Schizasterid Heart Urchins Host Microorganisms in a Digestive Symbiosis of Mesozoic Origin

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Because of their lifestyles, abundance, and feeding habits, infaunal marine deposit feeders have a significant impact on the ocean floor. As these animals also ingest microorganisms associated with their sediment and seawater diet, their digestive tract usually contains a diverse array of bacteria. However, while most of these microorganisms are transients, some may become part of a resident gut microbiome, in particular when sheltered from the main flow of digesta in specialized gut compartments. Here, we provide an in-depth analysis of the structure and contents of the intestinal caecum (IC), a hindgut diverticulum found exclusively in schizasterid heart urchins (Echinoidea: Spatangoida: Schizasteridae). Based on specimens of Brisaster townsendi, in addition to various other schizasterid taxa, our structural characterization of the IC shows that the organ is a highly specialized gut compartment with unique structural properties. Next generation sequencing shows that the IC contains a microbial population composed predominantly of Bacteroidales, Desulfobacterales, and Spirochaetales. The microbiome of this gut compartment is significantly different in composition and lower in diversity than the microbial population in the sediment-filled main digestive tract. Inferences on the function and evolution of the IC and its microbiome suggest that this symbiosis plays a distinct role in host nutrition and that it evolved at least 66 million years ago during the final phase of the Mesozoic.

Keywords: Echinoidea, microbiome, digestive tract, intestinal caecum, spirochete, Brisaster

INTRODUCTION

All animals are immersed in a microbe-filled world and interact with microorganisms across body surfaces exposed to the environment such as the integumentary, respiratory, urogenital, or digestive systems (McFall-Ngai, 2015). In many cases, microbial communities (or microbiomes) distinct from environmental bacterial communities reside on these surfaces or inside specific body compartments (Shade and Handelsman, 2012; Dubé et al., 2019; Zhang et al., 2019). The diverse metabolic activities of these microbiomes are often of critical physiological importance to the host, as is perhaps best exemplified by the many functions of the gut microbiome: protection against colonization by pathogens, development of the host's immune system, breaking down toxic components of the diet, digestion of refractory components, and provision of small organic molecules valuable to the host organism (Dubilier et al., 2008; Cavanaugh et al., 2013;
Sommer and Bäckhed, 2013). In fact, gut microbiomes are of such importance to their hosts that they have significantly influenced the evolution of metazoan digestive physiology as well as gut development and morphology (McFall-Ngai et al., 2013; Graf, 2016; Moran et al., 2019).

However, although digestive symbioses are known to occur in a wide range of metazoans, not all animals appear to establish a gut microbiome: in some species, the only microorganisms routinely found in the digestive tract are ingested with the food, and are therefore only transiently present (De Ridder and Foret, 2001; Hammer et al., 2019). One guild of animals for which this may generally be the case are marine deposit feeders, which often occur at high densities over large areas of the ocean floor (Jumars, 1993), and whose feeding and burrowing activities have significant biogeochemical consequences (Lopez and Levinton, 1987; Rosenberg, 2001). Most marine deposit feeders possess simple, tubular guts through which ingested material passes in a continuous, rapid flow (Penny and Jumars, 1987, 1990; Jumars, 2000), potentially hindering the establishment of a gut microbiome. However, even in such a scenario, theoretical considerations suggest that the potential for microbial activity might be particularly high in the hindgut, and that marine deposit feeders should be able to take advantage of this localized activity by retaining stable bacterial communities inside evacuations or pouches (Plante et al., 1989). Such hindgut diverticula have been observed in several marine deposit feeders, but evidence of microbial activity in these organs is sparse (Plante et al., 1990; Harris, 1993; Jumars, 1993).

In this study, we provide a detailed description of the structure and contents of such a hindgut compartment, the intestinal caecum (IC). This particular organ is found in some members of a widespread and abundant group of marine deposit feeders, the schizasterid heart urchins (Echinoidae: Spatangoida: Schizasteridae). Spatangoids are ubiquitous infauna on the continental shelf (Nichols, 1959; Moffitt et al., 2015), and as active burrowers play an important role in marine benthic ecosystem functioning through bioturbation (Bromley and Asgaard, 1975; Hollertz and Duchêne, 2001; Widdicombe et al., 2004; Lohrer et al., 2005; De Gibert and Goldring, 2008). Their activities result in transport of the organic-rich surface layer into deeper sediment layers and thus enhance benthic production both in shallow and deep sea habitats, while additionally increasing the diffusion of oxygen and uptake of carbon by the sediment community (Pequignat, 1970; Osinga et al., 1997; Hollertz, 2002; Vopel et al., 2007; Boon and Duineveld, 2012). Schizasterids, for example, possess a range of morphological features that permit burrowing in silty or muddy sediments, habitats that are frequently characterized by extreme hypoxia (Mortensen, 1951; Cheshire, 1963; Buchanan et al., 1980; Kanazawa, 1992; Schinner, 1993; Bromley et al., 1995; Thompson and Riddle, 2005; Sato et al., 2017). While the main digestive tract of spatangoids is packed with inorganic and organic particles taken up from the ocean floor, structural derivatives of the gut have been shown to be devoid of sediment: these functionally specialized organs include the primary and secondary siphon (De Ridder and Jangoux, 1982; Thorsen, 1998), the gastric and recto-intestinal caecum (Ziegler et al., 2010; Rolet et al., 2012; De Ridder and Saucède, 2020) as well as the IC discussed here. Given its position and the fact that the IC has been shown to contain a flocculent mass instead of sediment (Ziegler, 2014), we predict that this organ could be the site of a complex, yet undescribed digestive symbiosis. In a distantly related group of heart urchins, a similar structure was previously shown to contain nodules composed of a diverse array of microbial taxa that assist in digesting bulky inorganic or organic components forming part of the sediment diet (De Ridder et al., 1985; da Silva et al., 2006). This raises the question whether similar mechanisms could have independently evolved in schizasterids as well.

To better characterize the IC and its contents, we examined the well-studied and relatively easily obtainable schizasterid Brisaster townsendi using a diverse set of imaging techniques, including magnetic resonance imaging (MRI), micro-computed tomography (μCT), light microscopy (LM), scanning electron microscopy (SEM), and transmission electron microscopy (TEM), complemented by next generation sequencing (NGS). In addition, we obtained structural and genomic data on the IC from a representative of the schizasterid genus Abatus. This taxon has recently emerged as a model for ecological, biogeographical, and evolutionary research on benthic invertebrates of the Southern Ocean (Linse et al., 2008; Féral and Poulin, 2011; Michel et al., 2016; Fabri-Ruiz et al., 2017; Guillaumot et al., 2017). Finally, we compared our structural and volumetric observations of the IC in Brisaster and Abatus with other schizasterid taxa to establish inferences on the timing of origin and evolution of a potentially highly specialized organ found in an abundant group of marine deposit feeders.

**MATERIALS AND METHODS**

**Specimens**

Living individuals of Brisaster townsendi were obtained in May 2018 by otter trawl off the coast of Southern California on the San Pedro Escarpment (33°40.802 N, 118°19.764 W, 266 m depth to 33°41.016 N, 118°19.999 W, 275 m depth). This upper continental shelf soft sediment site lies within an area systematically monitored over the past decades with regard to various environmental and ecological parameters (Thompson et al., 1987; Thompson et al., 1993; Allen et al., 1999; Stull et al., 2001; Sato et al., 2017; Munakata and Markle, 2018). During the ca. 1 km long transect on a southwest-facing slope parallel to the coastline, several hundred individuals of B. townsendi were captured, in addition to numerous specimens of Strongylocentrotus fragilis and other infaunal and epibenthic animals. The B. townsendi sample was compared with eleven specimens of the closely-related schizasterid Abatus cordatus collected by SCUBA in the Golfe du Morbihan, Kerguelen Islands in November 2015 off Port-aux-Français (49°21.15 S, 70°13.083 E, 6 m depth) and in December 2015 off Île Suhm (49°30.90 S, 70°09.14 E, 10–15 m depth). Additional schizasterid species were obtained as ethanol-preserved specimens from various museum collections (Table 1). Species selection was aimed at complementing the list of schizasterid and selected outgroup taxa previously analyzed with regard to absence or presence of the IC (Ziegler, 2014).

| Specimen | Date       | Location                  |
|----------|------------|---------------------------|
| Brisaster townsendi | May 2018 | San Pedro Escarpment       |
| Abatus cordatus  | November 2015 | Port-aux-Français      |
| Abatus cordatus  | December 2015  | Île Suhm                  |
| Species | Specimen information (locality; test length; source; method) |
|---------|-------------------------------------------------------------|
| *Abatus cavernosus* (Philippi, 1845) | Kerguelen Islands, Indian Ocean; 27 mm; ZMB 5657; MRI Davis Station, Southern Ocean; 35 mm; unvouchered; dissection South Georgia, Southern Ocean; 38 mm; ZMH E7371; dissection Dundee Island, Southern Ocean; 45 mm; UB 2013.02.11.74; dissection |
| *Abatus cordatus* (Verrill, 1876) | Kerguelen Islands, Indian Ocean; 28 mm; ZSM 20011482; μCT Kerguelen Islands, Indian Ocean; 29 mm; ZMB 5437; MRI Kerguelen Islands, Indian Ocean; 33 mm; UB 2015.11.22.28; dissection Kerguelen Islands, Indian Ocean; 34 mm; NHMD; dissection Kerguelen Islands, Indian Ocean; 38, 41, 50 mm; UB 2015.12.8.42, 2015.11.22.28, 2015.11.22.28; dissection Kerguelen Islands, Indian Ocean; 42 mm; UB 2015.12.8.59; μCT + PTA Kerguelen Islands, Indian Ocean; four adults; UB 2015.11.28.14, 2015.11.28.18; NGS |
| *Brisaster antarcticus* (Döderlein, 1906) | Kerguelen Islands, Indian Ocean; 43 mm; NHMD; dissection Antipodes Island, Pacific Ocean; 51 mm; NIWA 114011; dissection Heard Island, Indian Ocean; 52 mm; AAD HIMS H117; dissection |
| *Brisaster capensis* (Studer, 1880) | Cape Peninsula, Atlantic Ocean; 48 mm; NHMD; dissection Cape Peninsula, Atlantic Ocean; 48 mm; ZMB 5855; dissection |
| *Brisaster fragilis* (Düben and Koren, 1844) | Tromsø, Atlantic Ocean; 25 mm; ZMB 2766; dissection Faroe Islands, Atlantic Ocean; 26 mm; NHMD; μCT Davis Strait, Atlantic Ocean; 30 mm; AM J.2630; dissection Bergen, Atlantic Ocean; 32 mm; ZSM 20011687; dissection Greenland, Atlantic Ocean; 45 mm; ZMH E7859; dissection Faroe Islands, Atlantic Ocean; 50 mm; NHMD; dissection |
| *Brisaster latifrons* (Agassiz, 1898) | Hokkaido, Pacific Ocean; 46 mm; MCZ 4407; dissection Sea of Japan, Pacific Ocean; 52 mm; UMUTZ Ecn-SI18-32; dissection Puget Sound, Pacific Ocean; 66 mm; unvouched; dissection |
| *Brisaster moseleyi* (Agassiz, 1881) | Isla de los Estados, Atlantic Ocean; 55 mm; unvouchered; dissection |
| *Brisaster owstoni* (Mortensen, 1950) | Sagami Bay, Pacific Ocean; 18, 23, 26, 28, 29 mm; MMBS Ec551, Ec659, 1888, 1908; 1921; dissection |
| *Brisaster townsendi* (Agassiz, 1898) | Long Beach, Pacific Ocean; 21, 22, 23 mm; ZMB 7440, 7441, 7442; μCT + PTA Long Beach, Pacific Ocean; 27, 28, 29, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 40, 41, 41, 42, 42, 42, 42, 42, 42, 42, 42, 44, 44, 44, 44, 45, 45, 46, 46, 46, 47, 47, 47, 48, 48, 48, 48, 49, 49, 49, 50, 50, 50, 50, 50, 51, 51, 51, 51, 51, 51, 53, 53, 54, 55, 55, 55, 56, 56, 57, 58, 58, 66 mm; unvouched; dissection San Luis Obispo, Pacific Ocean; 51 mm; CASIZ 92728; dissection Long Beach, Pacific Ocean; 42, 46, 47 mm TL; unvouched; TEM Long Beach, Pacific Ocean; three adults; unvouched; SEM Long Beach, Pacific Ocean; 42–56 mm TL; unvouched; NGS |
| *Genicopatagus affinis* (Agassiz, 1879) | Coronation Island, Southern Ocean; 51 mm; UB; dissection Coronation Island, Southern Ocean; 61 mm; UB; dissection |
| *Parapneustes cordatus* (Koecher, 1912) | Helment Bank, Southern Ocean; 40 mm; NHMD; dissection Lydian Island, Southern Ocean; 48 mm; UB; dissection |
| *Protenaster australis* (Gray, 1851) | Sydney Harbour, Pacific Ocean; 42 mm; unvouched; dissection |
| *Protenaster rostratus* (Smith, 1878) | Queensland, Pacific Ocean; 28 mm; QM GL271; μCT + PTA |
| *Pseudabatus nimrodi* (Koecher, 1911) | Davis Station, Southern Ocean; 40 mm; unvouched; dissection Davis Station, Southern Ocean; 52, 53 mm; unvouched; dissection Dumont D'Urville Station, Southern Ocean; 59 mm; UB; dissection Cape Armitage, Southern Ocean; 64 mm; SIO E3079; dissection |
| *Schizocosmus abatoides* (Clark, 1925) | Weddell Sea, Southern Ocean; 33 mm; ZMH E7342; dissection Dundee Island, Southern Ocean; 50 mm; UB; dissection |
| *Tripylaster philippii* (Gray, 1851) | Arica, Pacific Ocean; 33 mm; SIO E831; dissection Antarctica, Southern Ocean; 39 mm; NHMD; dissection Argentina, Atlantic Ocean; 44 mm; ZMH E7774; dissection Locality unknown; 57 mm; MCZ 3259; dissection Southern Argentina, Atlantic Ocean; 60 mm; ZMH E7775; dissection |
| *Tripylus excavatus* (Philippi, 1845) | Southern Argentina, Atlantic Ocean; 37 mm; ZMH E4181; μCT + PTA |
**Dissection and Photography**

In total, 68 specimens of *B. townsendi* (27–66 mm test length, TL) were vivisected and their internal organs observed using a SZX10 stereomicroscope (Olympus Scientific Solutions Inc., Waltham, MA, United States) with attached D5300 digital camera (Nikon Inc., Melville, NY, United States). Preparations were carried out by removing the aboral half of the test together with all gonadal tissue. The internal organization of each animal as well as color of the IC and its contents were surveyed. In addition, ten ethanol-fixed specimens of *A. cordatus* as well as specimens of further schizasterid species were dissected in order to document absence or presence as well as shape and size of the IC.

**Micro-Computed Tomography**

Three-dimensional (3D) analysis of internal organ systems was conducted using contrast-enhanced μCT. Three intact specimens of *B. townsendi* (21, 22, 23 mm TL) were formalin-fixed (4% in seawater) and, after several weeks of storage in fixative, were transferred to 70% ethanol through a graded series, and finally contrasted with 0.3% phosphotungstic acid (PTA) in 70% EtOH for 4 weeks (Ziegler et al., 2018; Ziegler, 2019). A single, ethanol-fixed specimen of *A. cordatus* (42 mm TL) as well as two formalin-fixed museum specimens (*Protenaster rostratus*, 28 mm TL and *Tripylus excavatus*, 37 mm TL) were treated in the same way. All contrast-enhanced samples were scanned using a SkyScan 1272 μCT system with a detector size of 4,032 × 3,280 px (Bruker microCT, Kontich, Belgium). In order to achieve higher isotropic voxel resolutions, a single intact IC from a formalin-fixed specimen of *B. townsendi* (22 mm TL) was excised together with adjacent digestive tract elements and scanned as well. For comparison, a single adult specimen each of *A. cordatus* (28 mm TL) and *Brisaster fragilis* (26 mm TL) was scanned using conventional μCT on a Phoenix Nanotom μCT system with a detector size of 2,304 × 2,304 px (GE Sensing & Inspection Technologies, Wunstorf, Germany). Reconstruction of the projection images was performed using the software provided with each scanner, i.e., NRecon 6.4 (Bruker microCT) and DatosX Reconstruction 1.5 (GE Sensing & Inspection Technologies). Visualization of two-dimensional (2D) and 3D data was carried out according to previously published protocols (Ziegler and Menze, 2013; Ziegler et al., 2018).

**Magnetic Resonance Imaging**

MRI data for one intact museum specimen each of *A. cavernosus* (27 mm TL) and *A. cordatus* (29 mm TL) previously obtained using a Pharmscan 70/16 MRI system (Bruker Biospin GmbH, Ettlingen, Germany) were integrated into the present study. Information on specimen preparation for MRI and scanning parameters has been published elsewhere (Ziegler and Mueller, 2011; Ziegler et al., 2014).

**Morphometric and Volumetric Analyses**

2D measurements (to nearest mm) of TL, width, and height as well as IC length and width (to the nearest 0.1 mm) were made using size-calibrated photographs gathered during specimen dissection. In several cases, test height was inferred using height-to-length ratios obtained through calibrated imagery of schizasterid test dimensions taken from the original species descriptions. Measurements were performed using Fiji/ImageJ 1.52s (Schindelin et al., 2012). Test volume (TV) was estimated by applying the geometric formula for an oblate spheroid, and IC volume (ICV) using the formula for a cylinder. The morphometric and volumetric values gathered using this approach were compared with 3D datasets obtained using MRI as well as μCT and were found to be congruent. The resulting data were plotted using Prism 8.3.1 (GraphPad Software Inc., San Diego, CA, United States).

**Light Microscopy**

During vivisection, ICs of selected larger specimens of *B. townsendi* were excised and their contents extracted using a plastic pipette. The liquid was placed on a glass slide, covered, and immediately observed using a BX51 light microscope (Olympus Scientific Solutions Inc.) with attached QIClick monochrome digital video camera (Teleyne QImaging Inc., Surrey, BC, Canada). A representative video was edited using Windows 10 Video-Editor software (Microsoft Corp., Redmond, CA, United States) and saved as an MPEG-4 file. In addition, semi-thin sections (1 μm) of excised ICs were obtained on an EM UC6 ultramicrotome (Leica Microsystems, Wetzlar, Germany) from epoxy resin-embedded samples originally prepared for TEM. The semi-thin sections were placed on slides, stained using toluidine blue, covered, and imaged using a BX51 light microscope with attached CC-12 digital camera (Olympus Soft Imaging Systems GmbH, Münster, Germany).

**Scanning Electron Microscopy**

For SEM, three adult specimens of *B. townsendi* were dissected immediately after collection while still on board the research vessel, and extracted ICs were fixed in 2.5% glutaraldehyde in Sorenson’s phosphate buffer at pH 7.2. Samples were held in fixative at 4°C for several days, then rinsed three times in the same buffer and postfixed in 1% OsO4 for 1 h. After the initial osmium treatment, they were rinsed once in buffer, once in buffer diluted to 50% with water, and once more in pure water. Tissues were then subjected to thio-carbohydrazide-mediated osmium binding (Kelley et al., 1975): they were immersed in a saturated solution of thio-carbohydrazide for 1 h, rinsed twice with water, then treated with 1% OsO4 in water for 1 h. Tissues were then rinsed twice in water, dehydrated through an ascending ethanol series (to 100%), critical-point-dried using a Samdri-PVT-3D system (Tousimis Research Corp., Rockville, MD, United States) with CO2 as the transitional fluid, and mounted on stubs using copper-conductive tape with conductive adhesive on both sides. Specimens were coated with gold/palladium using a Pelco SC-4 sputter-coater (Ted Pella Inc., Redding, CA, United States) and imaged with a FEI Quanta 200 scanning electron microscope (Thermo Fisher Scientific Inc., Waltham, MA, United States).

**Transmission Electron Microscopy**

For TEM, three specimens of *B. townsendi* (42, 46, 47 mm TL) were dissected immediately after collection while still on board the research vessel, and extracted ICs as well as pieces of intestinal wall were fixed in 2.5% glutaraldehyde in Sorenson’s phosphate buffer (pH 7.2). Fixed samples were held at 4°C for several days, then rinsed three times in buffer and postfixed in 1% OsO4 in water for 1 h. After the initial osmium treatment, the samples were rinsed once in buffer, then in pure water. Tissues were then subjected to thio-carbohydrazide-mediated osmium binding (Kelley et al., 1975): they were immersed in a saturated solution of thio-carbohydrazide for 1 h, rinsed twice with water, then treated with 1% OsO4 in water for 1 h. Tissues were then rinsed twice in water, dehydrated through an ascending ethanol series (to 100%), critical-point-dried using a Samdri-PVT-3D system (Tousimis Research Corp., Rockville, MD, United States) with CO2 as the transitional fluid, and mounted on stubs using copper-conductive tape with conductive adhesive on both sides. Specimens were then coated with gold/palladium using a Pelco SC-4 sputter-coater (Ted Pella Inc., Redding, CA, United States) and imaged with a FEI Quanta 200 scanning electron microscope (Thermo Fisher Scientific Inc., Waltham, MA, United States).
in 1% OsO₄ in buffer for 1 h. After osmium treatment, they were rinsed three times in buffer, dehydrated through an ethanol series of ascending concentrations (to 100%), transferred to propylene oxide, infiltrated, and then embedded in Araldite 502 epoxy resin (Electron Microscopy Sciences, Hatfield, PA, United States). Ultrathin sections (70 nm) were obtained with a diamond knife using an EM UC6 ultramicrotome (Leica Microsystems). Sections were mounted on Formvar-covered copper grids, stained with uranyl acetate and lead citrate, and examined using an EM 10 CR transmission electron microscope (Carl Zeiss AG, Oberkochen, Germany). Electron micrographs were digitally recorded from phosphor imaging plates using a Micron imaging plate scanner (DITABIS, Pforzheim, Germany). Composite overview images were created using the software AutoStitch (Brown and Lowe, 2007).

**Next Generation Sequencing**

For molecular inferences, ICs from 12 specimens of *B. townsendi* (42–56 mm TL) were extracted following dissection on the research vessel. The organs and their contents were each placed into sterile plastic vials stored on ice; within 3 h of collection they were transferred to -80°C for storage until analysis. For two of these specimens (43, 48 mm TL), small pieces of stomach, intestine, and rectum (and their contents) were also sampled and treated in the same way. ICs from *A. cordatus* were extracted from four adult specimens that had been directly fixed in 95% ethanol following collection. For all samples, nucleic acids were extracted using the FastDNA SPIN Kit for Soils (MP Biomedicals, Solon, OH, United States). The success of extractions was confirmed via gel electrophoresis (1% agarose) and quantified through comparison with the 1 kb exACTGene DNA ladder (Fisher Bioreagents, Pittsburgh, PA, United States).

Extracted samples were sequenced using a MiSeq NGS system with the Reagent Kit V2 (Illumina, San Diego, CA, United States) to generate paired-end sequences. The V4 variable regions of 16S rRNA genes were amplified using the barcoded primer pair 515F-806R (Apprill et al., 2015; Parada et al., 2016). Negative PCR controls were run during the sequencing reaction and none yielded positive results. The reaction mixture and thermocycling specifications were carried out in accordance with rules set forth by the Earth Microbiome Project (Caporaso et al., 2011). Analyses of microbiome data were carried out using QIIME 2 2016.8 (Bolyen et al., 2019). Raw sequence data were demultiplexed and quality-filtered using the *q2-demux* plugin followed by denoising with DADA2 via *q2-dada2* (Callahan et al., 2016), which included removing singleton sequences. The minimum sequence length found by DADA2 was 150 base pairs (bp), and the maximum was 254 bp, while the minimum threshold for quality control was 150 bp using a Phred quality score to represent confidence in assignment of the base call by the Illumina sequencer. Most sequences maintained a score above 30 until the 151 bp mark. Amplicon sequence variants (ASVs) were aligned with MAFFT via *q2-alignment* (Katoh et al., 2002). Aligned ASVs were then used in construction of a phylogeny using fasttree2 via *q2-phylogeny* (Price et al., 2010). Alpha diversity metrics in the form of a Shannon’s diversity index and operational taxonomic units (OTUs), beta diversity metrics in the form of Bray-Curtis and unweighted UniFrac dissimilarity (Lozupone and Knight, 2005), as well as Principal Coordinates Analysis (PCoA) were estimated via *q2-diversity* for each sample and visualized using the EMPeror software package (Vázquez-Baeza et al., 2013).

Taxonomy was assigned to the ASVs via *q2-feature-classifier* (Bokulich et al., 2018) using a naive Bayesian taxonomy classifier against the SILVA 97% database (Yilmaz et al., 2014). Kruskal-Wallis statistical comparisons for alpha diversity metrics and PERMANOVA beta diversity comparisons of unweighted UniFrac dissimilarity were made in QIIME 2 via *alpha-group-significance* and *beta-group-significance*, respectively. Differential abundance analysis was performed and visualized using a feature-table heatmap (Hunter, 2007) generated in QIIME 2 using the Bray-Curtis metric and UPGMA hierarchical clustering methods. As the initial numbers of taxa were highly complex, the visualizations were limited to the most abundant taxa (i.e., ≥ 2,000 OTUs for *B. townsendi* and ≥ 150 OTUs for *A. cordatus*) and the feature table was collapsed to level 5 taxonomy (i.e., family level) for *B. townsendi* and level 4 taxonomy (i.e., order level) for *A. cordatus*.

**RESULTS**

**Morphology of the Intestinal Caecum in *Brisaster townsendi* and *Abatus cordatus***

Adults of *B. townsendi* have two relatively short posterior petals and a deeply sunken petal in ambulacrum III (Figure 1a). The digestive tract shows the typical spatangoid form and is composed of a predominantly tubular gut that can be divided into esophagus, stomach, intestine, and rectum (Figure 1b). As in all other heart urchins, pharynx and Aristotle’s lantern are absent. The entire digestive tract is suspended from the relatively thin mesenteries. In addition to the four main gut compartments, the digestive tract also possesses a tubular primary and secondary siphon laterally connecting the posterior side of the stomach, a large and convoluted gastric caecum attached to the aboral side of the anterior part of the stomach, and a prominent IC. The IC is a kidney-shaped, almost cylindrical, blind-ending sac (Figure 1b) that is connected to the posterior part of the intestine through the short caecal canal (Figure 1c). The organ is located within a coil formed by the posterior part of the intestine and the rectum. In living specimens, the wall of the IC is semi-transparent and may bear dark (Figure 1c) or bright spots (Figure 1d), or no spots at all (Figure 1e). Seen through its semi-transparent wall, the contents of the organ either appear as a partially dark (Figures 1c,d) or bright mass (Figure 1e). In a subset of vivisected specimens (*N = 50*), 38 (76%) of the ICs had no, 6 (12%) had dark, and 6 (12%) had bright spots, while 31 (62%) had dark and 19 (38%) had bright contents.

In contrast to esophagus and stomach, the intestine and IC do not bear any large haemal ducts (Figures 1f,g). Due to its suspension from an inner mesentery that spans the posterior gut coil, the IC is surrounded by the coelomic fluid contained within the main perivisceral cavity, the somatocoele (Figures 1h,i).
The IC is present both in male and female specimens and no sexual dimorphism in its form was observed. A virtual, µCT-based horizontal section through the IC of B. townsendi at the level of the caecal canal shows that the organ’s contents are homogeneous, while the contents of stomach and intestine are heterogeneous (Figure 1j). The findings on the organization of the digestive tract in B. townsendi also apply to A. cordatus.

**Anatomy of the Intestinal Caecum in Brisaster townsendi**

A closer look at the gut contents reveals that the heterogeneous material observed within the esophagus, stomach, intestine, and rectum is primarily composed of sediment (Figure 2a), while the largely homogenous contents of the IC are organic, as can be inferred from their staining properties under contrast-enhanced µCT (Figures 2b,c). These observations also hold true in A. cordatus. At the anterior end of the IC, a centrally located, small lumen may be observed that bulges inward toward the center of the organ from the caecal canal (Figure 2d). While individual sediment grains inside the intestine can readily be discerned at 9.5 µm isotropic voxel resolution (Figures 2a,b), the contents of the IC cannot be differentiated in this as well as another scan acquired at 600 nm isotropic voxel resolution (Figure 2d). However, no sphincter or lip was found in the vicinity of the caecal canal that would prevent sediment from entering the IC.

The histology of the digestive tract wall follows the basic spatangoid pattern. From the gut lumen toward the somatocoel, an inner digestive epithelium is bordered by a middle connective tissue layer and finally an outer coelomic epithelium. In the intestine, the digestive epithelium is heavily folded (Figure 2e). The enterocytes of the digestive epithelium are simple columnar cells that measure about 30–40 µm in height and 5–10 µm in width. The connective tissue layer is well developed, ranging from 20 to 100 µm in thickness. Large parts of the extra-cellular matrix (ECM) are filled by a golden-brown granular material with intermittent coelomocytes and occasionally fibrocytes. In close proximity to the coelomic epithelium, the connective tissue layer contains a relatively wide haemal space of about 10–20 µm height. The coelomic epithelium itself exhibits a height of about 10–20 µm and is composed of a pseudostratified epithelium consisting of a basal layer of myoepithelial and a top layer of squamous epithelial cells each of about 5–10 µm in height and 5–10 µm in width. Seen from the somatocoelomic side, the outer wall of the intestine appears relatively smooth (Figure 1b).

The histological properties of the IC’s wall differ significantly from those of the intestine. The digestive epithelium is arranged in irregular bumps separated by depressions and which measure about 50 µm in height and 100 µm in
width (Figure 2f). As in the intestine, the enterocytes of the digestive epithelium are simple columnar cells, but here they measure about 70–90 μm in height and 5–10 μm in width. The connective tissue layer is weakly developed, with a thickness of 5–10 μm. The relatively small haemal space measures about 2–5 μm in height. The coelomic epithelium is characterized by a single layer of squamous cells with about 5–10 μm height and 5–10 μm width (Figure 2f). Seen from the somatocoelomic side, the outer wall of the IC appears entirely smooth (Figures 1c–e).

LM observations of the organic mass inside the IC obtained from living specimens show that the organ contains a microbial population composed of motile spirochetes each about 10 μm in length, as well as smaller spiral- and rod-shaped bacteria (Supplementary Video S1). In addition, ciliates can occasionally be observed (Figure 2g).
Ultrastructure of the Intestinal Caecum in *Brisaster townsendi*

SEM investigations reveal that the IC’s digestive epithelium is covered by a dense microbial population (Figure 2h). As also observed with LM (Figure 2g), the most obvious component of this bacterial mass are spirochetes that occur within the lumen of the IC and are not attached to the apical side of the digestive epithelium. Instead, the apices of the enterocytes are predominantly covered by smaller, rod-shaped bacteria (Figure 2i). The coelomic epithelial cells of the IC possess a collar of microvilli surrounding a single, long cilium (Figure 2j). In contrast to the digestive epithelial cells, the coelomic epithelial cells are not covered by a dense fringe of bacteria.

TEM analyses show that the digestive epithelium of the intestine is characterized by simple, non-glandular enterocytes that do not secrete mucous (Figure 3a). Apically, these non-ciliated enterocytes possess a dense zone of unbranched microvilli measuring about 1.75 × 0.15 µm, but no terminal web was observed. The non-muscular enterocytes join apically through upper adherens and lower tight junctions. In the apical region, numerous pinocytotic vesicles and electron-dense as well as clear vacuoles can be seen inside the cytoplasm. The centrally located nuclei are subspherical and measure about 4.5 × 3.5 µm. A Golgi apparatus is present lateral or apical to the nucleus, while located nuclei are subspherical and measure about 4.5 × 3.5 µm. However, no endocytosis of bacteria was observed. The centrally located nuclei are elliptical in shape and measure about 10 × 5 µm, their long axis oriented along the enterocytes’ main axis (Figure 3b). No prominent Golgi apparatus or RER cisternae are present. However, numerous glycogen granules and subspherical as well as elongated mitochondria are distributed throughout the enterocytes (Figures 3b,d–g). A well-developed epineurial basiepithelial nerve plexus can be found near the base of the digestive epithelium (Figure 3h). The enterocytes are strongly folded at their base, interdigitating through numerous protrusions with a basement membrane of about 3–4 µm thickness. The underlying ECM is characterized by a haemal space of about 2–10 µm thickness and is interspersed with coelomocytes and occasionally fibrocytes. A dense zone of circular and longitudinal striated collagen fibrils each measuring 40–80 nm in diameter borders the ECM and the basement membrane of the coelomic epithelium (Figure 3h). The coelomic epithelium of the IC rests upon a basement membrane of about 100–400 nm thickness. The relatively inconspicuous hyponeural basiepithelial nerve plexus is present between the cells of the coelomic epithelium (Figure 3h). The cells of the coelomic epithelium are podocytes, but only a small number of myofibrils can be observed in these cells. The podocytes possess subspherical nuclei measuring about 5 × 3 µm and scattered mitochondria as well as glycogen granules, while RER cisternae are few. The somata of the podocytes measure about 8 × 4.5 µm (Figure 3h). In cross-section, their pedicels have dimensions of 350–450 nm in height and 250–350 nm in width, while the foot processes measure 100–200 nm in height and 100–250 nm in width. The slit diaphragms between individual pedicels and foot processes are about 20–30 nm wide (Figure 3i). Similar to the intestine, a collar of microvilli surrounds a single cilium on each coelomic epithelial cell, as also observed with SEM (Figure 2j).

The Gut Microbiome of *Brisaster townsendi* and *Abatus cordatus*

LM, SEM, and TEM observations show that a variety of bacterial morphotypes occur inside the IC of *B. townsendi* (Figures 2g–i, 3d–g). These include straight or curved rod-shaped bacteria measuring from about 2.0 × 0.2 to 3.0 × 0.5 µm (Figures 4a–d), small spiral-shaped bacteria measuring from about 1.5 × 0.2 to 5.0 × 0.5 µm (Figures 4e–h), and larger spiral-shaped bacteria measuring about 9.5 × 0.4 µm (Figure 4i). No bacterial stacks, spores, capsules, or buds are observed, but occasionally small filamentous structures measuring about 1.0 × 0.1 µm can be seen (Figures 3e,g). These as well as the rod-shaped bacteria are usually found on or near the apical side of the IC’s digestive epithelium (Figures 3d–g), while the spiral-shaped bacteria are predominantly found free within the organ’s lumen (Figures 2g–i and Supplementary Video S1). The majority of organisms directly attached to the enterocytes are rod-shaped bacteria (Figure 4a), although spiral-shaped bacteria can occasionally be found attached to the digestive epithelium.
as well (Figure 4h). At the attachment sites of these two bacterial morphotypes to the enterocytes there is no sign of direct cytoplasmic interaction between host and bacterial cells (Figures 3e, 4a,h). Additional bacterial forms include curved rod-shaped bacteria with (Figure 4b) or without (Figure 4c) a dense cover of fimbriae as well as spiral-shaped taxa without any apparent fimbriae (Figures 4e–i). Some of the rod- as well as the spiral-shaped morphotypes have at least one polar flagellum (Figure 4c). In addition, some bacteria show clear inclusion bodies (Figures 4b,e,g). More frequently, glycogen granules are seen between the dense fringe of bacteria adhering to the enterocytes’ apex (Figure 3d) and sometimes these granules are also observed inside bacteria (Figure 4d). Where the IC lumen contains a substantial amount of glycogen granules, the adjacent digestive epithelium appears to show increased pinocytotic activity (Figure 3d).

To complement these structural findings, a total of 855,592 16S rRNA gene sequences were generated from four regions of the digestive tract (stomach, intestine, IC, and rectum) in B. townsendi as well as from the IC in A. cordatus, representing 2,738 OTUs across all samples. The overwhelming majority of OTUs in all gut compartments were identified as bacterial, with only a single nanoarchaeal lineage; this latter OTU was observed in all four A. cordatus IC samples, but not in any B. townsendi samples (Supplementary Tables S1, S2). The OTUs in B. townsendi and A. cordatus IC samples showed limited overlap at the 97% similarity cut-off, which roughly corresponds to species level. However, when viewed at a higher taxonomic level, there was more similarity as > 50% of the OTUs in both species’ IC microbiomes were members of the Bacteroidetes, sulfate-reducing Deltaproteobacteria lineages, and Spirochaetes. The most abundant taxon in most of the ICs of
**FIGURE 4** | Ultrastructural and genomic analyses of the microbial population found inside the intestinal caecum (IC) as well as selected gut compartments of *Brisaster townsendi* and *Abatus cordatus*. (a–i) TEM and SEM micrographs showing different bacterial morphotypes found inside the IC of *B. townsendi*. (j,k) Heatmaps depicting co-occurrence of the most abundant operational taxonomic units (OTUs) across four gut compartments with a minimum OTU frequency of 2,000 in *B. townsendi* (j) and from ICs with a minimum OTU frequency of 150 in *A. cordatus* (k); colored scales show relative abundance of taxa across all samples, clusters on the left depict Bray-Curtis metric across samples and on the top depict UPGMA hierarchical clustering. SILVA taxonomic identifications were collapsed to family level (j) and order level (k) or higher if taxonomic classification to those levels was not possible. fi, fimbria; fl, flagellum; gg, glycogen granule; ib, inclusion body; ic, intestinal caecum; in, intestine; re, rectum; st, stomach.

*B. townsendi* was most closely related to a yet uncultured member of the Bacteroidetes, the VC2.1 Bacc22 group (Supplementary Table S1). The most abundant taxon in ICs of *A. cordatus* was an unknown species of Gracilibacteria (Supplementary Table S2), which was present, but at lower abundance, in ICs of *B. townsendi* as well.

The most abundant OTUs (i.e., ≥ 2,000 representatives) in the *B. townsendi* IC samples showed little to no overlap with the other gut compartments. This resulted in clear splits in the hierarchical clustering (Figure 4j), clearly distinguishing taxa found in the IC samples of *B. townsendi* from those in the other three gut compartments of that species. Only a single OTU, identified as a yet unknown species of *Mesorhizobium* (Supplementary Table S1), showed meaningful overlap between the IC and the other gut compartments (14% frequency in the IC vs. 86% frequency in the rest of the gut).
While the total diversity of IC microbiomes in both species was primarily composed of Gram-negative, rod- or spiral-shaped, motile, obligately anaerobic taxa (e.g., *Desulfovibrio* sp., *Sulfurimonas*, *Sediminispirochaeta*), members of the bacterial candidate phylum Gracilibacteria were also present, often in high numbers of amplicons. The relatively large sample size of ICs from *B. townsendi* (N = 12) and *A. cordatus* (N = 4) enabled us to examine intraspecific variation among IC microbial communities as well: substantial intraspecific variation was observed in both schizasterid species (Figures 4j, k and Supplementary Tables S1, S2).

We also noticed major differences in community structure of the IC microbiome compared to other gut compartments. The IC microbiomes of *B. townsendi* and *A. cordatus* displayed much lower overall Shannon’s diversity indices than did the microbiomes of stomach, intestine, and rectum of *B. townsendi* (Figure 5A). The *B. townsendi* IC microbial communities were much less diverse than those of the other three *B. townsendi* gut compartments (p < 0.03 for all pairwise comparisons), and were relatively similar among all twelve sampled individuals. Similar patterns were observed for the number of OTUs, with an average of 69.8 (SD 30.9) in the *A. cordatus* and 61.9 (SD 12.1) in the *B. townsendi* IC samples compared to 660 (SD 38.2) in the stomach, 857.5 (SD 7.8) in the intestine, and 797.5 (SD 20.5) in the rectum of *B. townsendi*. Beta diversity comparisons confirmed that microbial species composition varied significantly across these five sets of samples [PERMANOVA, 999 permutations, pseudo-\(\text{F}(4, 17) = 8.782, p = 0.001\)]. In addition, pair-wise comparisons showed that beta diversity of the *B. townsendi* IC samples was significantly different from all other digestive tract compartments as well as from the *A. cordatus* IC samples (q < 0.05 for all). No significant differences were observed in comparisons between each of the three main gut compartments, i.e., stomach, intestine, and rectum (q > 0.3 for all).

PCoA analyses using unweighted UniFrac and Bray-Curtis datasets were visualized in the first three dimensions. Both plots revealed similarly distinct separation among sample types, so only the UniFrac results are shown here (Figure 5B). Axis 1 explained roughly half (47.0%) of the variability in the data and clearly distinguished the two species’ IC microbiomes from the other gut compartments of *B. townsendi*. The second axis (12.4%) distinguished the IC samples from *A. cordatus* and *B. townsendi*, while the third axis (9.1%) highlighted differences among stomach, intestine, and rectum samples, which were mostly overlapping. In addition, two main clusters were observed within the IC samples of *B. townsendi* (Figure 5B), further highlighting the significant variability among these samples despite their identical provenance.

**Distribution, Shape, and Volume of the Intestinal Caecum in the Schizasteridae**

Building on a study of several in- and outgroup taxa analyzed with regard to absence or presence of the IC (Ziegler, 2014), we examined additional species to provide a more comprehensive overview of the distribution of the IC among the Schizasteridae (Table 1). The IC is absent in the two extant species of the plesiomorphic schizasterid *Prosteraster* (Figure 6a). An IC can now be confirmed to occur in all extant species of *Brisaster*, including *B. moseleyi* (Figure 6b), *B. owstoni* (Figure 6c), and *B. townsendi* (Figures 1, 6d). The IC is absent in the only extant species of *Genicopatagus*, *G. affinis* (Figure 6e), but present in *Pseudabatus nimrodi* (Figure 6f). In addition, the organ is absent in *Schizocosmus abatoides* (Figure 6g), but present in the single extant species of *Tripylus*, *T. excavatus* (Figure 6h). Where present, the IC is always found to be suspended from the mesentery spanning the posterior gut coil. Based on contrast-enhanced CT data, the contents of the IC were found to be homogenous in *A. cordatus* and *T. excavatus*, as also observed for *B. townsendi* (Figures 1j, 2b–d).

In *Brisaster*, the IC is generally present in the form of a largely cylindrical, kidney-shaped structure (Figures 1f–i). This shape can be observed both in fresh as well as in formalin- or ethanol-fixed specimens (Figures 1b, 6d). While the IC takes this form in most species known to possess the organ, a few taxa differ in this regard. For example, the IC in *T. excavatus* is more rectangular in shape (Figure 6h), while a significant deviation from the general morphological pattern observed in *Brisaster* can be observed in...
**FIGURE 6** | Absence and presence of the intestinal caecum (IC) in selected schizasterid taxa; all images show results obtained using fixed specimens. **(a)** Protenaster australis, oral view. **(b)** Brisaster moseleyi, aboral view. **(c)** Brisaster owstoni, aboral view. **(d)** Brisaster townsendi, aboral view. **(e)** Genicopatagus affinis, aboral view. **(f)** Pseudabatus nimrodi, aboral view. **(g)** Schizocosmus abatoides, aboral view. **(h)** Tripylus excavatus, aboral view of a sliced volume rendering obtained using a contrast-enhanced µCT dataset with 10 µm isotropic voxel resolution. gc, gastric caecum; ic, intestinal caecum; in, intestine; re, rectum.

*P. nimrodi*, where the organ is present as a reduced, elongated structure (Figure 6f).

Data on the ICV for specimens of *B. townsendi* ranging from 22 to 67 mm TL reveal an exponential increase in ICV as a function of TL, and a linear increase in the percentage of TV occupied by the IC as a function of TL (Figure 7A). Calculations of ICV and TV among different species show that on average the ICV forms about 0.5–1% of TV (Figure 7B). As can be expected from the morphological findings, the ICV is much smaller as a function of TV in *P. nimrodi* (0.025%) and much larger in *T. excavatus* (1.5%), while on average few differences in ICV are observed among species of *Brisaster* and *Abatus* (Supplementary Table S3). For example, the ICV of *B. townsendi* and *A. cordatus* does not differ much in animals of comparable size: 14 µl (23 mm TL) vs. 14 µl (29 mm TL), 103 µl (42 mm TL) vs. 121 µl (42 mm TL), and 584 µl (66 mm TL) vs. 593 µl (50 mm TL), respectively.

**DISCUSSION**

Due to their deposit-feeding habits and frequent high densities, schizasterid heart urchins can have significant effects on the ocean floor and on biogeochemical fluxes between sediment and overlying water (Chesher, 1963; Brownell, 1970; Thompson et al., 1987; Penry and Jumars, 1987; Plante and Jumars, 1992; De Ridder, 1994; Bromley et al., 1995; Schatt and Féral, 1996; Mayer et al., 1997), the IC was only recently shown to be an organ unique to schizasterids (Ziegler, 2014). Here, we provide the first characterization of the schizasterid IC from macroscopic to ultrastructural levels based on observations of dozens of specimens of *B. townsendi*. These novel observations reveal a number of unique structural features of the organ.

To provide further insights into the microorganisms inhabiting the schizasterid digestive tract, we undertook a multimodal analysis of the gut of *Brisaster townsendi*. Members of the genus *Brisaster* are particularly suitable for study because of their established taxonomy, broad geographic distribution, and well-documented lifestyle (Mortensen, 1951; Ghioi, 1988-89; Kanazawa, 1992; Hood and Mooi, 1998; Walker and Gagnon, 2014). Most importantly, all species of *Brisaster* are characterized by the presence of the IC (Ziegler, 2014, this study), an enigmatic organ potentially harboring a distinct gut microbiome of unknown capacity.

**The Schizasterid Intestinal Caecum, a Highly Specialized Gut Compartment**

While the presence of a hindgut diverticulum in selected species of *Brisaster* and *Abatus* had been known for some time (Brownell, 1970; De Ridder et al., 1985; Penry and Jumars, 1987; Plante and Jumars, 1992; De Ridder, 1994; Bromley et al., 1995; Schatt and Féral, 1996; Mayer et al., 1997), the IC was only recently shown to be an organ unique to schizasterids (Ziegler, 2014). Here, we provide the first characterization of the schizasterid IC from macroscopic to ultrastructural levels based on observations of dozens of specimens of *B. townsendi*. These novel observations reveal a number of unique structural features of the organ.

The internal anatomy of both schizasterid species described in the present study is highly similar, and akin to their previously studied congener *B. latifrons* and *A. cavernosus*...
At the histological level, most of the digestive tract of *B. townsendi* resembles that of other heart urchins (Koehler, 1883; Hamann, 1887; Anisimov, 1982; De Ridder and Jangoux, 1993; Holland, 2013). Despite the absence of haemal ducts in both intestine and IC, our observations suggest that the small haemal space within the wall of the IC is continuous with the enlarged haemal space of the intestine via the tissues comprising the caecal canal and mesentery suspending the organ (Figures 1f–i, 2d), thus permitting an exchange of haemal fluid between intestinal and IC walls.

At the ultrastructural level, the intestinal tissues of *B. townsendi* resemble those of the distantly related heart urchin *Echinocardium cordatum* (De Ridder and Jangoux, 1993). As in *E. cordatum*, the intestinal ECM of *B. townsendi* is characterized by the presence of electron-dense granules (Figure 3a). These particles have previously been identified as a ferric phosphate precipitate caused by oxidative deposition under reducing conditions (Buchanan et al., 1980). In addition, the digestive epithelium of the intestine in both spatangoid species is characterized by the presence of a dense cover of microvilli. In contrast, the IC’s enterocytes possess only a small number of tiny microvilli (Figure 3c). This sparse cover of microvilli in the IC is likely related to the dense bacterial population attached to the organ’s digestive epithelium (Figure 3d). A further difference between intestinal and IC walls is the strong basal folding of the IC’s enterocytes (Figure 3b), indicative of an increased exchange between these cells and the underlying ECM (De Ridder and Jangoux, 1993). Furthermore, the presence of a pseudo-stratified coelomic epithelium in the intestine (Figure 3a) as opposed to a mono-layered coelomic epithelium in the IC (Figure 3b) suggests the possibility that these two tissues experience different mechanical loads. However, arguably the most unusual structural feature of the IC is the fact that the organ is lined by podocytes on its exterior, coelomic side (Figure 3h). The presence of this cell type is usually indicative of ultrafiltration mediated by a muscle-generated pressure differential (Ruppert, 1994).

However, in comparison to the intestine, the IC features only a small number of myoepithelial cells. If exchange does indeed occur across this barrier, it could be caused by pressure generated via the contraction of myoepithelial cells elsewhere in the digestive or mesenterial systems, or through diffusion.

In most marine deposit feeders, the main contents of the digestive tract are sediment particles. This is true for *B. townsendi* as well, as previously shown for its congener *B. fragilis* (Eichelbaum, 1910). However, the IC again is strikingly different from the rest of the digestive tract in this respect: although it is directly connected to the sediment-packed intestine by the caecal canal, we observed no sediment particles inside the ICs of *B. townsendi* and other schizasterids. Instead, the organ was filled with an organic mass composed of diverse microorganisms (Figure 2g).

**Composition of the Microbiome of the Schizasterid Intestinal Caecum**

An increasing number of studies are beginning to shed light on the gut microbiomes found in various echinoderm taxa, among them several deposit-feeding sea cucumber (Gao et al., 2014; Yamazaki et al., 2016; Wang et al., 2018; Pagán-Jiménez et al., 2019; Weigel, 2020) and sea urchin species (Thorsen, 1999; Thorsen et al., 2003; da Silva et al., 2006; Schwob et al., 2020). However, most of these analyses focus on microbial organisms found in the main digestive tract, while the present study provides data permitting a comparison between the microbiomes found inside the main gut compartments and a specialized digestive tract derivative.

Our genomic analyses show that the IC of *B. townsendi* contains a community of predominantly anaerobic bacteria with Bacteroidetes averaging 34.6% of the microbiome, in addition to Desulfobacterales (16.0%) and Spirochaetales (9.2%). Similar results were obtained for *A. cordatus*, with Desulfobacterales averaging 31.5% of its microbiome, followed by Bacteroidetes (14.0%) and Spirochaetales (9.5%), with the addition of
Rhizobiales (9.5%) and Lentisphaerales (7.0%). The most common phylotypes in A. cordatus were closely related to a recently described genus of the Desulfovacterae thus far known from a single psychrophilic species, Desulfoconvexus algidum, isolated from a fjord in the Svalbard archipelago (Konneke et al., 2013) and members of the bacterial candidate phylum Gracilibacteria, which recent genomic analyses suggest are often symbiotic, displaying limited metabolic capacity (Sieber et al., 2019).

For B. townsendi, the IC microbiome was found to be significantly different in composition and diversity from that of the rest of the gut (Figure 5A). Members of the sulfate-reducing Desulfovacterae were very abundant in the IC, in contrast to the intestine and rectum, where members of the Desulfolubacterae and Desulfuromonadaceae were prominent (Supplementary Table S1). Desulfovacterae are commonly found in marine sediments and are known to fully oxidize fermentative decomposition products such as acetate (Dhillon et al., 2003; Leloup et al., 2007; Jackson et al., 2014; Na et al., 2015). Another sulfate-reducing lineage abundant in the IC libraries were members of the genus Desulfoarba, a taxon within Desulfarticales recently discovered from a coal bed in central North America (An and Picardal, 2014). OTUs closely related to another uncultured group of Deltaproteobacteria, FW113, whose metabolic function is unknown, were also abundant. This group has also been found in anaerobic marine sediments and microbial mats (Li et al., 2006; Harris et al., 2013).

Confirming our microscopic results, OTUs most closely related to spirochetes were also very abundant in the ICs of B. townsendi and A. cordatus, but were not recovered in any of the other gut compartments. In contrast, a recent study found spirochetes primarily of the genus Spirochaeta in abundance in the main gut tissue of the schizasterid Abatus agassizii, and identified these taxa as part of its gut microbiome (Schwob et al., 2020). However, this particular species of Abatus does not possess an IC (Ziegler, 2014). In fact, the spirochete lineages found in the ICs of B. townsendi and A. cordatus differ significantly from those in A. agassizii, being predominantly related to the cultured taxon Sediminispirochaeta and the uncultured Spirochaeta group 2 clade (Supplementary Tables S1, S2). Sediminispirochaeta is a recently reclassified genus (Shivani et al., 2016) composed of three type strains isolated from microbial mats and oil fields (Fracek and Stolz, 1985; Magot et al., 1997; Dubinin et al., 2015). Spirochetes are known to form symbiotic associations with diverse metazoan taxa including cnidarians, arthropods, mollusks, echinoderms, and mammals (Paster and Canale, 1982; Margulis and Hinkle, 2006; Lawler et al., 2016; Van de Water et al., 2016; Jackson et al., 2018). Perhaps the best-studied example is the termite hindgut symbiosis, which includes free-living spirochetes in the gut fluids as well as spirochetes living on and in symbiotic flagellated protists (Smith and Arnett, 1974; Leadbetter et al., 1999; Iida et al., 2000; Graber and Breznak, 2004; Dröge et al., 2006, 2008; Brune and Dietrich, 2015). This scenario is similar to what was here observed in the schizasterid IC, where some spiral-shaped bacteria were attached to the IC’s digestive epithelium (Figure 4h) and others were found freely floating in the lumen of the organ (Figure 2g). In termites, spirochetes are known to facilitate the digestion of lignocellulose, providing metabolizable, small molecules to the host (Breznak and Brune, 1994). However, pure culture studies have shown that these spirochetes also participate in CO2-reducing acetogenesis (Leadbetter et al., 1999), thus providing a different pathway of carbon delivery to the host. Both endo- and ectosymbiotic members of the Bacteroidetes have additionally been identified in these termite gut associations (Wenzel et al., 2003; Noda et al., 2005). In the present study, members of the latter bacterial taxon were also found in the IC. For example, relatives of the genus Dracinibacterium, previously cultured from marine sediments in East Asia (Du et al., 2014; Gwak et al., 2015), were abundant in the organ. However, the most prominent OTU inside the IC was related to the uncultured VC2.1 Bac22 group, first described from a hydrothermal vent on the Mid-Atlantic Ridge (Reysenbach et al., 2000), but since found to be common in marine habitats as well as in association with various metazoan taxa such as annelids and cetaceans (Alain et al., 2002; Bik et al., 2016).

Finally, relatively low numbers of an OTU closely related to Sulfurimonas, a sulfur-oxidizing lineage of the Thiovulaceae (Epsilonproteobacteria), were observed in eight of the twelve (66.7%) IC samples from B. townsendi, but were particularly abundant in one sample (Supplementary Table S1). In addition to oxidizing reduced sulfur compounds, which would be in abundance in the IC due to the presence of sulfate-reducing bacteria, members of this group are known for reducing nitrate (Han and Pernar, 2015). However, these organisms require at least a partially oxic environment for their metabolism. While we did not measure oxygen levels in this study, low concentrations of O2 have previously been detected in the IC of B. latifrons (Plante and Jumars, 1992).

The strong and consistent distinction between the microbial communities found in the IC vs. other gut compartments (Figure 5B) suggests selection for specific microbial taxa in the IC. While the exact mechanisms of colonization or selection in the IC are yet to be determined, the bacteria in many animal-microbe symbioses are initially acquired from the environment and seawater diet of the host (De Ridder and Foret, 2001). As the initial microbial community passes through the gut, the transient microbiome may change due to the interactions of microbes with gut contents or their host as well as the conditions inside the digestive tract. Recent studies have suggested that in the digestive tract of the schizasterid A. agassizii (Schwob et al., 2020). However, in many marine deposit feeders that lack hindgut diverticula, most of this modified microbial community is likely washed out once the sediment reaches the anus. In turn, the IC found in B. townsendi is a discrete organ separate from the main flow of sediment, so that its inhabitants are protected from washout. We hypothesize that once the IC forms during ontogeny, it becomes initially seeded with bacteria from the host’s diet, and then its microbial community rapidly diverges in composition from that of the rest of the gut. Future studies using naıve animals would be important to address this topic, but conditions suitable for laboratory cultivation of B. townsendi have not been achieved thus far.
Function of the Microbiome of the Schizasterid Intestinal Caecum

The IC microbiome of *B. townsendi* is both consistent among individuals (although not completely uniform, suggesting some host or environmental variability) and strikingly different in composition and diversity from other gut compartments. These characteristics suggest that it has a distinct role in its host’s biology. However, some possible functions of the IC microbiome may be ruled out already on the basis of its position along the digestive tract, i.e., at the posterior end of the intestine, just anterior to the relatively short rectum. For example, it seems improbable that the IC is of importance in detoxifying gut contents, which are about to be shed from the body as feces via the anus. The organ also is unlikely to be involved in stockpiling beneficial microbes that could be used to inoculate the rest of the gut, for example to protect against the development of communities of pathogenic microbes or to facilitate digestive processes in other gut compartments.

Instead, the position of the IC is more consistent with a role in nutrition. It has previously been argued that conditions in the hindgut of marine deposit feeders are likely beneficial for bacteria (Plante et al., 1990), and that microbial activity in the posterior digestive tract should come at no or little cost to the host. If activities of such a hindgut microbial community produce molecules of use to the host that can be taken up across the gut wall, then selection might lead to the evolution of intestinal pouches to increase surface area for product uptake. Incidentally, the residence time of the microbial communities in such diverticula can be high, allowing them to become distinct from the microbiomes in gut compartments with only a short residence time. All of our data on the position, structure, and microbiome of the IC are consistent with this general scenario. What products of use to the host might then be generated by the schizasterid IC’s microbiome? We saw no evidence for phagocytosis of microbes by enterocytes of the organ, so it seems unlikely that *B. townsendi* is gardening bacteria for direct consumption. Instead, we hypothesize that the schizasterid IC’s microbiome converts dissolved organic compounds from the intestinal fluid into smaller, more easily metabolizable molecules that can then be used by the host. Intestinal fluid may be moved into the IC via pressure generated by muscular contraction in other gut compartments (for example, by peristalsis, which has been observed in spatangoids: De Ridder et al., 1985), by cilia near the entrance of the IC, or by diffusion. Dissolved organic molecules in the inbound liquid may fuel microbial activities inside the IC, much as occurs in the termite hindgut (Leadbetter et al., 1999). Products of these activities might be returned to the host in several ways. Small molecules like fatty acids might diffuse across the IC wall into the somatocoel or they might be taken up by IC enterocytes through pinocytosis and then transported into the haemal space of the organ. Once inside the haemal lacunae, such products could then be circulated directly to other parts of the body or be ultrafiltered from the IC’s haemal space across the podocytes lining the exterior of the IC.

A better understanding of these processes requires a detailed understanding of the organic substrates provided to the IC, and especially of the metabolic interactions of the diverse microbes in this specific gut compartment. At this point we can only speculate about these interactions. Although autotrophic members of the Desulfovibacteriaceae and Spirochaetales are known, it is more likely that they are functioning heterotrophically in the present context, as most described species of these groups as well as the Bacteroidetes are chemoorganotrophic carbon degraders and are most likely involved in fermentation of complex carbon substrates. The ensuing simpler molecules such as short-chained fatty acids might then serve as carbon source to the sulfate-reducing bacteria. This is reminiscent of a recently described association between a marine sediment spirochete and a sulfate-reducing *Desulfovibrio* strain that were studied in co-culture, and where the spirochete produces fermentation products for the sulfate reducer (Shivani et al., 2016). However, in echinoderms active sulfate reduction can lead to toxicity for the host in these types of symbioses (De Ridder and Forêt, 2001; De Ridder and Sauède, 2020; Schwob et al., 2020), but this may be ameliorated by sulfur-oxidizing bacteria observed in the IC that remove sulfide from the organ. Tests of these hypotheses will require a comprehensive functional characterization of the IC’s microbial community.

The present data show that each schizasterid IC contains a complex microbial consortium composed of taxa with overlapping and complementary metabolic activities. In this regard, the organ resembles another spatangoid hindgut structure, the recto-intestinal caecum (RIC), which is present in a distantly related group of heart urchins (De Ridder, 1994; De Ridder and Sauède, 2020). Morphologically and structurally, this organ differs from the IC (Ziegler, 2014) and in *E. cordatum* typically contains several small nodules predominantly composed of Deltaproteobacteria, Bacteroidetes, and Firmicutes (61.5, 23.1, and 15.4%, respectively; da Silva et al., 2006). However, the sulfur-oxidizing bacteria found inside the RIC were shown to be filamentous bacteria that form layers surrounding a detrital particle covered with sulfate-reducing microbes (De Ridder and Brignon, 2003). The bacteria on the outside of these nodules oxidize thiosulfate and sulfide originating from its core, thus providing the host with acetate and propionate (Temara et al., 1991, 1993; Brignon and De Ridder, 1998; Thorsen, 1998). In total, the processes inside the RIC contribute about 10% to the energy budget of the animal (Thorsen, 1999; Thorsen et al., 2003). The oxygen required to support part of this symbiosis is supplied through the somatocoel (Thorsen, 1999). Though neither the nodules nor the specific microbial taxa involved in this symbiosis are present in the schizasterid IC, similarly complex metabolic interactions are likely to have evolved among IC microbial residents. Whether the ciliates sometimes observed in the organ also play a role in these processes remains unclear at present.

**Evolutionary and Ecological Considerations**

Previous work has shown that the IC is absent in most extant higher schizasterid taxa as well as in more distantly related spatangoids (Ziegler, 2014). Thus, it seems probable that the IC evolved within a subclade of the Schizasteridae. However,
until now, taxon sampling was limited, making inferences on the evolutionary history of the IC difficult.

With the addition of observations reported in this study, a clearer picture emerges (Figure 8). The IC is present in all species of Brisaster, Tripylaster, Tripylus, and Pseudabatus—note that P. nimrodi has previously been placed in the genus Abatus, but is here tentatively re-assigned to Pseudabatus based on two recent phylogenetic analyses (Van Oosterom, 2012; Stara et al., 2016). However, the presence of the IC varies in the genus Abatus: the organ has been observed in the two species analyzed here (Table 1), but it is absent in four other species (Ziegler, 2014). In addition, the absence of an IC is here confirmed for the species Parapneustes cordatus, which in a previous study had been assigned to Tripylus (Ziegler, 2014). Furthermore, the IC is not present in any of the other extant schizasterid genera analyzed so far (Ziegler, 2014). An examination of the distribution of the IC in a phylogenetic context suggests that this structure may have arisen only once, in the common ancestor of a group of at least nine schizasterid genera including Brisaster, Tripylus, and Pseudabatus (Figure 8). If this hypothesis is correct, then the IC has been lost on several occasions: at least once within the genus Abatus, and more than once in the clade comprising Parapneustes and Genicopatagus as well as other genera (Figure 8). An alternative hypothesis, of course, is that the IC has originated more than once.

We note that there is variation in the form and relative size of the IC among the schizasterid species that possess the organ. While in most species, including B. townsendi and A. cordatus, it is a slightly curved cylinder constituting about 0.5–1.5% of TV, it is bean-shaped and shorter in A. cavernosus (Ziegler, 2014), more rectangular in T. excavatus, and reduced in P. nimrodi.

This variation in shape and size may be due to diversification.

![Figure 8](image-url)
after a single origin, or due to more than one independent origin of the IC. Though the latter scenario does not seem particularly parsimonious given the strong similarity in overall shape and location of the IC, it is worth noting that barriers to the evolution of digestive appendices may be low (Smith et al., 2017). Distinguishing among the different evolutionary hypotheses will require both an improved understanding of schizasterid phylogeny (David et al., 2005, 2016; Saucède et al., 2013), as well as further study of the distribution and characteristics of the IC within the clade.

Although the systematics of schizasterids are not yet fully resolved, the available data nonetheless allow us to infer the minimum age of the IC. Spatangoids are a relatively ancient sea urchin lineage that originated in the Mesozoic, more precisely the Early Cretaceous, approximately 135 mya (Kroh and Smith, 2010). About 40 million years later, the first schizasterids (members of the fossil genus Periaster) appeared during the early phase of the Late Cretaceous (Figure 8). However, the two oldest schizasterid taxa known to possess the IC, i.e., Brisaster and Tripylus, originated in the final phase of the Late Cretaceous, about 70–66 mya (Forbes, 1845; Lambert, 1933; Markov, 1994; Kroh and Smith, 2010). It is therefore very likely that at least in these two genera, the IC (and by inference, it’s microbiome) evolved prior to the Cretaceous-Paleogene asteroid impact that occurred about 65.5 mya. This cataclysmic event caused a significant reduction in metazoan diversity both in marine and terrestrial habitats (Renne et al., 2013), although various marine taxa, including spatangoids, apparently were not as strongly affected by the ensuing mass extinction (Smith and Jeffery, 1998; Ribeiro et al., 2011; Lowery et al., 2018). A better understanding of the physiological benefit of the IC’s digestive symbiosis might help to elucidate whether the presence of the organ in selected schizasterid taxa constituted an evolutionary advantage in a post-extinction ocean with reduced oxygen and nutrient levels (Henehan et al., 2019).

While schizasterids represent a relatively old spatangoid lineage, they nonetheless share with other heart urchins a specific infaunal lifestyle (Nichols, 1959; Gibbs, 1963; Bromley and Asgaard, 1975; Kanazawa, 1992). Correspondingly, ecological data show that schizasterids are frequently found in close association with other spatangoid taxa (McCauley and Carey, 1967; Lie and Kisker, 1970; Nichols, 1975, 2003; Thompson et al., 1987, 1993; Allen et al., 1999; Stull et al., 2001; Widdicombe et al., 2004; Macdonald et al., 2012). However, the physical and chemical parameters of the surrounding sediment may change significantly over short distances and slight depths, leading to zonation effects that provide heart urchins with potentially narrow ecological niches (Gibbs, 1963; Bromley et al., 1995; Enderlein, 2004). For example, in contrast to spatangoid taxa such as Echinocardium, Brissus, or Brissopsis that bury down to 10–20 cm depth (Kanazawa, 1992), species of Brisaster have been reported to be shallow burrowers that only dig down to 1–5 cm below the sediment surface (Gibbs, 1963; Brownell, 1970; Kanazawa, 1992). These observations could indicate that the schizasterid IC can only be found in species that inhabit a specific zone below the sediment surface where conditions of the surrounding sediment are suitable to support the IC’s digestive symbiosis. In fact, the reduction or even complete absence of the organ in schizasterids such as Pseudabatus, Amphipneustes, or Brachysternaster as well as most species of Abatus (Ziegler, 2014) could be related to a change in lifestyle, as these taxa have been observed to be secondarily semi-infaunal or even entirely epifaunal organisms (Larrain, 1985; Pearse and McClintock, 1990; David et al., 2005; Thompson and Riddle, 2005; Sumida et al., 2008; Schwob et al., 2020). A test for this hypothesized change in lifestyle would be a comparison of denuded tests of the different schizasterid species, as shape of the anterior ambulacrum correlates with burrowing depth and sediment structure (Mortensen, 1951; Chesher, 1963; Kanazawa, 1992; Bromley et al., 1995). In addition to providing a more comprehensive picture of the role of the IC and its microbiome, such data might aid our understanding of the notable evolutionary success of schizasterids in the Southern Ocean (Poulin et al., 2002; Schwob et al., 2020).

CONCLUSION
Our data suggest that the microbial inhabitants of the IC play a distinct, if not yet fully understood, nutritional role in the biology of their hosts, and in at least some schizasterid lineages have done so since the Mesozoic. Understanding exactly how the schizasterid IC’s microbiome interacts with the host organism will require further study of the constituent microorganisms and of the integrated holobiont. Functional metagenomics, cultivation, and physiological investigations of selected microbial lineages, paired with tracer studies of material transport from the IC to the host, are likely to be particularly effective approaches with which to address these open questions.

DATA AVAILABILITY STATEMENT
Raw and derived image data are available from the MorphoBank online repository (Ziegler et al., 2020). These data include two 3D MRI scans, two conventional and four contrast-enhanced 3D µCT scans, as well as 37 media files amounting to a total data volume of 2.7 GB. Bacterial 16S rRNA gene sequences were submitted to the Sequence Read Archive at the National Center for Biotechnology Information and have been assigned biosample accession numbers SAMN14692096 to SAMN14692121 under project number PRJNA627958.

ETHICS STATEMENT
Because the study of echinoderms and most other invertebrates is not regulated by the Animal Welfare Act (United States), approval for this study by the California State University Long Beach Institutional Animal Care and Use Committee was not required. Collections were conducted under Scientific Collecting Permit SC-7296 (California Department of Fish and Wildlife, Sacramento, CA, United States).
AUTHOR CONTRIBUTIONS

AZ and BP conceived the study. AZ, AG, JD, and BP carried out the experiments, analyzed the data, and wrote the manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2020.01697/full#supplementary-material

REFERENCES

Alain, K., Olagnon, M., Desbruyères, D., Pagé, A., Barbier, G., Juniper, S. K., et al. (2002). Phylogenetic characterization of the bacterial assemblage associated with mucous secretions of the hydrothermal vent polychaete Paralvinella palmiformis. FEMS Microbiol. Ecol. 42, 463–476. doi: 10.1111/j.1574-6941.2002.tb01035.x

Allen, M. J., Dieter, D., Mubarak, J., Weisberg, S. B., and Moore, S. L. (1999). “Megabenthic invertebrate assemblages of the mainland shelf of Southern California in 1994,” in Southern California Coastal Water Research Project 1997-1998 Annual Report, eds S. B. Weisberg and D. Elmore (Westminster, CA: Southern California Coastal Water Research Project), 113–124.

An, T. T., and Picardal, F. W. (2014). Desulfovicrobium induanensis gen. nov., sp. nov., a benzoeatoxidizing, sulfate-reducing bacterium isolated from water extracted from a coal bed. Int. J. Syst. Evol. Microbiol. 64, 2907–2914. doi: 10.1099/ijs.0.064873-0

Anisimov, A. P. (1982). Morphological and cytological characteristics of the alimentary canal epithelium of the heart sea urchin Echinocardium cordatum, Pennant (Echinodermata: Echinoidea: Spatangoidea). Biol. Morya 2, 39–46.

Aprill, A., McNally, S., Parsons, R., and Weber, L. (2015). Minor revision to V4 region SSU rRNA 806R gene primer greatly increases detection of SAR11 bacterioplankton. Aquat. Microb. Plankton 75, 129–137. doi: 10.3354/ame01753

Bik, E. M., Costello, E. K., Switzer, A. D., Callahan, B. J., Holmes, S. P., Wells, R. S., et al. (2016). Marine mammals harbor unique microbiotas shaped by and yet distinct from the sea. Nat. Commun. 7:10516. doi: 10.1038/srep10516

Bokulich, N. A., Karcher, B. D., Rideout, J. R., Dillon, M., Bolyen, E., Knight, R., et al. (2018). Optimizing taxonomic classification of marker-gene amplicon sequences with QIIME 2’s q2-feature-classifier plugin. Microbiome 6:90. doi: 10.1186/s40168-018-0470-z

Bolyen, E., Rideout, J. R., Dillon, M. R., Bokulich, N. A., Abnet, C. C., Al-Ghali, G. A., et al. (2019). Reproducible, interactive, scalable and extensible microbiome data science using QIIME 2. Nat. Biotechnol. 37, 852–857. doi: 10.1038/s41587-019-0209-9

Boon, A. R., and Duineveld, G. C. A. (2012). Phytopigments and fatty acids in the gut of the deposit-feeding heart urchin Echinocardium cordatum in the southern North Sea: selective feeding and its contribution to the benthic carbon budget. J. Sea Res. 67, 77–84. doi: 10.1016/j.seares.2011.10.004

Breznak, J. A., and Brune, A. (1994). Role of microorganisms in the digestion of lignocellulose by termites. Annu. Rev. Entomol. 39, 453–487. doi: 10.1146/annurev.en.39.010194.002321

Brigham, R. L., and De Ridder, C. (1998). Symbiotic relationship of Thiolithrix spp. with an echinoderm. Appl. Environ. Microbiol. 64, 3491–3495. doi: 10.1128/aem.64.9.3491-3495.1998

Bromley, R. G., and Asgaard, U. (1975). Sediment structures produced by a spatangoid echinoid: a problem of preservation. Bull. Geol. Soc. Denmark 24, 261–281.

Bromley, R. G., Jensen, M., and Asgaard, U. (1995). Spatangoid echinoids: deep-tier trace fossils and chemosymbiosis. Neues Jahrb. Geol. Palaeontol. Abhand. 195, 25–35. doi: 10.1127/njgpa/195/1995/25

Brown, M., and Lowe, D. G. (2007). Automatic panoramic image stitching using invariant features. Int. J. Comp. Vis. 74, 59–73. doi: 10.1007/s11263-006-0002-3

Brownell, C. L. (1970). The relationship between Morphology and Ecology in the Spatangoid Urrchin Brissura Latifrons. M.Sc. thesis, Oregon State University, Corvallis, OR.

Brune, A., and Dietrich, C. (2015). The gut microbiota of termites: digesting the diversity in the light of ecology and evolution. Annu. Rev. Microbiol. 69, 145–166. doi: 10.1146/annurev-micro-092412-155715

Buchanan, J. B., Brown, B. E., Coombs, T. L., Pirie, B. J. S., and Allen, J. A. (1980). The accumulation of ferric iron in the guts of some spatangoid echinoderms. J. Mar. Biol. Assoc. U. K. 60, 631–640. doi: 10.1017/s0025315400040327

Callahan, B. J., McMurdie, P. J., Rosen, M. J., Han, A. W., Johnson, A. J. A., and Holmes, S. P. (2016). DADA2: high-resolution sample inference from Illumina amplicon data. Nat. Methods 13, 581–583. doi: 10.1038/nmeth.3869

Caporaso, J. G., Lauber, C. L., Walters, W. A., Berg-Lyons, D., Lozupone, C. A., Turnbaugh, P. J., et al. (2011). Global patterns of 16s rDNA diversity at a depth of millions of sequences per sample. Proc. Natl. Acad. Sci. U.S.A. 108, 4516–4522. doi: 10.1073/pnas.100080107
Hollertz, K., and Duchêne, J. C. (2001). Burrowing behaviour and sediment reworking in the heart urchin Brissopsis lyrifera Forbes (Spatangoidea). Mar. Biol. 139, 951–957. doi: 10.1007/s002270010629

Hood, S., and Mooi, R. (1998). “Taxonomy and phylogenetics of extant Brisaster (Echinoidea: Spatangoidea),” in Echinodermata, San Francisco, eds R. Mooi and M. Telford (Rotterdam: Balkema), 681–686.

Hunter, J. D. (2007). Matplotlib: a 2D graphics environment. Comp. Sci. Eng. 9, 90–95. doi: 10.1109/mcse.2007.55

Iida, T., Ohkuma, M., Ohtoko, K., and Kudo, T. (2000). Symbiotic spirochetes in the termite hindgut: phylogenetic identification of ecosymbiotic spirochetes of oxymonad protists. FEBS Microbiol. Ecol. 34, 17–26. doi: 10.1111/j.1574-6941.2000.tb00750.x

Jackson, E. W., Pepe-Ranney, C., Debenport, S. J., Buckley, D. H., and Hewson, I. (2018). The microbial landscape of sea stars and the anatomical and interspecies variability of their microbiome. Front. Microbiol. 9:1829. doi: 10.3389/fmicb.2018.01829

Jackson, K. L. Whitcraft, C. R., and Dillon, J. (2014). Diversity of Desulfovibrioaceae and overall activity of sulfate-reducing microorganisms in and around a salt pan in a southern California coastal wetland. Wetlands 34, 969–977. doi: 10.1007/s11273-014-0560-z

Jumars, P. A. (1993). “Gourmards of mud: diet selection in marine deposit feeders,” in Mechanisms of Diet Choice, ed. N. E. Hughes (Oxford: Blackwell Scientific Publishers), 124–156.

Jumars, P. A. (2000). Animal guts as ideal chemical reactors: maximizing absorption rates. Am. Nat. 155, 527–543. doi: 10.1086/303333

Kanazawa, K. (1992). Adaptation of test shape for burrowing and locomotion in living starfish (Echinoidea: Spatangoida), “in Infauna communities off the coast of Washington. Proc. Linnean Soc. New South Wales 123, 119–139.

Linse, K., Walker, L. J., and Barnes, D. K. A. (2008). Biodiversity of echinoids and their epibions around the Scotia Arc, Antarctica. Antarctic Sci. 20, 227–244. doi: 10.1017/S0954004808001181

Lohrer, A. M., Thrush, S. F., Hunt, L., Hancock, N., and Lundquist, C. (2005). Rapid reworking of subtidal sediments by burrowing spatangoid urchins. J. Exp. Mar. Biol. Ecol. 312, 155–169. doi: 10.1016/j.jembe.2005.02.002

Lopez, G. R., and Levinton, J. S. (1987). Ecology of deposit-feeding animals in marine sediments. Quar. Rev. Biol. 62, 235–260. doi: 10.1086/415511

Lowery, C. M., Bralower, T. J., Owens, J. D., Rodriguez-Tovar, F. J., Jones, H., Smit, J., et al. (2018). Rapid recovery of life at ground zero of the end-Cretaceous mass extinction. Nature 558, 288–291.

Lozupone, C., and Knight, R. (2005). UniFrac: a new phylogenetic method for comparing microbial communities. Appl. Environ. Microbiol. 71, 8228–8235. doi: 10.1128/aem.71.12.8228-8235.2005

Macdonald, T. A., Burd, B. J., and van Roosdelaar, A. (2012). Facultative feeding and consistency of trophic structure in marine soft-bottom microbenthic communities. Mar. Ecol. Prog. Ser. 445, 129–140. doi: 10.3354/meps09478

Magot, M., Fardeau, M. L., Arnauld, O., Lanau, C., Ollivier, B., Thomas, P., et al. (1997). Spirochaeta smaragdinae sp. nov., a new mesophilic strictly anaerobic spirochete from an oil field. FEBS Microbiol. Lett. 155, 185–191. doi: 10.1111/j.1574-6968.1997.tb13876.x

Margulis, L., and Hinkle, G. (2006). “Large symbiotic spirochetes: ‘Clevelandinida, Cristatipina, Diplolyci, Hollandiina and Pellitina,” in The Prokaryotes: Proteobacteria: Dleta, Epsilon Subclass, eds M. Dworkin, S. Falkow, E. Rosenberg, K. H. Schleifer, and E. Stackebrandt (New York, NY: Springer), 971–982. doi: 10.1007/978-3-7074-8_43

Markov, A. V. (1994). Morphology, systematics and phylogeny of schizasterid sea urchins. Trudy Paleontol. Inst. Rossii. Akad. Nauk 258, 1–94.

Mayer, L. M., Schick, L. L., Self, R. F. L., Jumars, P. A., Findlay, R. L., Chen, Z., et al. (1997). digestive environments of benthic macroinvertebrate guts: enzymes, surfactants and dissolved organic matter. J. Mar. Res. 55, 785–812. doi: 10.1080/207300224093224427

McCauley, X. J. E., and Carey, A. G. Jr. (1967). Echinoida of Oregon. J. Fish. Res. Board Can. 24, 1385–1401. doi: 10.1139/f67-114

McCullough, S. W. (2016). Coral-associated bacterial diversity is conserved across two polar biomes. Polar Biol. 39, 913–923. doi: 10.1007/s00300-015-1873-y

Moffitt, S. E., Hill, T. M., Roopnarine, P. D., and Kennett, J. P. (2015). Response of seafloor ecosystems to abrupt global climate change. Proc. Natl. Acad. Sci. U.S.A. 112, 3229–3236.

McCullough, S. W. (2015). Giving microbes their due – animal life in a microbiologically dominant world. J. Exp. Biol. 218, 1968–1973. doi: 10.1242/jeb.115121

McKinney, M. L., McNamara, K. J., and Wiedman, L. A. (1988). Echinoids form the La Meseta Formation (Eocene), Seymour Island, Antarctica. Geol. Soc. Am. Memoir 169, 499–503. doi: 10.1130/MEM169-5-499

Mielke, L. N., David, B., Dubois, P., Leport, G., and De Ridder, C. (2016). Trophic plasticity of Antarctic echinoids under contrasted environmental conditions. Polar Biol. 39, 913–923. doi: 10.1007/s00305-015-1873-y

Moffitt, S. E., Hill, T. M., Roopnarine, P. D., and Kennett, J. P. (2015). Response of seafloor ecosystems to abrupt global climate change. Proc. Natl. Acad. Sci. U.S.A. 112, 4684–4689. doi: 10.1073/pnas.1417130112

Moran, N., Ochman, H., and Hammer, T. J. (2019). Evolutionary and ecological consequences of gut microbial communities. Annu. Rev. Ecol. Evol. Syst. 50, 451–475. doi: 10.1146/annurev-ecolsys-110617-062453

Mortensen, T. (1951). A Monograph of the Echinoidae, Volume 2, Spatangoidea II. C. A. Copenhagen: Reitzel, 593.

Murakata, N., and Markle, P. (2018). 2016-2017 Joint Water Pollution Control Plant Biennial Receiving Water Monitoring Report. Whittier, CA: County Sanitation Districts of Los Angeles County.

Na, H., Lever, M. A., Kjeldsen, K. U., Schulz, F., and Jørgensen, B. B. (2015). Uncultured Desulfobacteraceae and Crenarchaeotal group C3 incorporate 13C-acetate in coastal marine sediment. Environ. Microbiol. Rep. 7, 614–622. doi: 10.1111/1758-2229.12296

Nichols, D. (1959). Changes in the chalk heart-urchin Micraster interpreted in relation to living forms. Philos. Trans. R. Soc. B 242, 347–437. doi: 10.1098/ rstb.1959.0007

Nichols, F. H. (1975). Dynamics and energetics of three deposit-feeding benthic invertebrate populations in Puget Sound, Washington. Ecol. Monogr. 45, 57–82. doi: 10.2307/1942331
Temara, A., De Ridder, C., Kuenen, J. G., and Robertson, L. A. (1993). Sulphide-oxidizing bacteria in the burrowing echinoid *Echinocardium cordatum* (Echinodermata). *Mar. Biol.* 115, 179–185. doi: 10.1007/bf00346533

Thompson, B., Tsukada, D., and Laughlin, J. (1993). Megabenthic assemblages of coastal shelves, slopes, and basins off Southern California. *Bull. Southern Calif. Acad. Sci.* 92, 25–42.

Thompson, B. A. W., and Riddle, M. J. (2005). Bioturbation behaviour of the spatangoid urchin *Abatus ingens* in Antarctic marine sediments. *Mar. Ecol. Prog. Series* 290, 135–143. doi: 10.3354/meps290135

Thompson, B. E., Jones, G. F., Laughlin, J. D., and Tsukada, D. T. (1987). Distribution, abundance, and size composition of echinoids from basin slopes off Southern California. *Bull. Southern Calif. Acad. Sci.* 86, 113–125.

Thorsen, M. S. (1998). Microbial activity, oxygen status and fermentation in the gut of the irregular sea urchin *Echinocardium cordatum*. *Mar. Biol.* 132, 423–433. doi: 10.1007/s002270050408

Thorsen, M. S. (1999). Abundance and biomass of the gut-living microorganisms (bacteria, protozoa and fungi) in the irregular sea urchin *Echinocardium cordatum* (Spatangoida: Echinodermata). *Mar. Biol.* 133, 353–360. doi: 10.1007/s002270050474

Thorsen, M. S., Wieland, A., Ploug, H., Kragelund, C., and Nielsen, P. H. (2003). Distribution, identity and activity of symbiotic bacteria in anoxic aggregates from the hindgut of the sea urchin *Echinocardium cordatum*. *Ophelia* 57, 1–12. doi: 10.1080/003053403100012627

Van de Water, J. A. J. M., Melkonian, R., Junca, H., Voolstra, C. R., Rexynaud, S., Allemend, D., et al. (2016). *Spirochaetae* dominate the microbial community associated with the red coral *Corallium rubrum* on a broad geographic scale. *Sci. Rep.* 6:22777. doi: 10.1038/srep22777

Van Oosterom, J. T. (2012). *Gene Flow in East Antarctic Echinoderms and Resilience to Climate Change*. Ph. D. thesis, Deakin University, Melbourne, VIC.

Vázquez-Baeza, Y., Pirrung, M., Gonzalez, A., and Knight, R. (2013). EMPeror: a tool for visualizing high-throughput microbial community data. *GigaScience* 2:16. doi: 10.1186/2047-217X-2-16

Vopel, K., Vopel, A., Thistle, D., and Hancock, N. (2007). Effects of spatangoid heart urchins on O2 supply into coastal sediment. *Mar. Ecol. Prog. Series* 333, 161–171. doi: 10.3354/meps333161

Walker, D. E., and Gagnon, J. M. (2014). Locomotion and functional spine morphology of the heart urchin *Brissaster fragilis*, with comparisons to *B. latifrons*. *J. Mar. Biol. 2014*:297631. doi: 10.1155/2014/297631

Wang, L., Zhao, X., Xu, H., Bao, X., Liu, X., Chang, Y., et al. (2018). Characterization of the bacterial community in different parts of the gut of sea cucumber (*Apostichopus japonicus*) and its variation during gut regeneration. *Aqua. Res.* 49, 1987–1996. doi: 10.1016/are.13654

Weigel, B. L. (2020). Sea cucumber intestinal regeneration reveals deterministic assembly of the gut microbiome. *Appl. Environ. Microbiol.* 86:e00489-20. doi: 10.1128/AEM.00489-20

Wenzel, M., Radek, R., Brugerolle, G., and König, H. (2003). Identification of the ectosymbiotic bacteria of *Mixotricha paradoxa* involved in movement symbiosis. *Eur. J. Protistol.* 39, 11–23. doi: 10.1078/0932-4739-00893

Widdicombe, S., Austen, M. C., Kendall, M. A., Olsgard, F., Schaanning, M. T., Dashfield, S. L., et al. (2004). Importance of bioturbators for biodiversity maintenance: indirect effects of fishing disturbance. *Mar. Ecol. Prog. Series* 275, 1–10. doi: 10.3354/meps275001

Yamazaki, Y., Meirelles, P. M., Mino, S., Suda, W., Oshima, K., Hattori, M., et al. (2016). Individual *Apostichopus japonicus* fecal microbiome reveals a link with polyhydroxybutyrate producers in host growth gaps. *Sci. Rep.* 6:21631. doi: 10.1038/srep21631

Yilmaz, P., Parfrey, L. W., Yarza, P., Gerken, J., Pruesse, E., Quast, C., et al. (2014). The SILVA and “All-species Living Tree Project (LTP)” taxonomic frameworks. *Nucleic Acids Res.* 42, 643–648. doi: 10.1093/nar/gkt1209

Zhang, Z., Li, D., Xu, W., Tang, R., and Li, L. (2019). Microbiome of co-cultured fish exhibits host selection and niche differentiation at the organ scale. *Front. Microbiol.* 10:2576. doi: 10.3389/fmicb.2019.02576

Ziegler, A. (2014). Rediscovery of an internal organ in heart urchins (*Echinoidae: Spatangoida*): morphology and evolution of the intestinal caecum. *Organ. Divers. Evol.* 14, 383–395. doi: 10.1007/s13127-014-0178-2

Ziegler, A. (2019). Combined visualization of echinoderm hard and soft parts using contrast-enhanced micro-computed tomography. *ZooKeys* 15, 172–191. doi: 10.11646/zookeys.15.1.19

Ziegler, A., Bock, C., Ketten, D. R., Mair, R. W., Mueller, S., Nagelmann, N., et al. (2018). Digital three-dimensional imaging techniques provide new analytical pathways for malacological research. *Am. Malacol. Bull.* 36, 248–273. doi: 10.4003/006.036.0205

Ziegler, A., Faber, C., Mueller, S., Nagelmann, N., and Schröder, L. (2014). A dataset comprising 141 magnetic resonance imaging scans of 98 extant sea urchin species. *GigaScience* 3:21. doi: 10.1186/2047-217X-3-21

Ziegler, A., and Menze, B. H. (2013). “Accelerated acquisition, visualization, and analysis of zoo-anatomical data,” in *Computation for Humanity: Information Technology to Advance Society*, eds J. Zander and P. J. Mosterman (Boca Raton, FL: CRC Press), 233–261. doi: 10.1201/9781315216751-16

Ziegler, A., Moot, R., Rolet, G., and De Ridder, C. (2010). Origin and evolutionary plasticity of the gastric caecum in sea urchins (Echinodermata: Echinoidea). *BMC Evol. Biol.* 10:313. doi: 10.1186/1471-2148-10-313

Ziegler, A., and Mueller, S. (2011). Analysis of freshly fixed and museum invertebrate specimens using high-resolution, high-throughput MRI. *PLoS ONE* 6:27277. doi: 10.1371/journal.pone.0027277

Ziegler, A., and Menze, B. H. (2013). “Accelerated acquisition, visualization, and analysis of zoo-anatomical data,” in *Computation for Humanity: Information Technology to Advance Society*, eds J. Zander and P. J. Mosterman (Boca Raton, FL: CRC Press), 233–261. doi: 10.1201/9781315216751-16

Ziegler, A., Moot, R., Rolet, G., and De Ridder, C. (2010). Origin and evolutionary plasticity of the gastric caecum in sea urchins (Echinodermata: Echinoidea). *BMC Evol. Biol.* 10:313. doi: 10.1186/1471-2148-10-313

Ziegler, A., and Mueller, S. (2011). Analysis of freshly fixed and museum invertebrate specimens using high-resolution, high-throughput MRI. *Methods Mol. Biol.* 771, 633–651. doi: 10.1007/978-1-61779-219-9_32

Ziegler, A., Gilligan, A. M., Dillon, J. G., and Pernet, B. (2020). *Schiasterid Heart Urchins host Microorganisms in a Digestive Symbiosis of Mesozoic Origin*. *MorphoBank Project #3616*. doi: 10.7934/P3616

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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