دونون B Sea-Surface Temperature from Coral Skeletal Strontium/Calcium Ratios

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Seasonal records of tropical sea-surface temperature (SST) over the past 10⁵ years can be recovered from high-precision measurements of coral strontium/calcium ratios with the use of thermal ionization mass spectrometry. The temperature dependence of these ratios was calibrated with corals collected at SST recording stations and by ¹⁸O/¹⁶O thermometry. The results suggest that mean monthly SST may be determined with an apparent accuracy of better than 0.5°C. Measurements on a fossil coral indicate that 10,200 years ago mean annual SSTs near Vanuatu in the southwestern Pacific Ocean were about 5°C colder than today and that seasonal variations in SST were larger. These data suggest that tropical climate zones were compressed toward the equator during deglaciation.

Retrieval of climatic data from the geologic record is important for validating climate models and for understanding the causes of past climate fluctuations. In most climate models, ocean temperature is a crucial parameter because of its linkage with other climate variables such as atmospheric moisture content and temperature, the extent of cloud cover and atmospheric albedo, or the patterns of oceanic and atmospheric circulation. Earlier efforts to recover past ocean

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temperatures from the geologic record have generally yielded inconsistent results or have proven to be difficult to apply. For example, attempts to recover ocean temperature from the O isotopic composition of biogenic marine carbonate sediments (1, 2)have been hampered because we do not know the history of fluctuations in the ocean water ¹⁸O/¹⁶O ratio, which varies as a function of the volume of the planetary ice caps, or, in the case of sea-surface water, can be modified by rainfall or evaporation effects. Efforts have also been made to reconstruct past ocean temperatures from the records of foraminiferal shell assemblages preserved in deep-sea sediments (3) based on knowledge of the temperature controls on modern foraminiferal assemblages (4). Such studies suggest that tropical SSTs were 0° to 2°C less than at present during the last glacial maximum (LGM) ~18,000 years ago (3). These findings, however, are at variance with estimates of SST based on the elevation of past mountain snow lines,

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which indicate that tropical SSTs were 3° to 6.5° C lower than at present during the LGM (5–7).

One promising method of recovering past SST records involves measuring the Sr/Ca ratios in corals (8-10). Scleractinian corals secrete skeletons composed of aragonite (CaCO₃), which incorporates both Sr and Ca into its structure. The ratio of incorporation of Sr to Ca is controlled by two factors: the Sr/Ca activity ratio of the ocean water, and the Sr/Ca distribution coefficient between aragonite and seawater (11, 12). This latter factor depends on the temperature of the seawater in which the coral grew but is only a weak function of the chemical composition of the seawater. Because of the long residence times of Sr and Ca in the oceans, it is probable that the seawater Sr/Ca ratio has remained essentially constant over time scales of about 105 years. Thus, the Sr/Ca ratio of corals is a potential monitor of ocean temperature on these time scales. Earlier work confirmed that coral Sr/Ca ratios reflect SST to a precision of approximately $\pm 3^{\circ}$ C (2 σ) (Fig. 1) (8-10, 13). However, this precision is about the same as the typical seasonal SST range observed in the tropics where scleractinian corals are found. Longer term variations in global average SST, such as the difference between mean SST for modern times and that at the LGM, may be even smaller than this, perhaps less than 2°C (3). Thus, this resolution is inadequate for evaluating seasonal or long-term ocean temperature variability.

In this paper we test the hypothesis that significantly better estimates of SST may be obtained from coral Sr/Ca ratios by improving the measurement precision of this elemental ratio. We first show that considerable improvement in the precision of Sr/Ca measurements can be obtained through use of thermal ionization mass spectrometry (TIMS). Next we assess whether improved analytical precision of this ratio translates into improved accuracy in SST. To do this, we correlated coral Sr/Ca ratios with ocean temperature by measuring ¹⁸O/¹⁶O ratios on the same coral samples, using the knowledge that coral ¹⁸O/¹⁶O ratios are a proxy for ocean temperature. Because our corals were collected from ocean island sites adjacent to SST recording stations, the accuracy of the Sr/Ca temperatures was independently assessed by comparison with actual SST measurements. Finally, we discuss the application of this method to recovery of SST from corals that grew during the last deglaciation.

Improved precision in the measurement of coral Sr/Ca ratios was made possible through use of isotope dilution with a triple ${}^{42}Ca {}^{44}Ca {}^{84}Sr$ spike in combination with TIMS (14). Both Ca and Sr were loaded

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together onto a double Re filament without prior chemical separation from the coral matrix. For a typical sample size of about 5 to 10 µg of Ca and 100 to 200 ng of Sr, we found that the average uncertainty (15) on the coral Sr/Ca ratio measured this way was $\pm 0.02\%$ (2 σ) for an individual analysis. On the basis of four duplicate analyses of coral standard Tan-E, the external reproducibility (15) on the Sr/Ca ratio for typically sized samples is $\pm 0.03\%$ (2 SD). This precision is roughly two orders of magnitude better than obtained by earlier methods (Fig. 1) and translates into a possible analytical precision in temperature of ±0.05°C (2σ) based on the Sr/Ca and temperature relation shown by Smith et al. (9). With this greater degree of analytical precision, the remaining issue becomes the accuracy with which the coral Sr/Ca ratio reflects ocean temperature.

For the coral species Porites lobata, we recalibrated the Sr/Ca thermometer using the relation between coral Sr/Ca and ¹⁸O/ ¹⁶O ratios, using ¹⁸O/¹⁶O ratios as a proxy for SST (16, 17). We collected annually banded corals for calibration from ocean islands in the western and central Pacific. Corals were recovered from barrier reefs that were well washed with seawater from the open ocean to ensure that they grew under conditions representative of the open ocean and not of closed lagoons, which could have anomalous seawater Sr/Ca or ¹⁸O/¹⁶O ratios. Fluctuations in salinity were insignificant at these sites as determined from daily salinity records, confirming that these reefs were exposed to open ocean conditions. Corals from Noumea, New

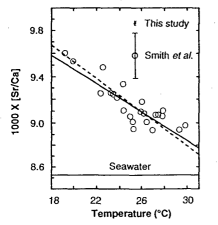


Fig. 1. Plot of Sr/Ca ratios in scleractinian corals against the temperature of the seawater from which they grew. The work of Smith *et al.* (9) is shown by open circles and the dashed line. Results of our work (solid line) indicate that some improvement in this correlation is possible, although the slope and intercept of the resultant linear relation (solid line) are similar to those determined in (9). The error bar for Smith *et al.* is the average (2σ) error The error bar for our work is ten times the average (2σ) error

Caledonia, and Papeete, Tahiti, were sampled parallel to the growth direction in time series with temporal resolution of approximately monthly intervals. We obtained both δ^{18} O (18) and Sr/Ca analyses from the same samples of coral by first extracting the oxygen (19) at 80°C by phosphoric acid dissolution in a quartz extraction line. The Sr and Ca concentrations were then obtained from TIMS analysis of the remaining residue.

For the New Caledonia site, both δ^{18} O and the Sr/Ca ratio vary cyclically with an annual periodicity and are strongly correlated (Fig. 2), yielding a temperature calibration (20) of the coral Sr/Ca ratio for the species *P. lobata*:

$$T (^{\circ}C) = 171.6 - 16013 \times (Sr/Ca)_{atomic}$$
(1)

Within error, this relation is similar to that derived by Smith *et al.* (9) for the genus *Porites* but has a slightly different slope and intercept, as well as improved precision (Fig. 1).

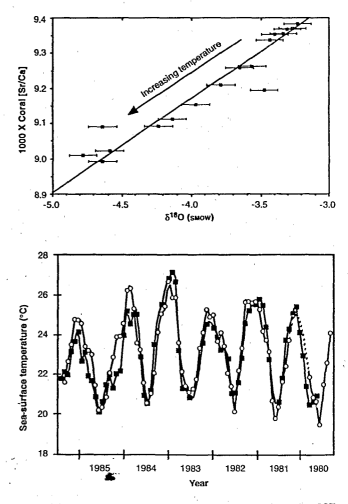
Using Eq. 1, we compared the calculated Sr/Ca temperatures with the actual SST

Fig. 2. Sr/Ca ratios versus δ^{18} O values from the same samples of coral. Horizontal error bars are the average (20) external analytical error on the 18O/16O ratio. Vertical error bars (height of the open square) are the average (2o) external analytical errors in the Sr/Ca ratio. External errors include uncertainties from the sample extraction process and are based on repeated analysis of carbonate standards (19).

Fig. 3. Comparison between actual SST (open circles) recorded at the temperature-recording station at Amedee Lighthouse, Noumea, New Caledonia, and Sr/Ca temperatures (filled squares) for samples of the coral species P. lobata, using Eq. 1. Actual SST data points represent 30-day averages of daily SST measurements. The coral sample size used for Sr/Ca analyses represents approximately monthly intervals of growth. As such, coral Sr/Ca temperatures represent the average ocean conditions over

recorded at New Caledonia and Tahiti-. (Figs. 3 and 4) as a test of the accuracy of the Sr/Ca thermometer. Matching of the Sr/Ca temperature and SST records was enabled by approximately superimposing the annual maxima in each record and assuming a constant coral growth rate throughout the year. For the Tahiti coral, the average difference between Sr/Ca temperature and actual 10-day average SST is 0.34°C. Thus, the resulting correspondence between the calibrated Sr/Ca temperatures and actual SST indicates that the Sr/Ca thermometer is both accurate and robust for at least these two quite different sites. These data suggest that coral Sr/Ca ratios are indeed a proxy for SST.

In order to obtain paleotemperatures from Sr/Ca ratios of ancient corals, we must assume that variations in the ocean Sr/Ca ratio have been small over the time period of interest. Because of the long ocean residence times of Sr and Ca [5.1 × 10° years for Sr and 1.1 × 10° years for Ca (21)] relative to ocean mixing times (~1000 years), temporal and geographic variations of Sr/Ca ratios in the open ocean are small



these intervals. The dashed portion of the temperature record represents a period for which SST records were not available.

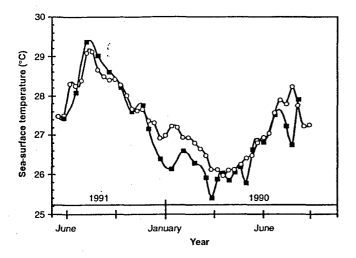


Fig. 4. Comparison between actual SST (open circles) recorded at the temperature recording station at Papeete, Tahiti, and Sr/Ca temperatures (filled squares) for samples of the coral species P. lobata, using Eq. 1. Actual SST data points represent 10-day averages of daily SST. Coral Sr/Ca temperatures represent the average ocean conditions over approximately 10-day intervals. The average difference between Sr/Ca temperature and the actual 10-day average SST is 0.34°C.

(8, 22), although the modern oceans do appear to show variations outside of analytical uncertainty (23, 24). Consequently, for time scales on the order of 10⁵ years. only small shifts in the ocean Sr/Ca ratio would generally be expected as a result of variations in river-water flux. Perturbations in the calcite compensation depth (CCD) (25) associated with Quaternary climate shifts could potentially cause more rapid changes in ocean Sr/Ca ratios. If such changes resulted in a 10% change in the area of sea floor suitable for calcite deposition, this would result in a maximum shift in the seawater Sr/Ca ratio of about 0.45% between glacial and interglacial periods, provided such CCD excursions did not cause wholesale dissolution of calcite sediments already on the sea floor. Errors in temperature introduced by uncertainties in the oceanic Sr/Ca ratio of this magnitude are less than about 0.5°C, which is about the same as the uncertainty of the Sr/Ca thermometer. In principle, this permits the recovery of paleo-SST records from corals that grew in the last 10⁵ years.

Fossil P. lobata corals that grew during the last deglaciation have been recovered from

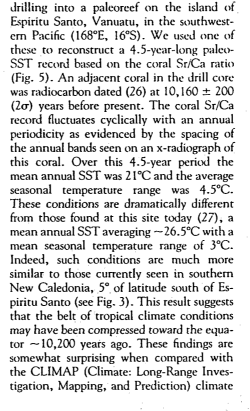
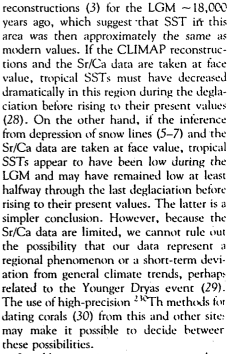


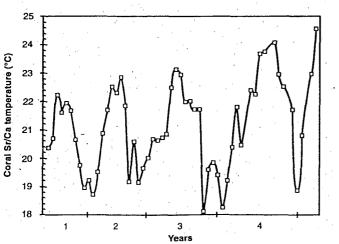
Fig. 5. A 4.5-year temperature record obtained from a fossil P. lobata recovered from drilling into a paleoreef on the island of Espiritu Santo (Vanuatu) in the southwestern Pacific (168°E) 16°S). This coral has been assigned an age of 10,160 \pm 200 (2 σ) years before present (26).



In addition to its use as an ocean thermometer, the correlation observed betweer coral Sr/Ca ratios and $\delta^{18}\text{O}$ may also make it possible to determine sea-surface δ^{18} O by removal of the temperature component o the coral δ^{18} O signal. Maps of sea-surface δ^{18} O generated in this way might be used to estimate variations in the volume of the planetary ice caps or to generate maps o sea-surface salinity (31, 32). The latte might in principle be used to recover pas patterns of rainfall and evaporation ove the tropical oceans. Finally, heavy rainfal events can result in short-term (week-lon: to month-long) depletions in surface-oceau water δ^{18} O. Such events may cause devia tions from the observed linear relation be tween Sr/Ca ratio and δ^{18} O. If so, it may be possible to assess the frequency of pas tropical storm events from time-series anal ysis of high-resolution coral Sr/Ca ratio and δ^{18} O records. Possible nonequilibrium fractionation of the O isotopes (17) it corals, however, may limit recovery of in formation about such rainfall or ice-volum variations.

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- A distribution coefficient is usually defined by an 11 equation of the form: $D_{St} = (Sr/Ca)_{mineral}$ $(Sr/Ca)_{flund}$, where $(Sr/Ca)_{flund}$ and $(Sr/Ca)_{flund}$ are the molar concentration ratios of these elements in the mineral and fluid, respectively.
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- We calculated 2σ error estimates from the mean square of the residuals of a linear regression through the data presented in (9) for the coral genus Porites.
- 14. Sr and Ca concentrations were measured by isotope dilution on a Finnigan MAT 262 solid source mass spectrometer, using static Faraday collection. An exponential mass fractionation (33) correction for ⁹⁴Sr/⁹⁶Sr and ⁴⁴Ca/⁴²Ca was used, correcting to an assumed (34) natural ⁸⁶Sr/⁸⁸Sr = 0 1194 and an assumed (33) natural ⁴²Ca/⁴⁴Ca = 0.31221 Spike isotopic composition corrections were based on gravimetric determinations for Ca, and on non-fractionation-corrected TIMS determinations of the isotopic composition of the Sr spike Results of 19 runs of Minnesota Isotope Laboratory standard ISC-1 Ca yielded an average 40Ca/42Ca = 151.030 ± 19 (2σ), whereas results of 10 runs of National Bureau of Standards (NBS) standard 987 Sr yielded an average ⁸⁷Sr/⁸⁶Sr = 0 710239 ± 10 (2o).
- 15. Each Sr or Ca analysis constitutes between 50 and 100 individual mass spectrometric scans (isotopic ratios). A standard error (SE) of the mean is calculated for each analysis; these SEs are in turn propagated to determine the SE for the Sr/Ca ratio. "Average uncertainty" in the sense used here refers to the average (2σ) SE for all of the Sr/Ca analyses reported in this paper. "External reproducibility" as it is used here refers to two standard deviations of the population, where the population constitutes the mean values of several different determinations of the Sr/Ca ratio of a standard.

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- δ^{18} O is the normalized deviation, in parts per thousand, of the sample 18 O/ 16 O relative to stan-18 dard mean ocean water (SMOW) (35).
- We performed δ^{18} O analyses on a Finnigan MAT 19. delta-E mass spectrometer. Average internal precision on the $\delta^{18}O$ analyses is reported at 0.07 per mil (2 σ) on a typical 5-mg sample of coral. Five runs of standard NBS 18 yielded an average δ^{18} O $(SMOW) = 7.18 \pm 0.06$ per mil (2 σ), whereas results of nine runs of standard NBS 19 yielded somewhat poorer external reproducibility, with an average δ^{18} O (SMOW) = 28.75 ± 0.10 per mil (20)
- 20 The Sr/Ca temperature calibration made use of the $\delta^{18}O$ versus temperature calibration for the coral genus Porites generated by McConnaughey (17) This equation may be expressed as: $\delta^{18}O_c - \delta^{18}O_w = 0.594 - 0.209 T$ (°C), where $\delta^{18}O_c$ and $\delta^{18}O_{\omega}$ are the O isotopic compositions of the coral and seawater, respectively. The value of $\delta^{18}O_{\omega}$ for one seawater sample from the Amedee Lighthouse, Noumea, New Caledonia, calibration site was +0 52 \pm 0.06 per mil (2 σ). Average seawater $\delta^{18}\text{O}$ was assumed to be +0.6 per mil for this site based on published compilations:
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