A stable isotope study of organic cycling and the ecology of an anchialine cave ecosystem

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ABSTRACT. Stable carbon and nitrogen isotope data, complemented with other geochemical parameters, were used to identify the sources of organic matter that support the food web of an anchialine cave ecosystem in the northeastern Yucatan Peninsula, Mexico. Anchialine caves, common along tropical karstic and volcanic coastlines, are completely or partially inundated by highly stratified layers of fresh and marine waters. Stable isotope data from the cave fauna, the particulate organic matter (POM) from the cenote pool and from the cave, the forest soil and the cave sediments indicated that at least 3 sources of nutritive organics could support the anchialine food web. These sources were: (1) soil from the overlying forest; (2) freshwater algae from adjoining open water pools; and (3) chemoheterotrophic nitrifying bacteria living in the cave. Production of nitrate and a decrease in $O_2$ along the halocline provided geochemical evidence of nitrification. Stable nitrogen isotope data defined 2 to 2.5 trophic levels in the food web. Furthermore, it was found that troglobitic (cave-limited) species residing in the water column are capable of preferentially feeding on specific organic reservoirs. This study presents the first extensive description of the ecological and biogeochemical relationships of the anchialine cave ecosystem.

KEY WORDS: Anchialine Organic cycling Nitrification Niche partitioning Yucatan Peninsula Cave

INTRODUCTION

In many tropical karstic regions, coastal groundwater consists of a fresh to slightly brackish lens overlying seawater that intrudes from the coast (Iliffe 1992a). Within these 'underground estuaries' in areas such as the Yucatan Peninsula of Mexico, vast cave networks, technically termed anchialine, meander through the Pleistocene bedrock (Back et al. 1979). Mixing of the water masses at the halocline produces a corrosive solution capable of dissolving additional limestone (Back et al. 1986) (Fig. 1a). The dissolution of limestone under such conditions has created extensive cave systems that serve as conduits for groundwater flow and provide habitat for a unique community of troglobitic (cave-limited) organisms.

The recognition of anchialine caves as a significant biological habitat has only occurred over the past 2 decades. Developments in diving technology have made it possible for trained cave-diving scientists to investigate the biology of the anchialine habitat. Consequently, our knowledge of the taxonomy and biogeography of troglobitic organisms in anchialine caves has increased tremendously (Yager 1981, Iliffe 1983, Iliffe et al. 1984b, Bowman et al. 1985, Hart et al. 1985, Manning et al. 1986). These discoveries have provided the foundation for the investigation of anchialine cave ecology and biogeochemistry: the focus of the present study.

The world's most extensive anchialine cave networks are found in the Yucatan Peninsula of Mexico, a flat limestone plain completely devoid of surface rivers and streams. Due to this absence of surface streams, the underground aquifer is the primary reservoir for domestic, agricultural and industrial needs. This water is accessible principally through pools called cenotes.
Fig. 1 (a) Schematic diagram of a typical cenote-cave system. Cave passages are primarily formed by mixing corrosion at the depth of the halocline, while cenotes result from the collapse of the ceiling. The caves serve as conduits for the groundwater flowing to the coast. Typically, water emerges from a spring on one side of the cenote pool and returns below ground through a siphon on the opposite side. (b) Map of the Yucatan Peninsula (Mexico) showing the location of Mayan Blue Cenote relative to several other sites.

that pockmark the karst landscape of the peninsula and wells bored into the limestone caprock (Buck & Hanshaw 1974).

At the present time, this coastal aquifer is pristine, but as the population and economy continue to grow, the groundwater quality will likely be compromised. Injection of wastewater or sewage into the groundwater is a primary concern. Dissolved oxygen (DO) levels in groundwater are inherently low due to the absence of photosynthesis in the aphytic subterranean environment and isolation from atmospheric exchange (Iliffe et al. 1984a). Increasing the input of organic material stimulates bacterial productivity and respiration, further depleting DO and producing zones of anoxia spreading out from the source of contamination. For example, Iliffe et al. (1984a) investigated Government Quarry Cave, an anchialine cave in Bermuda, to assess the damage caused by such organic pollution. Domestic wastes dumped into the cave pool caused widespread anoxia, eradication of the macrofauna and high accumulations of nutrients. According to their calculations, doubling the natural input of organic matter into such caves can result in anoxic conditions. Considering the small quantities of organic material naturally present in anchialine caves (Fichez 1991), doubling the size of the organic matter reservoir is not difficult to do.

Most aquatic cave ecosystems are regarded as detritus-based (Dickson 1975), but chemoautotrophic primary production has been documented in several caves. The ecosystem in Movile Cave, a thermomineral cave in Romania, is supported entirely by chemoautotrophic sulfide-oxidizing bacteria floating at the water's surface in air pockets and coating the walls of the underwater cave (Sarbu et al. 1996). Dissolved H₂S concentrations supporting this production are exceptionally high, ranging between 500 and 1300 μM. Additionally, several submarine caves in Italy receive input from warm sulfurous springs and have associated chemoautotrophic bacterial mats (Southward et al. 1996). Stable carbon isotope ratios of animal tissue from that study indicated that the bacterial contribution to the food chain varies with species from 0 to 100%. Because anchialine caves are not typically associated with geothermal sources, we inferred that their ecology would be governed by different biogeochemical principles.

We hypothesized that the major reservoirs of organic matter supporting the anchialine ecosystem would be: (1) algal detritus transported by groundwater flow from the cenotes; (2) soil particulates percolating from the tropical forest soil through the cracks and fissures of the limestone bedrock into the caves; and (3) coastal-borne particulate organic matter (POM) transported into the cave systems with tidally exchanged seawater. We also expected that algal and soil detritus would support the community above the halocline, while the coastal-borne POM would nourish the community below the halocline. We recognized the potential for chemoautotrophic production of organic material by methanogenesis, hydrogen sulfide or regenerated ammonia, but assumed it would be insignificant in terms of its contribution to the nutritive organic pool.

MATERIALS AND METHODS

Study site and collection procedures. The study site was in and around Mayan Blue Cenote, a karst window in the Systema Naranjal anchialine network,
Mexico (Fig. 1b). The orientation of the 18.47 km long Systema Naranjal is perpendicular to and about 5 km inland from the Caribbean coast (J. Coke pers. comm.). Samples were collected during expeditions in July 1993, March and August 1994, and March 1995. The samples collected included fauna, flora, sediments, POM, bacteria and water from the caves and open water cenote pools, as well as vegetation and soil from the tropical forest. Fauna were either collected individually in plastic vials or with a diver-towed 0.3 m diameter plankton net with a 93 μm mesh. The fauna were collected from 5 different passages accessed from Mayan Blue Cenote (Pohlman 1995). Sediments, soil, bacteria and flora were collected in sterile Whirl-Pak plastic sacks, while POM samples were collected with a diver-operated Wilden M.025 pneumatic pump onto precombusted Whatman GF/F filters. Water samples were collected in 50 ml Nalgene bottles that were acid cleaned and filled with distilled water before the dive. Water samples collected for δ13C-DIC (dissolved inorganic carbon) and DIC concentration analyses were fixed with a saturated HgCl2 solution immediately after the dive. Scientifically trained cave divers, following standards set by the National Speleological Society-Cave Diving Section (NSS-CDS), collected all samples.

Troglobitic fauna. A diverse and unique assemblage of troglobitic fauna have been identified from the caves surrounding Mayan Blue Cenote. The freshwater lens above the halocline is habitat for 1 species of fish and 6 species of Crustacea, while the seawater portion of the cave supports 4 species of Crustacea (Kornicker & Iliffe 1989, Iliffe 1992b, L. Kornicker pers. comm.). The benthic fauna includes 2 species of Isopoda (Iliffe 1992b). However, because several of these species are either rare and/or small they were not collected during this study. The species collected from Mayan Blue are listed in Table 2 (see Results) with habitat information and stable isotope data. The fauna were identified to the most specific taxonomic level possible—usually species.

Stable isotopes. Fauna, flora, POM, sediments and soils collected for stable isotope analysis were stored either on dry ice or in an L-C Lab Line drying oven held between 60 and 70°C. In the laboratory, these samples were acidified to remove inorganic carbon. The samples were analyzed by a modified Dumas combustion method that converts organic carbon and nitrogen to CO2 and N2 for mass spectral analysis (Macko 1981). The nitrogen gas was analyzed on a Nuclide 3-60-RMS isotope ratio mass spectrometer and the CO2 gas by a Finnigan MAT 251 or 252 stable isotope mass spectrometer.

For δ13DIC analysis, the CO2 was removed from solution with 85% o-phosphoric acid and separated on a Varian 3400 Gas Chromatograph (GC). The GC was coupled to a combustion oven and a Finnigan MAT 252 stable isotope mass spectrometer, where the CO2 was analyzed.

Stable carbon and nitrogen isotope ratios are reported according to the standard formula:

$$\delta X = \left(\frac{R_{\text{sample}}}{R_{\text{standard}}} - 1\right) \times 10^3 \%$$

where δX is either δ13C or δ15N, and R is either 13C/12C or 15N/14N. The standards for carbon and nitrogen are, respectively, PeeDee Belemnite limestone fossil and atmospheric dinitrogen gas. The precision for both instruments was ±0.2%.

Dissolved inorganic carbon. DIC concentrations were measured coulometrically (Dickson & Goyet 1991). Triplicate samples (10 ml) were injected into a stripping chamber containing 0.1 N H2SO4 that converted the DIC to CO2. The evolved CO2 entered a coulometer cell (UIC Inc., Model 5011) where it was titrated to a constant endpoint. The integrated titration current measured by the coulometer was converted to μM CO2. Precision of standards prepared from NaCO3 was ±2.8 μM.

Dissolved inorganic nitrogen (DIN), methane and total dissolved sulfide (TDS). DIN (NO3−, NO2− and NH4+) was measured on a Technicon AA-II Autoanalyzer (Koroleff 1970, Aminot & Kerovell 1982). Methane was measured by the headspace phase equilibrium method with a Shimadzu gas chromatograph equipped with a flame ionization detector (McAuliffe 1971). TDS was determined spectrophotometrically in the field against a standard curve generated by the Cline (1969) method. Sensitivity range for the curve was between 2 and 40 μM. Because of the lack of laboratory facilities in the field, the standard curve was generated 1 wk prior to the field work.

Organic carbon and nitrogen analysis. Sediments and algae collected from the pool and cave were analyzed for percent organic carbon and nitrogen. Sample preparation was conducted according to the methods described by Nieuwenhuize et al. (1994) and the analysis carried out with a Carlo Erba NA-1500 elemental analyzer. This technique is particularly useful for analyzing carbonate-rich sediments like those found in the caves.

Multiprobe parameters. Salinity, temperature, pH and DO were measured with a Hydrolab Datsasdone III multiprobe. Accuracy was ±0.2 g l−1 for the salinity, ±0.15°C for the temperature, ±0.02 units for pH and ±0.2 mg l−1 for DO. The probe was equipped with an independent battery pack and data logger which made it possible to transport the probe into the underwater cave for profiling. The unit was carried by the first diver with the probes leading to avoid measuring an artificially mixed water column.
Biogeochemical properties of the water column

Constituents of the water column affected by biogeochemical activity also showed dramatic variations in the halocline. DIC concentrations were much higher in the brackish region of the water column with a maximum of 8.01 mM near the cave ceiling at a depth of 10 m (Fig. 4a). Below the halocline, DIC varied between 2.16 and 2.70 mM. The stable carbon isotope values of DIC (Δ13C) also reflected a mixing of 2 distinct water types as Δ13C values as negative as -15.0% near the cave ceiling increased to -3.0% at the cave floor (Fig. 4b). The most dramatic changes of Δ13C occurred in the halocline.

DIN concentrations were similar above and below the halocline, but displayed unexpected behavior within the halocline (Fig. 5). Ammonium (NH₄⁺) concentrations were consistently low, ranging between 0 and 0.02 μM. Nitrite (NO₂⁻) concentrations ranged between 0.02 and 0.07 μM above and below the halocline, but displayed a 0.70 μM peak within it. Nitrate (NO₃⁻) was most abundant, varying between 2.92 and 5.79 μM above and below the halocline, but showed a pronounced peak increasing from 6.32 to 18.64 μM in the halocline. Although this is a 1 point peak, a similar nitrate peak of 23 μM measured by Stöessell et al. (1989) at the depth of the halocline corroborates the 18.64 μM nitrate peak measured during this study.

Near hypoxic conditions existed throughout the water column with dissolved oxygen (DO) values ranging between 1.4 and 1.7 mg l⁻¹ in water masses above and below the halocline (Fig. 2c). In the mixing zone of

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**RESULTS**

**Physical properties of the water column**

Temperature and salinity profiles from a Mayan Blue cave passage illustrate the sharp division between the brackish and seawater components of the water column (Fig. 2). Over a 3 m vertical range the salinity changed from 3.2 g l⁻¹ to a nearly full seawater concentration of 34.3 g l⁻¹. The temperature of the 2 water masses differed by 0.8°C, with the deeper seawater being slightly warmer at 26°C. Non-conservative mixing between temperature and salinity indicated that water has cooled slightly since entering the groundwater (Fig. 3a).

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**Fig. 2.** Vertical profiles of (a) temperature, (b) salinity, (c) dissolved oxygen (DO) and (d) pH from Mayan Blue cave system, March 1995.

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**Fig. 3.** Property salinity plots for (a) temperature, (b) dissolved oxygen (DO) and (c) pH from Mayan Blue cave system, March 1995. Dashed lines between salinity end members represents the conservative mixing distribution.
the halocline, however, there was a decrease in the amount of DO as concentrations dropped as low as 0.9 mg l⁻¹. DO exhibited non-conservative mixing through the water column with depressed concentrations in the halocline (Fig. 3b).

The pH of the brackish portion of the cave water column was nearly constant at 6.9. The pH in the mixing zone dropped as low as 6.6 and then increased to a typical seawater value of 7.2 below the halocline (Fig. 2d). When salinity was plotted against pH, it was evident that mixing was non-conservative due to hydrogen ion production in the halocline (Fig. 3c).

Methane and TDS in the cave waters were low. Two samples collected for methane in the halocline were 72 and 180 nM, while TDS was below the detection limit of 2 μM throughout the water column.

**Organic carbon composition of sediments**

The percentage of organic carbon in the sediments decreased dramatically with distance into the cave. At the cave entrance, where epilithic algae and vegetative detritus covered rock surfaces and the bottom of the pool, the sediments were rich with organic carbon. The percent organic carbon at the cave entrance ranged from 1.9 to 24.4% (n = 4). In the twilight zone (defined as <60 m from the entrance), however, the amount of organic carbon in the sediments dropped dramatically. Organic carbon in these sediments ranged between 0.1 and 1.9% (n = 8). The sediments in the dark portion of the cave (defined as >60 m from the cave entrance) where the troglobites reside contained even less organic carbon. In this area, organic carbon ranged from 0.1 to 1.9% (n = 12). Sediments in the dark portion of the cave consisted almost entirely of calcium carbonate.

**Stable carbon and nitrogen isotopes of the organic reservoirs**

**Forest.** Fresh leaves collected from the forest floor had mean δ¹³C and δ¹⁵N values of −27.8 ± 0.9 (n = 8) and +2.2 ± 0.9 (n = 8), respectively (Table 1). These values are typical of C-3 plant tissues (Boutton 1991, Nadelhoffer & Fry 1994) and, as would be expected, were nearly identical to the stable isotope values for

<table>
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<th>Location</th>
<th>Source</th>
<th>δ¹³C (‰)</th>
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<th>δ¹⁵N (‰)</th>
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<tr>
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<td>−24.1 ± 6.0</td>
<td>10</td>
<td>7.8 ± 0.6</td>
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the soil. δ{}^{13}C and δ{}^{15}N of the soil collected from the forest floor around the Mayan Blue study site averaged -27.8 ± 0.4 (n = 8) and +2.2 ± 0.8 (n = 8), respectively. The soil samples were separated into different size fractions (<125 μm and 125 to 500 μm) to examine possible isotopic differences between the size classes, but none were found (t-test, p < 0.01).

Pool particulate organic matter (PPOM). The living and non-living suspended material from the cenote that enters the cave is collectively referred to as PPOM to represent a single surrogate product of the cenote pool. PPOM had δ{}^{13}C and δ{}^{15}N values of -35.3 ± 2.0 (n = 4) and +6.7 ± 0.3 (n = 4), respectively (Table 1). Similar δ{}^{13}C values for POM have been reported from aquatic systems where the vegetation and phytoplankton assimilate isotopically light biogenic CO₂ (Rau 1978, Routton 1991).

Soil particulate organic matter (SPOM). Although water in the cenote pool flows into the cave, the δ{}^{13}C of the cave POM was drastically different from that found in the pool. Cave POM had a δ{}^{13}C of -24.1 ± 2.0 (n = 7) through the entire water column (Table 1). This material is classified as soil particulate organic matter (SPOM) because the forest soil, which has a similar δ{}^{13}C of -27.8, is most likely the primary source for SPOM. The discrepancy of 3.7% between the two can be attributed to enrichment of 13C that occurs during soil degradation (Boutton 1991). As soil degrades, it also becomes enriched with the heavier 15N isotope (Nadelhoffer & Fry 1994). For that reason, the SPOM, with δ{}^{15}N of +8.1 ± 1.8 (n = 6), is heavier with respect to nitrogen than the soil organic matter (+2.2 ± 0.8).

Cave benthic organic matter (CBOM). Sedimentary organic matter from the cave, hereafter referred to as CBOM, had δ{}^{13}C and δ{}^{15}N values of -24.1 ± 6.0 (n = 10) and +7.8 ± 0.6 (n = 9), respectively (Table 1). Because these values are nearly identical to its source. SPOM, it is presumed that SPOM sinks to the sediment and becomes CBOM.

Troglobitic fauna. The carbon and nitrogen stable isotope values of the troglobitic fauna displayed a wide range of values (Table 2). Carbon stable isotope values ranged between -21.2 and -42.6‰, while the nitrogen stable isotope values ranged between +4.1 and +13.4‰.

**DISCUSSION**

Nitrification and carbon cycling in anchialine caves

Our initial hypothesis was that soil from the forest, particulate matter in the pool and particulate matter from the Caribbean Sea would support the anchialine ecosystem, but upon closer inspection we found that production of organic matter by chemoautotrophic bacteria could be a significant food component. The presence of the small nitrite peak and dramatic nitrate peak in the mixing layer that coincided with a DO minimum (Fig. 5) suggests that nitrification is occurring. Nitrification is a chemoautotrophic process whereby ammonium (NH₄⁺) is sequentially oxidized, first to nitrite (NO₂⁻) and then nitrate (NO₃⁻). Nitrification is a common microbial event in aphotic oceanic environ-

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<th>δ{}^{15}N</th>
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ments where DO is low (Gundersen 1966, Carlucci & McNally 1969). Both of these environmental conditions exist in anchialine cave systems. Because nitrate production is believed to be strictly biogenic (Ward 1986), the presence of nitrate in the mixing zone of the cave water column can only be explained by nitrification.

The role of nitrification in the ocean is important for nitrogen cycling, but has generally been considered unimportant as a food source. Unusually high rates of nitrification at intermediate salinities in the mixing region of the Mississippi River (USA) plume have been reported (Pakulski et al. 1995), but nitrification as a mechanism for carbon fixation is inefficient. The ratio of carbon fixed to ammonium oxidized varies from 0.07 to 0.18 (Feliatra & Bianchi 1993). Consequently, the contribution of nitrifying bacteria to the total organic pool of the Mississippi River plume is most likely small even though the rates of nitrification are high. However, in anchialine systems, where organic matter is in short supply, production considered insignificant in other ecosystems could prove substantial.

Chemoautotrophic nitrification in anchialine caves can be likened to 'old' production in oceanic systems where remineralized nitrogen is recycled by the phytoplankton. Excretion by troglobitic metazoans, transformation of organic matter by bacteria and ammonification by heterotrophic microbes are the likely sources of ammonia supporting the nitrifiers. The pulse of hydrogen ions at the halocline indicates that such heterotrophic activity occurred (Fig. 2d).

A model for carbon cycling that includes nitrification is presented in Fig. 6. The 2 unequivocal sources of organic matter available for the anchialine food web were SPOM and PPOM. The source of SPOM was forest soil that percolated into the system through the cracks and fissures of the limestone bedrock. Its presence was corroborated by the similarity of the stable carbon isotope signatures between the POM collected in the cave and the surface soils collected from the forest. PPOM must have entered the ecosystem as water from the cenote flowed into the cave. Although the PPOM isotope signature was not reflected in the cave POM, it was observed in some of the troglobitic species.

The similarity between the cave POM and the forest soil does not preclude the presence of PPOM as a food source in the cave. Although the SPOM overwhelmed the isotopic signature of the PPOM in the cave POM, it is entirely possible that the pelagic crustaceans selectively fed on the less abundant PPOM fraction. Del Giorgio & France (1996) proposed this as a mechanism to explain the differences in the isotopic signature of the zooplankton and bulk POM of numerous freshwater and marine environments. PPOM is primarily algae, which has a higher proportion of nitrogen than vascular plants. Thus, it makes sense that the fauna would preferentially consume the PPOM despite the fact that it is present in small quantities.

The unusually light δ¹³C of the fauna provides additional evidence that nitrifying bacteria may be present in the caves. Peterson et al. (1980) reported that nitrifying bacteria were −28 to −30‰ lighter than their carbon source. Given the −11 to −15‰ range of DIC measured in the mixing zone where the nitrate peak was observed (Fig. 4b), one would expect that their stable carbon isotope values of the nitrifying bacteria would range somewhere between −35 and −45‰. δ¹³C values in this range were measured in the troglobitic fauna. Individuals from the atyid shrimp species Typhlatya mitchelli (−42.4‰) and the thermosbaenacean species Tulumella unidentes (−42.6‰) were more negative than any of the POM samples we measured, and are more negative than what one would expect from algae in the cenote. The lightest δ¹³C POM value from the pool was −38.3‰. This value fits into the range of −33 to −38‰ that one would expect for pure algae utilizing the pool DIC (Peterson & Fry 1987). Recognizing that there is an enrichment in stable carbon isotope values of about 1‰ per trophic level transfer (Haines & Montague 2003), it is possible that the pelagic crustaceans selectively fed on the less abundant PPOM fraction.
1979), the values measured from individual T. mitchelli and T. unidens are impossible to obtain if the only food sources were SPOM and PPOM. These individuals must have been supplementing their diet with a 13C-depleted source of organic carbon.

The variability in stable carbon isotope values of the suspension-feeding, pelagic crustacean species may be explained by specialized feeding behavior Typhlatya mitchelli and Tulumella unidens employ specialized feeding mechanisms that should enable them to graze small algal cells and bacteria, particularly nitrifying bacteria which can be as large as 3 μm (Ward 1986). T. mitchelli has setal tufts on its first and second pereiopods that allow it to ‘sweep’ minute particles from the water (Fig. 7). The individual setae have an array of filaments and saw-like structures that could ensnare and entangle bacteria or other small algal particles. Likewise, T. unidens has highly setose maxillules that also should allow it to gather small particles, although less effectively than T. mitchelli (M. Wicksten pers. comm.). On the other hand, mysids use their thoracic endopods to collect larger food particles (Shram 1986). Assuming that the soil particles are larger than the bacteria and algae, the ability to select differently sized food particles that are isotopically distinct may explain the differences in the stable carbon isotope signatures of the 3 suspension-feeding crustacean species.

Methanotrophic and sulfide-oxidizing bacteria may also be significant food sources. Groundwater methanotrophic bacteria have extremely light δ13C, ranging between −55 to −90‰ when oxidizing biogenic methane (Grossman & Coffman 1989), while sulfide-oxidizing bacteria from a nearby sinkhole, Cenote Angelita, had a δ13C of −34.3‰. Assimilation of these 13C-depleted stable carbon isotope sources could help explain the faunal isotope ratios. Concentrations of both methane and sulfide in the Mayan Blue system were low (see ‘Results’), but low concentrations of these compounds can support chemosynthetic production by methanotrophic and sulfide-oxidizing bacteria (Dando et al. 1985, Ward & Kilpatrick 1990).

Contrary to our original hypothesis, coastal-borne POM appears to be a negligible source of organic food for the organisms in the Mayan Blue cave system. A POM sample collected in the seawater portion of that cave had a δ13C of −24.3‰. This value is nearly identical to the mean δ13C of POM from the upper cave waters (−24.7‰) and is within the range reported for marine waters (−22.0 to −24.4‰) by Williams & Gordon (1970). This similarity between the seawater POM and soil POM makes it difficult to surmise the origin of the POM in the seawater of the cave. However, due to the proximity of the overlying soil horizon and the 6 km distance of the Mayan Blue cave system from the coast, we believe the POM in the seawater portion of this cave is derived principally from the soil. We still believe cave systems closer to the coast, which have a more recent input of coastal seawater, will contain a larger reservoir of marine-derived particulates for the euhaline troglobitic community. However, in systems far from the coast like Systema Naranjal, the marine influence is slight. The dearth of organic matter in the cave seawater in caves around Mayan Blue accounts for the fact that near-microscopic ostracods are the largest consumers of particulates below the halocline.

**Trophic structure of the food web**

A dual isotope approach was used to resolve the relative contributions of the measured organic sources to each species and to define the trophic structure of the
food web (Fig. 8). Based on a 2 to 3\% increase in $\delta^{15}N$ per trophic level transfer between consumer and consumed (DeNiro & Epstein 1981), there appear to be 2 to 5 trophic levels in the food web. Typhlatya mitchelli, Tulumella unidens, Antronmysis cenotensis (a mysid) and isopods constitute the lower trophic level and show source feeding preferences that agree with those suggested by the stable carbon isotope data. $T.$ mitchelli and $T.$ unidens were 2.4 and 1.7\% heavier, respectively, than the PPOM with regard to nitrogen, while similar with regard to carbon. These relationships suggest a strong, but not complete, dependence on PPOM. Large standard deviations of the stable isotope data for the $T.$ mitchelli and $T.$ unidens populations (Table 2) attest to the flexibility of these species to feed on SPOM and nitrifying bacteria, along with PPOM. A. cenotensis is intermediate between PPOM and SPOM in terms of its stable carbon isotope values and is slightly heavier in nitrogen than some of the other pelagic crustaceans. Both trends suggest that the mysid $A.$ cenotensis relies more so on SPOM. The isopods occupy the benthos and thus are reflective of the CBOM (Fig. 6).

The upper trophic level members—amphipods, the remipede Speleonectes tulumensis, the shrimp Creaseria morleyi and the blind cave fish Ogilbea pearsei—are all predators and/or scavengers, so their high trophic ranking is expected. The $\delta^{15}N$ values of the amphipods and $O.$ pearsei are the most positive. Therefore, they are placed half a trophic level higher than $S.$ tulumensis and $C.$ morleyi since they are able to feed on members of both the same and lower trophic levels. Amphipods have been observed scavenging on the exuviae of other crustaceans (possibly even other amphipods), while $O.$ pearsei is the top predator. $S.$ tulumensis, a crustacean from the ancient class Remipedia, is the lone predator/scavenger in the seawater. It is presumed to have fed on ostracods and crustaceans that have passed through the halocline.

**Comparison with the ecology of other anchialine caves of the Caribbean**

Similarities in habitat and community structure with other anchialine caves in the Caribbean suggest that our observations may not be site specific. For instance, salinity, DO and temperature profiles reported from an anchialine cave in Cuba (Cueva de los Carboneros) by Yager (1994) resemble those from Mayan Blue and other anchialine caves we investigated in the Yucatan. As well, sites in the Bahamas, the Dominican Republic and Cuba share 4 common genera of Crustacea with those found in the Yucatan Peninsula (Holsinger 1989). Investigations in other cave systems and continued research in Systema Naranjal will provide a more complete understanding of the fundamental biogeochemical forces that drive anchialine cave ecology. The system model presented in this paper is certain to be altered and improved as important factors we have not yet considered are examined, and spatial and temporal patterns are more closely considered. Finally, the stability of anchialine cave ecosystems makes them suitable models for studying the dynamics of other oligotrophic ecosystems.

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