Skeletal reorganisation affects geochemical signals, exemplified in the stylasterid hydrocoral *Errina dabneyi* (Azores Archipelago)

Max Wisshak1,*, Matthias López Correa1, Helmut Zibrowius2, Joachim Jakobsen3, André Freiwald1

1GeoZentrum Nordbayern, Universität Erlangen-Nürnberg, Loewenichstr. 28, 91054 Erlangen, Germany
2Station Marine d’Endoume, Rue Batterie des Lions, 13007 Marseille, France
3Rebikoff-Niggeler Foundation, Rocha Vermelha, 9900-451 Horta, Faial, Azores, Portugal

ABSTRACT: The pure white fans of the stylasterid *Errina dabneyi* are a conspicuous feature on the upper bathyal slopes in Azorean waters and were documented and recovered alive with the aid of a submersible in the southern Faial Channel. Etched vacuum-epoxy-casts of the aragonite skeleton, studied by scanning electron microscopy, reveal the 3-dimensional internal architecture comprising coenosarc canal meshwork, dactylopores, gastropores and ampullae. Near the surface, the canals are narrow and interconnected in a regular 3-dimensional meshwork. Deeper inside the canals are less abundant, more irregular and wider. This pattern implies that the skeletal architecture is modified during growth, with more central canals being enlarged by dissolution, and other canals, gastropores and dactylopores in turn being filled with aragonite reprecipitates. The skeleton is primarily composed of irregular spherulitic aggregates and overprinting during growth is evident from ghost structures in the form of successive semi-spherulitic infill of former canals. Due to differential dissolution and reprecipitation, this internal rebuild process inevitably involves an alteration of initial geochemical signatures such as stable isotope ratios ($\delta^{18}$O and $\delta^{13}$C), trace element signals and the distribution of radiogenic isotopes of carbon and uranium. This has to be taken into account when applying radiometric dating techniques and when using stylasterids as a geochemical archive.

KEY WORDS: Hydrozoa · Stylasteridae · Microstructure · Biomineralisation · Coenosarc canals · Geochemical signals · Stable isotopes · Epoxy-casting technique

INTRODUCTION

Coral skeletons, both solid calcareous and proteinaceous structures, are promising geochemical archives, for instance with respect to (palaeo)temperatures deduced from their oxygen and carbon stable isotope composition (reviewed in Swart & Leder 1996). This is based on their sessile nature, an often distinct incremental growth pattern and comparatively low growth rates, especially in deep water, with some species reaching centennial and even millennial life spans (Druffel et al. 1995, Risk et al. 2002, Andrews et al. 2005, Roark et al. 2009). The skeletal aragonite of deep-water scleractinians can be precisely dated with U-Th and radiocarbon techniques to reconstruct the timing of ventilation changes in deep and intermediate water masses (e.g. Cheng et al. 2000, Frank et al. 2004).

However, the use of corals as geochemical archives to reconstruct past temperature and productivity regimes in the deep oceans also presents various methodological and interpretational pitfalls, most of which are related to biological fractionation processes known as ‘vital effects’ (e.g. Smith et al. 2000, Cohen et al. 2006). This applies to many azooxanthellate cold-water scleractinians, including the intensely studied *Lophelia pertusa* Linnaeus, 1758 (Blamart et al. 2005, Lutringer et al. 2005). On a quest for other suitable deep-water coral species, geochemical research now...
includes primnoid (Heikoop et al. 2002, Sherwood et al. 2005, Noé et al. 2007), coralliid (Weinbauer et al. 2000) and isidid octocorals (Thresher et al. 2009), as well as zoanthids (Druffel et al. 1995) and antipatharians (Love et al. 2007, Williams et al. 2007).

Another promising, but up to now little studied, group of corals are calcareous hydrocorals, specifically stylasterids. In a pioneering paper, Weber & Woodhead (1972) found that tropical shallow-water hydrozoans, contrary to most scleractinians, actually precipitate their aragonitic skeleton close to equilibrium with seawater. They investigated representatives of the stylasterid genera *Stylaster* and *Distichopora*, alongside *Millepora*. Results by Mienis (2008) indicate that this is also true to some degree for deep-water *S. gemmoscens* (Esper, 1794) from the Rockall Trough (NW Atlantic). Likewise, preliminary results by Andrus et al. (2007) show the deep-water *S. erubescent* Pourtalès, 1868 from the Blake Plateau (N Atlantic) records stable isotopes in equilibrium with (or in uniform offset from) seawater, showing a sinusoidal variation related to apparent incremental banding.

In the present study, we critically evaluate the suitability of stylasterids for geochemical analyses—exemplified by *Errina dabneyi* (Pourthalès, 1871) from the upper bathyal slope of the Azores Archipelago. This evaluation is based on an investigation of its skeletal architecture, the deduced growth mechanisms and its stable isotope signatures (δ18O and δ13C). In fact, stylasterids are characterised by a complicated inner architecture that has to be taken into account. This inner architecture was first analysed and illustrated in a remarkable way by Moseley (1879, 1881), then revisited by Cairns (1983) as the basis for systematic revision requirements. The oldest representatives of the genus *Errina* are known from the Danian (65 to 61 million yr) deeper water coral limestone of Faxe, Denmark (Bernercker & Weidlich 1990, Lindner et al. 2008). At present, 26 species of *Errina*, most from deeper waters, are known worldwide (Cairns 1983). *E. dabneyi* seems to be endemic to the Azores Archipelago and the adjacent Mid-Atlantic Ridge region (Zibrowius & Cairns 1992).

**MATERIALS AND METHODS**

**Documentation and sample recovery.**

Colonies of *Errina dabneyi* were video- and photo-documented in situ in the southern Faial Channel during several dives with the research submersible ‘LULA’ between 2003 and 2008 (Fig. 1, Table 1). A total of 3 specimens were recovered during Dives 117 (2006; see Figs. 4–7A–C,E), 126 (2007) and 136 (2008; see Figs. 2B,C & 3A–D) in close to 500 m water depth with the aid of a manipulator and a sampling basket; specimens are herein referred to by their respective dive numbers.

**Growth rate analysis.** An estimation of the growth rate was obtained via 2 time series of in situ still images, taken over 2 and 5 yr, respectively, from selected *Errina* colonies (Table 1). One of the photographed specimens (Specimen 136) was subsequently recovered for exact scale determination and later stable isotope analyses, while the other was left in situ for further long-term growth pattern monitoring.

**Microstructure analysis.** *Errina* skeletons were vacuum-epoxy-impregnated with Araldite BY158 and Aradur 21 prior to sectioning, grinding and polishing. The skeletal architecture and microstructure was studied by staining with Feigl solution for discrimination of the carbonate mineral phase and CamScan scanning electron microscopy (SEM) of polished gold sputter-coated samples after treatment with 0.1 mol l–1 HCl for 30 s. To visualise the internal canal system, the resin-impregnated samples were treated with ~1 mol l–1 HCl for up to 1 h in order to remove part or all of the skeletal carbonate, leaving only the casts of the canals which were then studied with SEM.

**Stable isotope analysis.** MicroMill (New Wave) technology was applied to obtain high-resolution carbonate powder samples from epoxy resin-embedded (resin non-soluble in phosphoric acid) and slow-speed
sawed (Bühler IsoMet 1000) sample blocks for the stable isotope analyses (δ\(^{18}\)O and δ\(^{13}\)C). Individual tracks were ~4 mm in length and 100 µm wide with an average spacing of 130 to 140 µm. To enhance the faint growth increments, as a guide for placement of MicroMill tracks, a polished and briefly etched sample was gold sputter-coated, photographed and the contrast increased with Adobe Photoshop. The 20 mm long transect across the mid-plane of a basal trunk of a medium-sized (12 cm height) *Errina dabneyi* colony (Specimen 117) comprised 147 consecutive sampling tracks. Additionally, this and 2 other colonies (Specimens 126 and 136) were sampled along their surface (~3 mm wide and very shallow spot samples) with a hand-held dental drill to screen isotopic differences between contemporaneously grown proximal adult and distal juvenile surfaces. The carbonate powders were reacted with 100% phosphoric acid (density > 1.9) at 75°C using a Kiel III online carbonate preparation line connected to a Thermo-Finnigan MAT 252 mass spectrometer. All values are reported in per mil relative to Vienna-PeeDee Belemnite (V-PDB). The reproducibility (1σ) is better than ±0.04‰ for δ\(^{13}\)C and ±0.05‰ for δ\(^{18}\)O.

### RESULTS AND DISCUSSION

#### Habitat characteristics and distribution

*Errina dabneyi* is a common faunal element on the upper bathyal slope of the southern Faial Channel and was encountered on most of the dives that targeted waters deeper than ~350 m down to the submersible’s diving limit of 500 m (Fig. 1, Table 1). The highest population density was found at around 450 m water depth, where this coral occurs on bedrock outcrops that often also bear aggregations of the deep-sea oyster *Neopycnodonte zibrowii* Gofas, Salas and Taviani in Wisshak et al., 2009 and the cyrtocrinoid *Cyathidium foresti* Cherbonnier and Guille, 1972 (Wisshak et al. 2009a,b). Most of the previous samples of *E. dabneyi* from the Azores, including the type material, were recovered from 215 to 500 m water depth (Zibrowius & Cairns 1992). Further material stems from the Mid-Atlantic Ridge to the southwest from 1400 to 2200 m water depth (Zibrowius & Cairns 1992), but this much greater depth seems strange given the considerable divergence from confirmed records and in situ (submersible) observations in the Azores. *E. dabneyi* was not among the abundant stylasterid fauna sampled (by the cruise SEAMOUNT 2 in 1993) on Great Meteor and other seamounts situated a few degrees south of the Azores archipelago.

The pure white, densely branched, flabellate colonies are up to ~40 cm in size and are predominantly found on the more current-exposed ridges and cliff tops, with their concave side facing the current (Fig. 2). Quite often, solitary scleractinians, mainly *Desmophyllum dianthus* (Esper, 1794) (= *Desmophyllum cristagalli* Milne-Edwards and Haime, 1848) and *Caryophyllia cyathus* (Ellis and Solander, 1786), were found attached to the stylasterid fan facing downstream (Fig. 2B,C). The base and the current-facing side of the fan is less often colonised by scleractinians but is frequently overgrown by sponges (Fig. 3).

<table>
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<tr>
<th>Dive no.</th>
<th>Date (dd.mm.yy)</th>
<th>Start Coordinates</th>
<th>Depth (m)</th>
<th>End Coordinates</th>
<th>Depth (m)</th>
<th>Specimens</th>
<th>Photo/video-documentation</th>
<th>Water samples (depth)</th>
<th>Logger deployment/recovery</th>
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<td>475</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
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<tr>
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<td>13.09.06</td>
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<td>500</td>
<td>38°29.218’N 28°37.404’W</td>
<td>495</td>
<td>X</td>
<td></td>
<td></td>
<td>(= 136)</td>
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<tr>
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<td>17.09.06</td>
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<td>490</td>
<td>38°29.400’N 28°37.400’W</td>
<td>350</td>
<td>117 (isotope transect)</td>
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<td></td>
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<tr>
<td>125</td>
<td>24.09.07</td>
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<td>495</td>
<td>38°29.510’N 28°37.400’W</td>
<td>350</td>
<td>X</td>
<td>X (500 m)</td>
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<td>495</td>
<td>38°29.218’N 28°37.404’W</td>
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<td>X (= 136)</td>
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A symbiont regularly found occupying the more distal branches on both sides of the fans is the prosobranch gastropod *Pedicularia sicula* Swainson, 1840 (Fig. 4A–C). Representatives of the genus *Pedicularia* are common obligate symbionts on stylasterids worldwide, including in the NE Atlantic. The snail typically deposits a layer of lime under which the sealed-off stylasterid skeleton cannot grow in thickness, leading to a characteristic attachment scar (Fig. 4C) on which the foot of the snail adheres efficiently (Zibrowius & Cairns 1992, Goud & Hoeksema 2001).

**Growth rates**

While pronounced destruction and loss of branches was visible in both monitored colonies (Fig. 3C,F), the extent of skeletal growth at the outer margin of the fan was minute and of the magnitude of only a few millimetres per year. One monitored colony (Specimen 136; Figs. 2B,C & 3A–D) was sampled after the last photo was taken, allowing for direct measurement of the areas where the colony had remained intact; in places, branches had grown 4 to 6 mm in length per year. In the case of the other monitored colony (not sampled; Fig. 3E,F), the partial damage was possibly caused by an unfortunate manoeuvre with the submersible only minutes after the first photo was taken. In the 5 yr and 2 mo since, the colony regenerated branches <2 cm in length, even though precise scaling was impossible because the colony was left *in situ* for further long-term observations. Potential natural hazards for the fragile colonies were captured on film when a crab walked across a colony (Specimen 136; Fig. 3B, arrow) and a large ray almost hit the same colony during Dive 126 (Fig. 3D), possibly distracted by the lights or electromagnetic field of the submersible. For statistically sound growth rates of *Errina dabneyi*, further *in situ* time series—preferentially supported by calcein or manganeese tagging experiments—are needed to gain a better understanding of how they vary with respect to colony size, position, and recovery from partial mortality.

The only other stylasterid studied with respect to growth rate is *Errina novaezelandiae* Hickson, 1912. Chong & Stratford (2002) obtained growth rates (branch lengthening) from calcein tagging and stereo photography and found positive values of 3.1 ± 0.6 and 8.0 ± 0.9 mm yr⁻¹, respectively. They also recorded considerable negative growth caused by breakage and bioerosion, leading to a net growth rate of 2.7 ± 1.1 mm
yr^{-1}. For the same species, Miller et al. (2004) found highly variable rates with a net 7 mm yr^{-1} and a very high maximum rate of 680 mm yr^{-1}, largely independent of colony size. However, this New Zealand species is primarily calcitic in mineralogy (Cairns & McIntyre 1992) and was studied in a shallow-water fjord setting within the reach of scuba diving (15 to 40 m). In this setting, higher nutrient fluxes can be presumed to exist than on the bathyal slopes in Azorean waters, which may explain the higher growth rates.

Skeletal architecture

Errina dabneyi colonies grow as pure white, uniplanar (and upstream slightly concave) fans up to 40 cm wide. The dactylopores and gastropores are crowded on terminal branches (Fig. 4D,E), whereas the more basal stems bear only scarce coenostreal papillae (Fig. 4F).

The vacuum-epoxy casting technique, adopted and modified from microbioerosion investigations (after Wisshak 2006, modified from Golubic et al. 1970), proved useful for the SEM study of the 3-dimensional internal architecture of the coenosarc canal meshwork, dactylopores, gastropores and ampullae (Figs. 5 & 6). The method is simple, time-efficient, cheap and bears good potential for studying permeable canal systems, pores and the like in calcareous skeletons.

In the peripheral layer of the branches, near the surface, the canals are ~10 µm wide and branch in a regular pattern with a mesh width of around 100 µm (Fig. 5). Deeper inside, the canals are more irregular and wider, some running along the central axis of the branches (Fig. 5D,G,I). In the thinner peripheral branches, the gastropores with the central gastrostyle are primarily oriented distally at oblique angles (Fig. 5A). They are connected to the canal meshwork at their walls and particularly at their base (Fig. 5B,C). In a more proximal position and in the more peripheral
layer, the branches contain subspherical to irregularly shaped ampullae, 300 µm and more in diameter, which are connected to the wider canals (Fig. 5D,F).

**Skeletal reorganisation**

Analysis of the patterns of the 3-dimensional coenosarc canal meshwork indicates that the skeletal architecture undergoes modification during growth. This follows simple geometric reasons. With growth in thickness of the branches, the formerly superficial meshwork of narrow canals is, with progressing growth, located deeper inside where then, however, wider canals are observed. Accordingly, some of the central canals have become enlarged by dissolution of skeletal material, whereas at the same time other canals have been filled-in with aragonite reprecipitates. Jointly, during this process the dactylopores and gastropores are entirely resorbed. Likewise, the ampullae are filled or remain open but embedded while new active ampullae are formed closer to the surface.

This process of transformation is recorded in the microstructure of the stylasterid skeleton (Fig. 6). The aragonitic nature of the skeleton was positively tested via Feigl solution and is primarily composed of full- and semi-spherulitic aggregates (Fig. 6A,B), the typical microstructure of stylasterids (Fenninger & Flajs 1974). This structure is overprinted during growth, as indicated by ghost structures in the form of successive semi-spherulitic in-fill of former canals (Fig. 6B–E). Where side branches were filled with reprecipitates (Fig. 6D), the plug-like ending can be seen from inside the canal (Fig. 6F). Still to be resolved is whether the dissolved skeletal material is actively transported by the live tissues to the sites of reprecipitation, potentially a more efficient biomineralisation process.

The proximal thicker trunk of the colony often consists of brownish, dead coenosteum that may undergo rejuvenescence by new, white coenosteum spreading over from the more peripheral living regions. Distinct growth breaks are thus formed which are characterised by partly preserved peripheral canal meshwork (Fig. 6G,H). Sometimes encrusting epibionts and traces of microbioerosion are thus immurated by the overflow of new coenosteum. This indicates that the coenosteum, once dead (i.e. devoid of live tissues in the canal meshwork), is not or only partly subjected to further restructuring. These growth breaks may have the appearance of cyclic increments; however, this is not their actual nature. In the denser centre of the basal

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Fig. 4. *Errina dabneyi* and *Pedicularia sicula*. SEM images illustrating morphological details of Specimen 117 and its symbiont. (A) Symbiotic prosobranch gastropod *P. sicula* and (B) close-up of the early ontogenetic shell of a juvenile. (C) Characteristic attachment scar of *P. sicula*. (D) Terminal branch of *E. dabneyi* bearing many dactylopores. (E) Typical circular gastropore and its central gastrostyle. (F) Coenosteum of a more proximal branch devoid of dactylopores and gastropores but bearing many small coenostreal papillae.
Fig. 5. *Errina dabneyi*. SEM images of Specimen 117 that was vacuum-resin-embedded and decalcified with hydrochloric acid, illustrating the 3-dimensional internal architecture. (A–C) Terminal branch bearing a number of gastropores and dactylopores, integrated into a dense meshwork of coenosarc canals. (B) Sectioned gastropore exposing the gastrostyle. (C) Gastropore base with radiating connections to the canal meshwork. (D–F) Sub-terminal branch without gastropores or dactylopores but showing ampullae and recording the development of wider canals in the centre, replacing the former fine meshwork of peripheral canals. (E) Peripheral coenosarc meshwork. (F) Ampullae. (G–I) Proximal stem (only partially etched, retaining part of the skeletal matrix; SEM composite image) showing denser central part with irregularly formed and oriented large canals and the fine meshwork restricted to the periphery. (H) Peripheral and (I) central meshwork.
trunk, only a faint incrementation can be detected (enhanced in Fig. 7B). The centre of the trunk is, however, often found intensely bored by hadromerid sponges (Fig. 6I) which may also extend upwards into the living parts of the colony.

Stable isotope signature

In order to evaluate to what degree *Errina dabneyi* precipitates its stable isotope composition of aragonite in equilibrium with the ambient seawater, a complete seasonal cycle of *in situ* temperatures was logged in high resolution, and the stable isotope composition of the ambient seawater in 500 m water depth was monitored (Wisshak et al. 2009a). Temperature was stable with an annual mean of 12.3 ± 0.25°C, monthly means differed by <0.4°C, short-term fluctuations were on the order of 1°C (min. = 10.86°C, max. = 13.52°C) and the δ¹⁸Owater varied between 0.47 and 0.64‰ with a mean of 0.54 ± 0.06‰ relative to V-SMOW (Vienna-Standard Mean Ocean Water) (Wisshak et al. 2009a). Using these background data, an expected mean equilibrium at 2.40‰ δ¹⁸O relative to V-PDB can be calculated via the equation of Grossman & Ku (1986): T = 20.6 – 4.34 (δ¹⁸Oarag – δ¹⁸Owater), where T is temperature (°C). Considering the standard deviation of both parameters, a skeletal δ¹⁸O range of 2.29 to 2.52‰ would be expected. For the unlikely case that the extreme temperature and δ¹⁸Owater values are reflected in the iso-

Fig. 6. *Errina dabneyi*. SEM analysis of the microstructure of the aragonitic skeleton (Specimen 117). (A,B) Irregular spherulitic aggregates. (B–E) Ghost structures in form of successive semi-spherulitic infill of former canals; former tunnel delineated by dashed line and direction of infill indicated by arrows. (F) Terminus of a side branch filled with reprecipitates as seen from inside a canal. (G,H) Apparent incrementation in the periphery of the basal stem, formed by new skeletal material being formed on dead coenosteum, retaining part of the former peripheral meshwork. (I) Spiculae of a boring hadromerid sponge, commonly found tunnelling the dense centre of the basal stems.
Fig. 7. Errina dabneyi. Stable isotope ($\delta^{18}O$ and $\delta^{13}C$) analysis of 3 skeletons (Specimens 117, 126 and 136). (A) Medium-sized colony (Specimen 117, 490 m) with the position of surface samples and the orientation of the high-resolution transect. (B) MicroMill transect in a longitudinal section across the basal stem of the colony. (C) Stable isotope transect showing an overall ontogenetic decrease in $\delta^{13}C$ after a rapid initial increase, and constant fluctuations in $\delta^{18}O$ at both sides of the growth centre with poor high-resolution symmetry. (D) Cross plot of $\delta^{18}O$ versus $\delta^{13}C$ of terminal branches versus more proximal surfaces in 3 colonies sampled during Dives 117, 126 and 136 in ~500 m water depth. (E) Cross plot of $\delta^{18}O$ versus $\delta^{13}C$ from the transect showing low correlation and moderate range. Note the depleted $\delta^{13}C$ values of the centre and the zone beyond the growth break.
topic record, a maximum \( \delta^{18}O \) range of 2.06 to 2.83\%o would be expected (Fig. 7C–E).

The actual mean oxygen stable isotope values of *Errina dabneyi* (1.5 ± 0.4\%o, \( \delta^{18}O \) V-PDB; Specimens 117, 126 and 136; \( n = 182 \)) show a negative offset from the expected equilibrium by ~0.9\%o, with the heaviest values closest to, but still 0.4\% offset and outside, the extreme equilibrium range (Fig. 7D,E). The overall range (~0.9\%o) of the oxygen values is moderate. Hence, unlike shallow-water *Millepora*, *Stylaster* and *Distichopora* which were reported to precipitate aragonite in equilibrium with seawater (Weber & Woodhead 1972), *E. dabneyi* is prone to a moderate biological fractionation process leading to an average 0.9\%o negative offset in \( \delta^{18}O \). However, none of the environmental settings for the previously studied milleporid and stylasterid hydrocorals comprised such constant conditions as encountered in the upper bathyal Azorean waters; therefore, seasonal variations could have complicated a concise equilibrium assessment. Weber & Woodhead (1972) also considered all their sampling sites as uniform with respect to the ambient seawater \( \delta^{18}O \) composition and a less accurate temperature equation was used. It is therefore possible that similar biological fractionation might still be unveiled in the previously investigated hydrocorals. Indeed, the negative 0.9\%o shift in \( \delta^{18}O \) from the expected equilibrium values for skeletal aragonite has also been found in *S. gemmascens* from the southwestern Rockall Trough (Mienis 2008, her Fig. 7.3). But the absolute range (~0.7\%o) of skeletal \( \delta^{18}O \) in *S. gemmascens* roughly coincides with the estimated range calculated from the ambient temperatures of 6.2 to 9.8\°C, and with the range observed for the Azorean *E. dabneyi*. A similar, possibly constant, negative offset from equilibrium has also been invoked by Andrus et al. (2007) for *S. erubescens* from the Blake Plateau (NW Atlantic).

Another prerequisite for the application of an organism as a reliable high-resolution geochemical archive is the reproducibility of the signal across coeval time series of incremental banding. Hence sample transects on both sides of the growth axis should show a clear symmetry as an essential pattern. But here, the above outlined factor of skeletal reorganisation comes into play. The high-resolution transect across the main trunk of Specimen 117 shows a mean of 1.6 ± 0.3\%o, \( \delta^{18}O \) V-PDB with short-term fluctuations of up to 0.5\%o (Fig. 7C,E). In light of the logged stable temperature, this short-term amplitude must be considered far too high for reliably mirroring temperature fluctuations.

The basal trunk shows in its geochemical pattern 3 different zones: (1) the \( \delta^{13}C \)-depleted centre, (2) the somewhat symmetric layers on either side of the centre, and (3) following a growth break, a rejuvenescent zone with higher \( \delta^{18}O \) and very low \( \delta^{13}C \) composition (Fig. 7B,C). The centre is characterised by a single point with a depleted \( \delta^{13}C \) composition of 0.7\%o V-PDB, which we interpret as a buried relict of the earliest ontogeny. The immediately adjacent central layers show much heavier \( \delta^{13}C \) compositions of 1.8 to 2.0\%o V-PDB. To both sides of the centre, a slight decrease in \( \delta^{18}O \) was found but no high-resolution symmetry can be recognised. The \( \delta^{13}C \) values exhibit a mean of 1.6 ± 0.2\%o V-PDB with maximum values clustering just below 2.0\%o V-PDB and an overall range of close to 1.5\%o. This range is largely due to a general ontogenetic decrease in \( \delta^{13}C \) following the rapid ontogenetic increase in the earliest ontogeny, and shows a moderate degree of symmetry. This overriding ontogenetic trend is visible despite the skeletal reorganisation, and is coherent with the observed general difference between proximal and distal surface portions (see below). As a consequence of this ontogenetic imprint, the overall correlation between \( \delta^{18}O \) and \( \delta^{13}C \) is poor (\( r^2 = 0.26 \) and 0.38 for the 2 sides), and the slopes of the regressions are considerably inconsistent in different ontogenetic parts of the skeleton, most likely as a result of metabolic effects on the \( \delta^{13}C \) values, thus complicating an application of the lines technique (after Smith et al. 2000) for overcoming the ‘vital effect’. Conspicuous non-symmetric drops of the \( \delta^{13}C \)-composition below the ontogenetic trend to values between 0.6 to 1.5\%o are considered as remnants of former depleted surface compositions, which have only partly been modified by the skeletal reorganization. This also implies that the reorganization is not affecting each portion of the skeleton in the same way and with the same efficiency. On one side of the transect, an outer 3 mm of younger skeleton was precipitated following a distinct growth break. This layer is also visible as whitish zone on the right side of the basal trunk (Fig. 7A). Its oxygen isotope values are slightly higher than in the adjacent adult trunk (Fig. 7C). The respective much lower \( \delta^{13}C \) depicts the variability in the external layer, prior to the modification during the ubiquitous skeletal reorganization.

Further evidence for the skeletal reorganization comes from a comparison between the isotopic composition of the surface layer, from contemporaneous proximal (adult) to distal (juvenile) portions (Specimens 117, 126 and 136, \( n = 35 \); Fig. 7D), with the isotopic composition across the internal layers from the high-resolution MicroMill transect in the basal trunk (Specimen 117, \( n = 147 \); Fig. 7E). While the composition of oxygen isotopes shows similar ranges and absolute values between 1.2 and 2.0\%o \( \delta^{18}O \) V-PDB in all skeletal portions, there are strong differences for carbon isotopes. The most depleted \( \delta^{13}C \) is encountered in the distal terminal branches, with similar
extreme values of ~−0.3‰ δ13C V-PDB. Likewise, there is a clear difference between these surface layers in distal portions and contemporaneous proximal surface layers, which exhibit the heaviest external δ13C-values in each colony. Even heavier δ13C compositions are encountered in the internal layers—a fact which clearly highlights later alteration of the isotopic composition during the skeletal reorganization, which has been verified by our microstructural investigations.

**Implications for geochemical signals**

In conclusion, the present investigation on *Errina dabneyi* suggests that the internal reorganisation of the coenosarcs canal meshwork with considerable differential dissolution and precipitation alters not only initial stable isotope signals, but most likely also trace element signatures, limiting the value of stylasterids with a similar growth pattern as high-resolution geochemical archives. Stable isotope and trace element transects must be expected to have undergone a certain degree of signal averaging and modification. However, since the major zone of reprecipitation—the infill of the dense peripheral canal meshwork—is steadily moving outwards, following the zone of initial precipitation, overall trends in the geochemical signals may be partially retained, such as the ontogenetic decrease in δ13C as shown above. The bulk geochemical composition of the skeleton might also bear meaningful environmental signals, such as the bulk δ18O when considering the relatively constant negative offset of 0.9‰ observed in *Errina dabneyi* and other stylasterids. However, it needs to be taken into account that stylasterid architecture and the degree of skeletal reorganisation may vary among different species, for instance the shallow-water species *Distichopora vervoorti* and *Pedicularia vanderlandi*, which has more robust branches and poorly defined pore rows (Lindner et al. 2004).

The skeletal reorganisation likely also affects the distribution of radiogenic isotopes such as 14C, 210Pb, 230Th and 238U. Hence, radiometric dating as a prerequisite for time-referencing the geochemical archive must be expected to yield underestimated ages. This has been suggested for *Paragorgia arborea* (Linnaeus, 1758), which is very likely affected by a similar skeletal reconstruction, leading to a potential modification of its radiocarbon composition (Sherwood & Edinger 2009, their Fig. 6). Furthermore, precise time-referencing of geochemical transects through the basal stem of a stylasterid will be complicated by the potential occurrence of growth breaks (no cyclic increments as discussed above) of unknown duration close to the periphery.

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**LITERATURE CITED**


