and others, 2001b; Cramer and others, 2009). Pearson and Palmer (2000) used planktic foraminiferal $\delta^{11}\text{B}$ to estimate surface water pH and atmospheric $p\text{CO}_2$ over the past 60 Myr. Whereas their Neogene pH and $p\text{CO}_2$ estimates are similar to late Pleistocene and Holocene records, their Paleogene pH values are as low as 7.4, and $p\text{CO}_2 > 3000$ ppm. These estimates have been questioned for various reasons, including the use of the same $\delta^{11}\text{B}/\text{pH}$ calibration curve for various sizes and extinct foraminifera, uncertainties in the $\delta^{11}\text{B}$ composition of seawater in the past (the ocean residence time of boron is only ~ 14 – 20 Myr, Spivack and Edmond, 1987), and uncertainties in the DIC concentration in seawater, which Pearson and Palmer (2000) used to pair the pH estimates for estimating $p\text{CO}_2$ (Demico and others, 2003). Much more effort is required to solve these issues before Cenozoic pH can be estimated accurately from $\delta^{11}\text{B}$.

Upwelling areas are sources of CO_2 to the atmosphere, and $\delta^{11}\text{B}$ can help to evaluate the relative strength of these sources in the past. For instance, Sanyal and others (1997) and Sanyal and Bijma (1999) studied Holocene and glacial upwelling in the eastern equatorial Pacific and off northwest Africa, respectively. Whereas glacial pH was 0.2 units higher off northwest Africa, pH remained constant in the eastern equatorial Pacific, indicating that northwest African upwelling was a smaller source of CO_2 to the atmosphere than it is today. The data generated for these reconstructions are sparse, and many more cores from all major upwelling areas are needed to accurately quantify the effect of upwelling on atmospheric $p\text{CO}_2$.

Another study focused on the transition from the LGM into the Holocene in the western equatorial Pacific (Palmer and Pearson, 2003). Between 15,600 and 13,800 years before present, surface water pH dropped below modern values, indicating that this area temporarily turned into a strong

source of CO_2 to the atmosphere. Palmer and Pearson (2003) explained this observation with increased upwelling intensity in the eastern equatorial Pacific, which could have played an important role in the deglacial $p\text{CO}_2$ rise.

$\delta^{11}\text{B}$ in Benthic Foraminifera

The study of $\delta^{11}\text{B}$ in benthic foraminifera is in its infancy. The first studies by Sanyal and others (1995; 1997) used mixed benthic foraminifera; their reconstructions for glacial conditions yielded higher alkaline pH (+0.3 to +0.5 pH units, $\geq +100$ $\mu\text{mol kg}^{-1}$ in bottom water [CO_3^{2-}]) compared to Holocene bottom water (Fig. 7). Although such high bottom-water pH could explain the lower glacial atmospheric $p\text{CO}_2$, the data are inconsistent with deep-sea calcite preservation records (Sanyal and others, 1995; Anderson and Archer, 2002), which cast doubt on the validity of benthic foraminiferal $\delta^{11}\text{B}$. The concerns were subsequently corroborated by results of newly emerging deep-sea CO_3^{2-} proxies, such as size-normalized test weights of planktic foraminifera (e.g., Broecker and Clark, 2004), benthic foraminiferal Zn/Ca (Marchitto and others, 2005), and benthic foraminiferal B/Ca (Yu and Elderfield, 2007), which consistently indicate changes on the order of at most ± 10 to 30 $\mu\text{mol kg}^{-1}$, depending on location and water depth. A likely explanation for the unreasonable and highly scattered benthic $\delta^{11}\text{B}$ data is the use of mixed species for the reconstructions. Whereas epifaunal benthic species should reflect bottom-water pH, the contribution of infaunal benthic species would introduce a porewater pH signal. Alternatively, matrix or vital effects could cause a wide range of $\delta^{11}\text{B}$ offsets from the borate curve, analogous to the offsets observed between planktic foraminifera and corals (Fig. 6). Because the desired parameter is bottom-water pH, studies need to focus on an epifaunal benthic species, such as *P. wuellerstorfi*. However, the low abundance of benthic foraminifera in sediment samples, and the supposedly low boron concentration, seemed to prohibit $\delta^{11}\text{B}$ analyses of single benthic species.

The first attempt to measure $\delta^{11}\text{B}$ in the single epifaunal benthic species *Planulina wuellerstorfi* found high B concentrations of ~ 25 ppm (Hönisch and others, 2008). This result was unexpected because planktic foraminifera contain only 7–10 ppm boron. Recent ICP-MS analyses comparing B/Ca ratios in multiple benthic foraminiferal species indicate that *P. wuellerstorfi* may be the benthic species with the highest B concentration (Yu and Elderfield, 2007). This allows $\delta^{11}\text{B}$ analysis with as few as 10 tests. Hönisch and others (2008) calibrated *P. wuellerstorfi* $\delta^{11}\text{B}$ from core-tops across a wide depth range (1000–4500 m) on the Walvis Ridge. A large negative offset was found from $\delta^{11}\text{B}$ of borate, but this offset was constant across the entire depth range and does not indicate a dissolution effect. Application to last glacial sediments and a downcore study on the Sierra Leone Rise revealed a much smaller $\delta^{11}\text{B}$ change compared to the previous mixed benthic species reconstruction. This puts benthic foraminiferal $\delta^{11}\text{B}$ in line with other deep ocean carbonate chemistry proxies, and although laboratory calibration over a much larger pH range should be acquired, $\delta^{11}\text{B}$ analysis in benthic foraminifera now appears promising.

TRACE ELEMENT pH AND [CO₃²⁻] PROXIES*Boron (B/Ca)*

Analyzing B/Ca ratios in foraminiferal tests to reconstruct paleo-pH is an exciting new approach that may allow for a much faster sample throughput than analysis of $\delta^{11}\text{B}$ (Yu and Elderfield, 2007). Similar to the $\delta^{11}\text{B}$ proxy, B/Ca is based on the equilibrium reaction between boric acid and borate in seawater. $\delta^{11}\text{B}$ studies indicate that borate is the species incorporated into marine carbonates, and because the seawater borate concentration increases with pH, B uptake should increase at higher pH. Coretop studies indicate that B/Ca in foraminiferal calcite is strongly influenced by ambient seawater pH, CO₃²⁻, or temperature (Yu and others, 2007; Foster, 2008), although the studies disagree on the specific control. Yu and others (2007) have suggested that Mg/Ca temperature estimates can be used to extract seawater pH changes from planktic foraminiferal B/Ca records. A study of B/Ca in benthic foraminifera indicates that carbonate saturation in the deep sea influences the uptake of B in benthic foraminifera (Yu and Elderfield, 2007; Foster, 2008). A potential complicating factor in validating new paleoproxies is that carbonate chemistry and temperature often change in unison. Culture experiments under controlled laboratory conditions have been carried out to better quantify the effects of pH, CO₃²⁻, temperature, salinity, and B concentration on B/Ca in planktic foraminifera (Allen and others, 2009).

Uranium (U/Ca)

U/Ca in planktic foraminifera shows promise as a proxy for seawater [CO₃²⁻] (Russell and others, 1996; Russell and others, 2004). In culture experiments, U/Ca in *O. universa* and *Globigerina bulloides* d'Orbigny decreases with increasing [CO₃²⁻], with a species-specific offset (Russell and others, 2004). A similar relationship has been reported in corals, but the results were also consistent with a temperature control for U/Ca (Min and others, 1995; Shen and Dunbar, 1995). In contrast, the *O. universa* culture experiment indicated no relationship between U/Ca and temperature at 15–25°C (Russell and others, 2004). Down-core U/Ca records in planktic foraminifera were used to reconstruct Holocene-last glaciation surface water [CO₃²⁻], and the results show good agreement with $\delta^{11}\text{B}$ and shell weights (Russell and others, 2004).

NEODYMIUM ISOTOPES (ϵ_{Nd})

An exciting new geochemical proxy is neodymium isotope ratios measured in foraminifera, which incorporate the seawater Nd isotope ratio as they calcify. This promises to become a powerful new tool for reconstructing past ocean structure.

Seawater Nd isotopes (¹⁴³Nd/¹⁴⁴Nd, expressed as ϵ_{Nd}) are influenced by continental source rock region, and can be used to track changes in ocean circulation because Nd has a short residence time (~500–1000 years) relative to ocean mixing (~1500 years) (Goldstein and Hemming, 2003 and references therein). The variability of ϵ_{Nd} in the oceans is due to the distribution of old vs. young continental crust,

particularly near regions of deepwater formation. The most extreme end-members in the modern ocean are the deep North Atlantic (with low ϵ_{Nd} , ~-13.5, derived from old continental crust) and North Pacific (with high ϵ_{Nd} , from 0 to -4, acquired from young volcanic arcs) (Goldstein and Hemming, 2003, and references therein). Variations in ϵ_{Nd} with water mass are seen both between ocean basins and in depth profiles (von Blanckenburg, 1999; Vance and others, 2004), belying its use as a water mass tracer.

Marine apatite (especially fish debris) and authigenic Fe-Mn crusts and nodules are well established archives of past seawater ϵ_{Nd} and are useful on million year timescales, but are not applicable to high-resolution studies (e.g., Frank, 2002 and references therein). Dispersed Fe-Mn oxyhydroxide coatings on marine sediments have been exploited for this purpose (Rutberg and others, 2000; Piotrowski and others, 2005; Gutjahr and others, 2008; Piotrowski and others, 2009). However, they are subject to contamination from volcanogenic detritus (if in close proximity to volcanic islands) or pre-formed oxides (near continental margins) (e.g., Frank, 2002; Bayon and others, 2004; Piotrowski and others, 2005). Furthermore, these archives provide no records of surface-ocean circulation.

In contrast to ϵ_{Nd} sedimentary archives, foraminiferal ϵ_{Nd} offers the potential to conduct higher-resolution studies and to reconstruct surface-ocean circulation. Pioneering work on ϵ_{Nd} in core-top and plankton-tow foraminifera, along with seawater, bulk sediment, and oxide leaches, has shown that cleaned planktic foraminifera accurately record the ϵ_{Nd} of surface water, and are not measurably contaminated by Fe-Mn coatings or any post-depositional overprint (e.g., Vance and Burton, 1999; Burton and Vance, 2000; Vance and others, 2004). ϵ_{Nd} in planktic foraminifera may be complicated by vertically migrating species, such as during addition of a gametogenic crust (e.g., *Globigerinoides sacculifer*), if they migrate between water masses. The impact of vertical migration may be tested for by comparing ϵ_{Nd} of *Gln. sacculifer* with that of a shallow, mixed-layer species that does not produce a gametogenic crust (e.g., *Gln. ruber*). Therefore, accurate species-specific measurements of ϵ_{Nd} in planktic foraminifera could allow for the reconstruction of the shallow surface circulation and thermocline structure (Vance and others, 2004).

As a paleocirculation proxy, Nd isotopes hold a major advantage over nutrient-like proxies ($\delta^{13}\text{C}$, Cd, Ba, and Zn) in that ϵ_{Nd} is not fractionated biologically or during most low-temperature processes at Earth's surface; therefore, it is not subject to vital effects, nor should it respond to changes in the mechanism or rate of Nd uptake. Any variability in foraminiferal ϵ_{Nd} (including interspecies variability) reflects only the seawater variability in which the foraminifera calcify. Another advantage of ϵ_{Nd} over the nutrient-like proxies is its usefulness over much longer timescales; ϵ_{Nd} can be used as a water mass tracer as long as the assumption of constant end-member compositions is valid. The ϵ_{Nd} of end-member water masses is set by the surrounding continents at the sites of deepwater formation. Therefore, the end-member ϵ_{Nd} could be altered by a significant change in location of deepwater formation or composition of adjacent exposed continental crust.

Although foraminiferal ϵ_{Nd} appears to reliably record changes in water-mass mixing (e.g., Vance and Burton, 1999; Vance and others, 2004; Klevenz, 2008), ongoing studies will determine its ultimate utility in paleoceanographic applications. Little is known about the behavior of dissolved Nd in the oceans or its incorporation into foraminiferal calcite. Seawater ϵ_{Nd} exhibits quasi-conservative behavior (Goldstein and Hemming, 2002 and references therein). Although open ocean ϵ_{Nd} varies only with water mass, the concentration of Nd is depleted in surface waters relative to the deep ocean due to scavenging from the particle-rich surface waters. Therefore, there is a discrepancy between the behavior of Nd concentration and its isotope ratios in the oceans. The mechanism of Nd incorporation into foraminiferal tests is entirely unknown, and the apparent distribution coefficient for Nd into foraminiferal calcite is orders of magnitude higher than expected (e.g., Vance and others, 2004). There is also debate about adequate cleaning techniques and the presence of an unidentified refractory contaminant phase, which may or may not affect ϵ_{Nd} . The use of ϵ_{Nd} in foraminifera is still being developed, and calibration studies of planktic and benthic foraminiferal ϵ_{Nd} to seawater ϵ_{Nd} should shed light on some of these issues (Vance and others, 2004).

CONCLUDING REMARKS

The three-dimensional variability of the modern ocean is relatively well constrained by direct measurements of myriad characteristics such as temperature, pH, and nutrient content (e.g., Levitus and Boyer, 1994). In contrast, reconstructing ocean conditions through geologic history relies on inferences based on geochemical proxy records. Isotope and trace element composition of calcareous foraminifera has provided much of the foundation for reconstructions of past ocean and climate conditions, paleocirculation patterns, the carbon cycle, paleoproductivity, and marine carbonate chemistry, and also has provided an important stratigraphic tool for correlation and chronologies.

In this paper, we reviewed the geochemical proxies that utilize foraminiferal carbonate tests, including common uses of the proxies for reconstructions through time— $\delta^{18}\text{O}$, $\delta^{13}\text{C}$, trace elements (Mg, Cd, Ba, Zn, B, U), $^{87}\text{Sr}/^{86}\text{Sr}$, $\delta^{26}\text{Mg}$, $\delta^{11}\text{B}$, and ϵ_{Nd} . Employing these proxies requires an understanding of a wide range of associated factors that ultimately determine the geochemical signatures recorded in the foraminiferal calcite; discussions of these issues were also covered here.

Several of the geochemical proxies (B/Ca, U/Ca, $\delta^{26}\text{Mg}$, $\delta^{11}\text{B}$, ϵ_{Nd}) are relatively new and are in the development stage for use with foraminifera. As these proxies become better understood and analytical techniques are fine-tuned, they will provide important additions to our geochemical proxy toolbox with the potential to broaden the scope of reconstructions in the near future.

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