A New foodweb based on microbes in calcitic caves: The Cansiliella (Beetles) case in Northern Italy

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Abstract:

The troglobitic beetle, Cansiliella servadeii (Leptodirini), has specialized mouthparts modified for browsing and feeding under percolating water on moonmilk, a speleothem formation in Grotta della Foos, Italy. Results from analyses of stable isotopes of carbon and nitrogen suggest that it acquires and assimilates dissolved allochthonous organic carbon, inorganic nitrogen, and possibly phosphorus and other nutrients from the microbial fauna associated with moonmilk.

Keywords: Cansiliella servadeii, cave, moonmilk, stable isotopes, food web, bacteria, microorganisms

INTRODUCTION

In air-filled cave and karst habitats, troglobites and troglobiophiles rely on an influx of allochthonous (surface-derived) organic matter from dripwaters or sheeeflow on moist cave-wall surfaces. This habitat type is referred to as the cave hygropetric (Ske, 2004). Hydrological isolation for some cave hygropetric habitats may restrict the influx of organic matter, making a habitat nutrient-limited overall. Consequently, food limitation over extended periods of time has served as a major driver for evolutionary adaptation for troglobites (e.g., Ske, 2004). One highly evolved, troglobitic beetle, Cansiliella servadeii Paoletti 1980 (Coleoptera: Leptodirini, formerly Bathyscininae), has been found at a depth of ~450 m below the surface in Grotta della Foos, northeastern Italy.

The genus Cansiliella consists of the species C. servadeii, C. toniellai Paoletti 1972, and C. montiscenesi (Piva, 2000). Cansiliella, as well as other beetles from the genus Hadesia, and possibly the recently described genera Radziella, Velobitodromus, Croatodirus, Nauticella, Tartariella, and Kircheria, have peculiar, semi-aquatic feeding behavior morphological structures that differ greatly from the majority of other troglobitic Leptodirini (Jeannel, 1924; Remy, 1940; Paoletti, 1973;1980; Moldovan et al., 2004; Ske, 2004; Giachino & Vailati, 2006; Perrau & Pavicevic, 2008; Paoletti et al., 2009). The mouthparts of Cansiliella spp. are adapted for a semi-aquatic feeding behavior and show modified hoe-shaped mandibles and spoon-like galeae with fine, dense setae for filtering (e.g., Moldovan et al., 2004; Paoletti et al., 2009). The food sources for these beetles have not been thoroughly studied yet, although the beetles do not feed upon guano or cricket eggs, carrion, or particulate organic matter such as woody debris. They cannot be easily captured by guano, rotting meat, cheese, or other baits routinely used for troglobite collection in caves. Ske (2004) hypothesized that Cansiliella may have a microphagous feeding habit or filter fine particles of allochthonous organic material from water.

The C. servadeii beetles from Grotta della Foos are essentially found associated with moonmilk, a secondary carbonate speleothem, that forms from percolating waters containing little to no surface-derived organic matter (Figure 1) (Engel, unpublished results). Moonmilk is a hydrated, spongy to powdery assemblage of microcrystalline carbonate minerals,
resembling toothpaste or marzipan (Hill & Forti, 1997). Moonmilk is usually rich in microbial biomass that may or may not be involved in moonmilk formation (e.g., Borsato et al., 2000; Northup et al., 2000; Northup & Lavoie, 2001; Mulec et al., 2002; Van de Kamp et al., 2004; Cáhaveras et al., 2006; Blyth & Frisia, 2008), and the microbes are intermixed with carbonate minerals rather than form a separate, distinct biofilm on the surface (e.g., Borsato et al., 2000). We are currently characterizing the diversity of microbes and melofauna from the moonmilk, of which we determined there was \( \sim 10^5 \) microbial cells/ml and \( \sim 10^4 \) meiofauna cells/m\(^2\) (Paoletti et al., 2009).

Based on the assumption that animal tissues reflect the assimilated isotopic composition of their food (Ponsard & Arditi, 2000), we evaluated the food web position of \( C. \) *servadeii* and other cave animals with respect to the moonmilk and to the cave water using stable carbon and nitrogen isotope ratio analyses. Despite recent investigations that diverse microbial activities in non-sulfur based caves can support diverse food webs and trophic level interactions (e.g., Simon et al., 2003; Gerić et al., 2004; Opsahl & Chanton, 2006), moonmilk microbial communities have not been previously considered a food source for invertebrates, especially troglobitic cave beetles. In the present study, we assess whether \( C. \) *servadeii* consumes organic matter, such as microbial cells, in moonmilk as its main food source. This work extends our current understanding of subsurface food webs (Gibert & Deharveng, 2002) and may provide clues as to the unique evolutionary and morphological adaptations of \( C. \) *servadeii* and other troglobites in cave hypogrotic habitats.

**MATERIALS AND METHODS**

**Sample collection and microscopy**

Grotta della Foos formed within Monte Ciaurlec which is underlain by Cretaceous and Triassic limestone units (Gasparo, 1971; Beggio, 2009). The cave contains over 2600 m of passages. Specific geologic details regarding the cave can be found in Gasparo (1971). Four sampling locations within the cave were chosen, although \( C. \) *servadeii* was collected only from location "Site B" (Figure 1). Basic sampling details are withheld because the \( C. \) *servadeii* population size is not yet known and the cave is not protected; the beetles are at risk of over-collection. Foraging and feeding behaviors of \( C. \) *servadeii* were documented using digital and video cameras. A video of \( C. \) *servadeii* from the cave can be viewed at http://geol.lsu.edu/aengel/publications.htm and https://sites.google.com/site/maurizioguidopaolletti/download-area/videos.

Other cave invertebrates were collected throughout the cave, including the saprofagous beetles \( Orostygia pretneri \) and \( Bryaxis persici, \) the carabid beetle \( Anopthalmus nivalis charon, \) the aquatic isopod \( Monolistra tavalensis, \) and the terrestrial isopod \( Androniscus noduliger \) (Figure 1). For comparison, samples of surface soil and invertebrates from the soil and forest litter near the cave entrance were collected, including five different isopod species (\( Protracheoniscus politus amoenus, Armadillidium vulgare, \) Trachelipus sp., Trichoniscus sp., and \( Ligidium germanicum). \) A few \( A. \) *noduliger* specimens were also acquired from small dolines at the top of Monte Cavallo at an altitude of 2000 m. All specimens were identified by M.G. Paoletti (unpublished data).

\( C. \) *servadeii* and \( O. \) *pretneri* individuals were analyzed with scanning electron microscopy (SEM). Samples were either air-dried or washed in alcohol and covered with a thin gold layer. A Philips XL30 ESEM®-TEM environment (E-)SEM was used to observe non-dehydrated samples that were mounted on a thermostatically regulated holder. Experimental conditions were normally at 5°C at 4.5 to 6.5 Torr in the instrumental chamber. The ESEM was capable of X-ray fluorescence (XRF), induced by the electron primary beam and detected in Energy Dispersive Spectroscopy (EDS) mode, to determine the elemental composition of moonmilk.

**Water analyses**

Basic geochemical analyses of percolating water samples were done, including pH, temperature, and specific conductance using standard electrode methods. Water was filtered to 0.2 μm, and major anions and cations were measured by single column ion chromatography. The amount of dissolved organic carbon in cave water was determined by the dichromate odyometric method. Briefly, 50 ml of water were supplemented with 1 ml of 1 M K\(_2\)Cr\(_2\)O\(_7\), and subsequently acidified with 25 ml 96% H\(_2\)SO\(_4\). The absorbance at 600 nm was read spectrophotometrically and compared against a standard curve obtained with increasing D-glucose concentrations.
Carbon and nitrogen stable isotope systematics

Biological processes result in distinct isotopic values based on the differential utilization and isotopic discrimination (called fractionation). For the most part, metabolic discrimination between the lighter and heavier isotopes is predictable and can be used to track food sources among individuals within a community and estimate trophic level status. Comparisons of carbon (C) and nitrogen (N) isotope compositions are routine for terrestrial and aquatic food webs (e.g., DeNiro & Epstein, 1981; Post, 2002; West et al., 2006). Given its typical adaptations, we did not assume that *C. servadei* could leave the cave, and because we had not observed individuals in other parts of the cave, beetle C and N isotopic compositions were assumed to represent its local food source. However, we recognized that tracing an assimilated primary food source of a mobile animal is difficult because the isotopic values of an individual could be biased by its most recent feeding pattern and by tissue turnover rates for that individual (Post, 2002). As we know very little about the tissue turnover rates of *C. servadei* and the other invertebrates, we used whole individuals and attempted to use only adults to minimize possible differences in body tissue composition. Moreover, because it was unclear whether or not the beetle guts were completely purged prior to isotopic analyses, we analyzed the dissected gut from *Cansiliella* specimens separately.

The stable organic carbon (C) and nitrogen (N) isotopic composition of the invertebrates was done from the same sample, but the organic C and N isotope compositions from the soils and moonmilk were analyzed separately. Whole invertebrate specimens were dried, crushed, and weighed. Stable organic C isotope compositions from soil and moonmilk were done after removing inorganic carbonate with a solution of 1N HCl for 16 hr at room temperature while monitoring reaction pH (Midwood & Boutton, 1997). Because the acidification procedure could result in the loss of N-acid soluble organic matter and enhanced ingassing of NH₃, the N isotope analysis of the soils and moonmilk were performed on non-acidified samples. Samples were rinsed to pH 5.5 in DI H₂O, dried at 50 °C, and ground to a fine powder.

An aliquot from each sample was flash combusted on an EA 1110 Carlo Erba elemental analyzer coupled on line to an isotope ratio mass spectrometer (IR-MS) Delta plus Finnigan MAT that was operated in the continuous flow mode using He as a carrier gas. Combustion was in an oxygen atmosphere in a quartz reactor packed with Cr₂O₃ and CoOAg to form CO₂, N₂, NOx, and H₂O. The gases were passed through a copper furnace (650°C) to reduce NOx to N₂, and water was subsequently removed by MgCl₂. Finally, CO₂ and N₂ were separated by gas chromatography (4 m, 1/ 4-inch Poraplot Q column) and introduced to the IR-MS via the Finnigan ConFlo II open split interface with reference gases inserted as pulses of pure standard gases. Isotope compositions are expressed in standard delta notation (δ, units of ‰) relative to known standards (Pee Dee Belemnite for C and AIR for N). Precision of individual isotope ratio measurements, as measured by internal laboratory organic standards, was ±0.2‰ for δ¹³C and ±0.1‰ for δ¹⁵N values. Isotope results were normalized using IAEA-CH6, IAEA-CH7, and USGS24 for C and IAEA-N1, IAEA-N2 and USGS25 for N.

RESULTS AND DISCUSSION

*Cansiliella servadeii* behavior

*C. servadeii* were associated with active moonmilk deposition at Site B (Figures 1 and 2A). Individuals were observed submerged in the percolating waters that flow over the moonmilk (up to 1 to 40 mm water depth) (Figure 2). *C. servadeii* occurred in densities of 2 to 10 specimens/m² (with a mean of 3.8 specimens/m² on five different carbonate surfaces) (Figure 2B), and at least 20 specimens were observed partially to totally...
submerged in the percolating waters between 4 to 20 mm depth (Figure 2C). Engaged mouthparts denote feeding behavior, and when seen associated with the moonmilk, *C. servadeii* stopped from time to time with the head prone and mouthparts engaged at the water-moonmilk interface (Figure 2 D, E). *C. servadeii* spends considerable time (about 49%) preening its head, legs, and antennae; typically, this behavior is to move organic particles towards the mouth. Only on a few occasions were beetles found away from the percolating waters, and they moved much faster than in the percolating waters. We did not observe specimens inside stagnant water bodies within the cave or in stream passages. Additionally, beetles were not or rarely observed with engaged mouthparts (grazing or feeding) on the limestone host rock (Figure 2A). The movement and behavior can be readily observed from video (http://geol.lsu.edu/aengel/publications.htm).

![Environmental scanning electron photomicrographs of *C. servadeii* and *Orostygia pretneri* mouth parts.](image)

(A) Frontal view of *C. servadeii* mouth with a piece of moonmilk in mouth (arrows). Scale bar is 100 μm. (B) Frontal view of *O. pretneri* head and mouth parts. Scale bar is 100 μm (figure is intentionally turned upside down for comparison to panel A). (C) Left mandible (m sx) of *C. servadeii* showing wear on the tip and edge. Inset drawing is of the whole mandible. Scale bar is 20 μm. (D) Left mandible of *O. pretneri* for comparison to *C. servadeii* mandible. Inset drawing is of the whole mandible. Scale bar is 20 μm. (E) *C. servadeii* mouth part showing microbial cells (arrow) on the top right mandible. Scale bar is 100 μm. (F) *C. servadeii* top left mandible with microbial cell layer on the galea. Scale bar is 5 μm.
Although C. servadeii mouthparts have been previously described (Paoletti, 1972, 1973, 1980), we confirmed that the left and right mandibles are hoe-shaped (Figure 3A and C), and that the right branch had 4-5 short lobes and the left branch had two larger teeth. Carbonate particles up to 80 μm wide were found inside C. servadeii mouths (Figure 3A), and the mouthpart structures were consistently worn and abraded compared to those of O. pretleri (Figure 3C). Based on the behavior we observed, the most likely cause for abrasion would come from browsing relatively rigid substrate. The surfaces of C. servadeii galeae and mandibles had rod-shaped structures that we interpreted as adhered microbial cells (Figure 3D and E). From the size of the mouthparts, we estimated that it would be possible for C. servadeii to browse easily from ~10 to 60 μm depths into the moonmilk (Figure 2D and Figure 3A), although larger particles were also seen in their mouths (Figure 3A). Galeae were covered by rows of spines that resemble a brush or rack (Fig. 4 A,B,C). The spines were ~1-1.5 μm apart, which was appropriate spacing for bacterial cells to collect (Figure 4 D).

Physicochemistry of cave water and moonmilk

The percolating waters within Grotta della Foos were 8.8°C at a pH of 8.25, dissolved oxygen content of 10.8 mg/L, and conductivity of 225 μS/cm. Dissolved constituents were: Na⁺, 0.2 mg/L; K⁺, <0.05 mg/L; NH₄⁺, 0.3 mg/L; Ca²⁺, 45.2 mg/L; Mg²⁺, <0.05 mg/L; Cl⁻, 1.2 mg/L; HCO₃⁻, 149.5 mg/L; NO₃⁻, 0.3 mg/L; NO₂⁻, 4.5 mg/L; PO₄³⁻, 0.2 mg/L; SO₄²⁻, 3.5 mg/L. The dissolved organic carbon in the water had a mean value of 10.11 mg/L. The elemental composition of the moonmilk (by % weight) had 39% Ca, 39% O, 19% Al, 15% C, 3% Si, 2% Fe, and <1% each of Mg, P, S, and K. The organic C composition of the moonmilk was 0.06-0.52 %, with 0.01-0.05 % organic N. The bulk of the organic C was considered to be microbial biomass based on the lack of photosynthetically derived organic matter detected in the moonmilk (Beggio, 2009). With a C/N ratio for the moonmilk of ~9.4, and ~9.8 for the water, both the moonmilk and water have high, nutritionally balanced content and can support high bacterial growth efficiency (Del Giorgio & Cole 1998; Farjalla et al., 2009).

Stable isotope analyses

To examine the food web structure and trophic interactions occurring in Grotta della Foos for C. servadeii, we performed isotope ratio analyses to evaluate the status of the beetle in comparison with the habitat water, moonmilk, and other cave and surface animals (Figure 5). The δ¹³C value for water dissolved inorganic C was -17.1‰, and the δ¹³C value for organic C in water was -25.8‰. Moonmilk organic C averaged -24.2 ‰ (n = 10). The average δ¹³C value for C. servadeii bodies was -20.7‰ (n = 8). The separated gut contents (n = 3) had consistent isotopic values to the full body for several beetles (n = 3), suggesting that the full body measurements also represented the food source. C. servadeii δ¹³C values were similar to those for M. lavalensis (unpaired t-test P = 0.17), but
not to Site B A. noduliger and A. noduliger from Site D, or other invertebrates from the cave (Figure 5). The distinct position of C. servadeii in C and N isotope space can be explained by the other dietary habitats of other cave invertebrates. For instance, A. noduliger is a terrestrial isopod, and like O. pretneri, consumes plant and especially animal-derived organic detritus.

Animals occupying higher trophic levels should have heavier isotopic compositions compared to their diet, especially with ascending trophic level structure. For the C isotope system, an enrichment of 0.5 to 3‰ is expected for a consumer relative to its diet (e.g., Post, 2002), being attributed to preferential loss of the lighter isotope during respiration by the consumer, preferential uptake of 13C during digestion, or metabolic fractionation of tissue synthesis (e.g., lipids tend to have lighter composition compared to other tissues). The data would thus be compatible with C. servadeii obtaining its organic C either from the dissolved organic pool from the percolating water in the hygropetric habitat or from the moonmilk, as both have more negative δ13C compositions.

The N isotope values allow for the evaluation of possible food sources. The δ15N values for the percolating waters, containing mainly inorganic nitrate, was -10.9‰, with moonmilk having values around +2‰. Sustained microbial growth likely causes the δ15N values for the moonmilk to become positive if the microbial communities were utilizing organic N in the moonmilk, such as from beetle excrement (e.g., Macko & Estrep, 1984; Hoch et al., 1992; Lehnmann et al., 2002). The positive δ15N composition of the moonmilk is likely due to increased microbial degradation of isotopically depleted organic matter that causes the nitrate pool to become increasingly more depleted. With ascending trophic level and limited microbial recycling of N, the expected δ15N values of animals should increase by at least 3‰ (Peterson & Fry, 1987; Ponsard & Arditi, 2000; Post, 2002), which is attributed to enrichment in tissue content relative to food source due to excretion of 15N-depleted recalcitrant material (e.g., urea, ammonia). Because the body of C. servadeii had a mean δ15N composition of -4.8‰, our data appear to exclude the possibility that moonmilk is an exclusive source of food for C. servadeii, as the moonmilk δ15N composition is more positive than the insect. The δ15N composition of the percolating water, in which the beetle is constantly submerged, is a more probable candidate for the N-based nutrition of C. servadeii.
**Cansiliella food web interpretation**

In conclusion, although the specialized mouthparts for Cansiliella have been known for many years, little has been done to understand why Cansiliella is different from other troglobitic Leptodirinae (e.g., Paolotti, 1973, 1980; Moldovan et al., 2004; Sket, 2004). The feeding behavior of *C. servadiei*, and direct manipulation and association with moonmilk in the Grotta della Foos habitat, suggest that the insect may make use of the moonmilk for its nutrition. However, the stable C and N isotope data provide some evidence against supporting this hypothesis, particularly with respect to the δ15N composition of the beetles. Part of the inconsistent evidence is because of the unique microbial communities in the moonmilk, which are currently being described (Engel et al., unpublished data). Microbial recycling of isotopically variable C and N sources by those microbes, as well as the isotopic influence of living submerged in water with a unique isotopic composition, may play a role in the *C. servadiei* isotopic compositions. Nevertheless, wherever *C. servadiei* is obtaining its organic C and N for its diet directly from the water, or from the moonmilk, or both, the constant browsing and squeezing of the moonmilk would cause the observed, abraded beetle mouthparts. Further work will help us to understand the microbial communities in the moonmilk associated with *C. servadiei*, and will uncover additional unique explanations for morphological adaptations of other troglobites in cave hygropetric habitats.

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