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
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Sophistication in a seemingly simple creature: a review of wild holothurian nutrition in marine ecosystems

Joséphine Pierrat^{1*} , Alexandre Bédier², Igor Eeckhaut³, Hélène Magalon^{1,4} and Patrick Frouin^{1,4}

¹UMR ENTROPIE (IRD, CNRS, Univ. Reunion, Ifremer, Univ. New Caledonia), University of La Réunion, St-Denis, 97 400, France

²Nutrina Production, Le Port, 97 420, France

³Biology of Marine Organisms and Biomimetism Lab, University of Mons, Mons, 7000, Belgium

⁴Labex Corail, Perpignan, 66 000, France

ABSTRACT

Holothurians are marine invertebrates that are among the most widespread benthic megafauna communities by both biomass and abundance in shallow-water and deep-sea ecosystems, their functions supporting important ecological services worldwide. Despite their simple appearance as sea cucumbers, holothurians show a wide range of feeding practices. However, information on what and how these animals eat is scattered and potentially confusing. We provide a comprehensive review of holothurian nutrition in coastal and deep-sea ecosystems. First, we describe morphological aspects of holothurian feeding and the ultrastructure of tentacles. We discuss the two processes for food capture, concluding that mucus adhesion is likely the main method; two mucous cells, type-1 and type-2, possibly allow the adhesion and de-adhesion, respectively, of food particles. Secondly, this review aims to clarify behavioural aspects of holothurian suspension- and deposit-feeding. We discuss the daily feeding cycle, and selective feeding strategies. We conclude that there is selectivity for fine and organically rich particles, and that feeding through the cloaca is also a route for nutrient absorption. Third, we provide a wide description of the diet of holothurians, which can be split into two categories: living and non-living material. We suggest that Synallactida, Molpadida, Persiculida, Holothuriida and Elasipodida, ingest the same fractions, and emphasise the importance of bacteria in the diet of holothurians.

Key words: holothurians, tentacle ultrastructure, bud epidermis, mucous cell, trophic mode, selective feeding, dietary, proteobacteria, plastic particles

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* Address for correspondence (Tel: +292692888677; E-mail: josephine.pierrat@univ-reunion.fr).

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I. INTRODUCTION

Dating back 460 million years, holothurians (commonly called sea cucumbers) are ubiquitous marine echinoderms belonging to the class Holothuroidea. An initial classification was established by Pearson (1914), who attempted to organise the species, under the name *Holothuria* L., into groups based on their gross morphology. Pawson & Fell (1965) subsequently proposed a classification based on tentacle, body, and calcareous ring morphology, defining five orders: Dendrochirotida, Apodida, Molpadida, Elasipodida, and Aspidochirotida. Until recently, this was the most widely used classification for holothurians. When the first broad-scale molecular phylogenetic analyses of Holothuroidea were completed, these deeply altered the previous classification through the suppression of the order Aspidochirotida, which was revealed to be polyphyletic. Species previously under Aspidochirotida have since been placed into three new orders: Synallactida, Persiculida (in part), and Holothuriida (Miller *et al.*, 2017). There are more than 1,752 accepted holothurian species (WoRMS, 2020), with new species being described each year.

Holothurians have a global distribution, colonising all biotopes of the ocean from the polar front (Lawrence & Guille, 1982; Féral & Magniez, 1985; Gutt, 1990; Post *et al.*, 2017; O'Loughlin, Bardsley & O'Hara, 2020) to the tropical zone (Sloan & von Bodungen, 1980; Wiedemeyer, 1994; Asha *et al.*, 2015; Resueño & Angara, 2020), with most species inhabiting the tropical Indo-West Pacific region (Conand, 1990). They have also colonised all depths, from shallow-water (Jaquemet, Rousset & Conand, 1999; Dissanayake & Stefansson, 2010; MacTavish *et al.*, 2012; Lee *et al.*, 2017) to hadal zones (Iken *et al.*, 2001; Jamieson *et al.*, 2011). They are among the most widespread benthic megafauna species in terms of biomass and abundance in many ecosystems, especially in the hadal zone, considered 'the kingdom of Holothuroidea' (Beliaev & Brueggeman, 1989; Kuhn *et al.*, 2014), in coral

reefs and lagoons (Uthicke, 1999; Wolfe & Davey, 2020), and in sheltered marine shallow habitats (Conde, Diaz & Sambrani, 1991).

Some species of holothurians are considered as luxury food (bêche-de-mer, trepang or hai-som), medicines, and aphrodisiacs in many Asian countries (Conand, 1990; Lovatelli *et al.*, 2004; Shiell & Uthicke, 2005; Toral-Granda, Lovatelli & Vasconcellos, 2008). More than 70 species of holothurians are commonly harvested (Purcell *et al.*, 2016), predominantly from the Indo-Pacific region (Kinch *et al.*, 2008; Conand, 2018). In some locations, populations of highly commercially valuable species have been decimated to a point that fishing regulations and regulatory measures alone may be insufficient to restore populations (Friedman *et al.*, 2011). This overexploitation is linked to a shift from traditional to semi-industrial fisheries (Conand, 2001). Countries of the Indian Ocean, West Pacific and Latin America have active fisheries (Conand, 2018), and South-East Asia is considered the main world market (Rahman & Yusoff, 2017). The world fishery of holothurians quadrupled between 1955 and 2012 to satisfy the increasing Asian market for 'bêche-de-mer' (Rahman, Yusoff & Arshad, 2015). Consequently, many countries in the Indo-Pacific have prioritised sea cucumber aquaculture in their development plans (Jimmy, Pickering & Hair, 2012). In addition, restocking, sea ranching, and sea farming have been described as potential alternatives to reduce pressure on wild holothurian populations and their worldwide overexploitation. Three species have been registered recently in CITES (Convention on International Trade in Endangered Species of Wild Fauna and Flora) Appendix II: *Holothuria (Microthele) fuscogilva* Cherbouner, *Holothuria (Microthele) nobilis* (Selenka), and *Holothuria (Microthele) whitmaei* Bell, each with high commercial value but declining natural stocks (Di Simone, Horellou & Conand, 2019).

Due to their global distribution favouring many local studies, and their importance in Asian culture, holothurians have gained increasing importance in recent decades. A search on

Google Scholar using the key words ‘sea cucumber’ for general publications and ‘bêche-de-mer’ for those focusing on fisheries and aquaculture, shows that the number of articles on both topics has increased considerably since the 1990s (Fig. 1).

The increasing literature provides scattered data about holothurian feeding, making it difficult to obtain a broad view on this topic. This review intends to assess our understanding of holothurian feeding. The nutritional aspects considered herein include feeding (the act of consuming food), digestion and absorption. To our knowledge, only one review has been published previously on the nutrition of holothurians (edited by Jangoux & Lawrence, 1982), about 40 years ago. There is no comprehensive review covering the nutrition traits of holothurians in relation to their ecology, and collating information on their feeding patterns. While much information exists on the feeding of farmed holothurians, this relates only to controlled conditions. Thus, our aim herein is to analyse holistically the nutrition of wild holothurians.

II. MORPHOLOGICAL AND PHYSIOLOGICAL ASPECTS

(1) Tentacle structure and movements during ingestion

The number, size, and structure of tentacles varies among holothurian orders. Based on their structure, Massin (1982) defined five types of tentacles: dendritic, peltate, pinnate, digitate, and peltodendritic. The first four of these are common and widely used in the description of tentacle morphology of holothurians. Species within Dendrochirotida possess dendritic tentacles whereas those of Molpadida possess digitate tentacles (Levin, 1989). Peltate tentacles, described as “cauliflower-like structures” (Boulard, Massin & Jangoux, 1982, p. 134) or “a nasturtium leaf with a central short stalk giving

off horizontal branches” (Hyman, 1955, *cf.* Cameron & Fankboner, 1984, p. 193), are found within Elasipodida, Synallactida, Persiculida, and Holothuriida. In some cases, the tentacle structure shows variations among species within the same order, such as for Apodida species, which are found in coastal and deep-sea ecosystems and possess pinnate or digitate tentacles. Combinations of these four main types of tentacle can be found in the literature: (i) a ‘peltodendritic tentacle’ which is, according to Massin (1982), a combination of the shaft of a peltate tentacle and distal end of a dendritic tentacle; (ii) a ‘peltatodigitate tentacle’, which is a combination of the shaft of a peltate tentacle and distal end of a digitate tentacle (Miller *et al.*, 2017). Peltodendritic tentacles are uncommon, while peltatodigitate tentacles are found in Apodida (Miller *et al.*, 2017).

Tentacle morphology may also differ between early and adult stages of the same species (Cameron & Fankboner, 1984). Generally, a tentacle possesses several shafts that end with one or more discs, composed of numerous apical papillae (Fig. 3A, B). The apex of each papilla is characterised by the presence of buds (Fig. 3C) with various cell types. The main difference in tentacle structure is the degree of branching of the main tentacle stalk (Fig. 2), from an unbranched (digitate form), to a slightly branched (peltate form), to a highly branched (pinnate form) to an ultra-branched (dendritic form) structure. These differences reveal an adaptive radiation facilitating habitat and feeding specialisations (Sokolova, 1958; Hansen, 1975). From an ecological perspective, differences in tentacular morphology among sympatric holothurian species with overlapping bathymetric ranges may allow resource partitioning to avoid competitive interactions (Roberts & Moore, 1997).

A comprehensive review by Massin (1982) described the feeding mechanisms of deposit-feeding holothurians. When the animal initiates feeding, the tentacles expand into the water column, and are pressed onto or into the sediment surface where they perform investigatory movements (Boulard *et al.*, 1982). The tentacular movement is thought to be driven by the hydrostatic pressure of the ambulacral fluid of the water-vascular system and the mesothelial muscles (Boulard *et al.*, 1982). Tentacles can also cooperate to capture large debris such as fragments of seagrass or macroalgae. In this case, when the desired food elements are located, the extremities of the tentacle retract first, thus entrapping the particles between the buds (Fankboner, 1978; Levin, 1989). Sokolova (1958) described how tentacle morphology can influence the collection of particulate food in deep-sea holothurian species; well-developed marginal processes on discs are capable of picking up single food particles from the sediment, whereas species with undifferentiated discs indiscriminately ingest the upper layer of the sediment. After collection, the tentacle continues to contract, bending towards the mouth and penetrating it. At this ingestion stage, mouth size is the limiting factor (Myers, 1977), although a proportion of captured particles is lost during transport to the mouth (Powell, 1977; Levin, 1989). As the tentacle penetrates the pharyngeal cavity, particle removal is facilitated by wiping

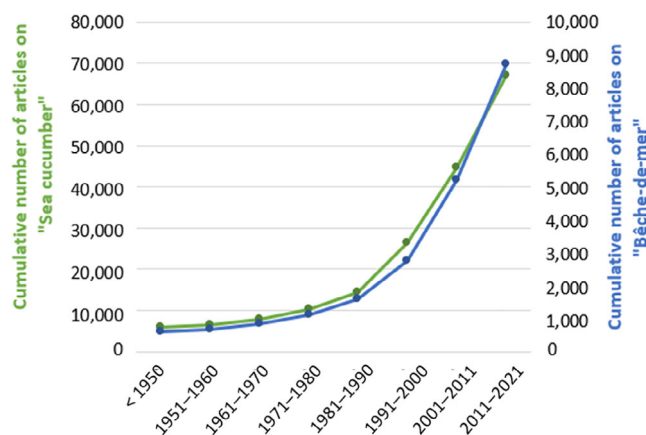


Fig 1. Cumulative number of articles on “sea cucumber” for general publications and “bêche-de-mer” for those focusing on fisheries and aquaculture since 1950. Note the different scales for the two axes.

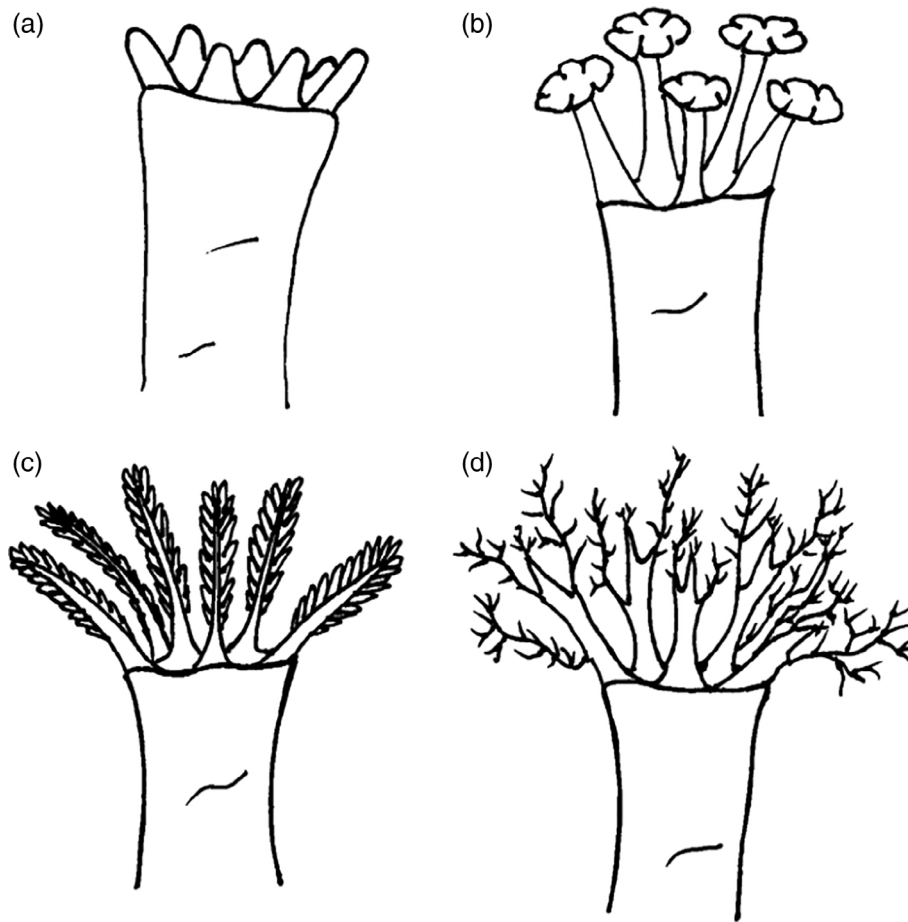


Fig 2. Examples of holothurian tentacle morphology. (A) Digitate (unbranched); (B) peltate (slightly branched); (C) pinnate (highly branched); (D) dendritic (ultra-branched).

the tentacle against the bulging pharynx wall (Cameron & Fankboner, 1984). Finally, the tentacle withdraws from the mouth and extends again to continue investigating for food sources.

(2) Food-capture mechanism

Understanding the mechanism(s) of capture of food particles in holothurians was approached *via* studies in functional morphology integrating data concerning the ultrastructure of the tentacles to determine the composition of the tentacle parts in contact with the ingested sediment. These studies, mainly carried out in the 1980s and 1990s, are unfortunately few in number. Sixteen species were investigated, mainly Dendrochirotida (13 species; Smith, 1983; McKenzie, 1987), two former Aspirochirotida (Boulard *et al.*, 1982; Cameron & Fankboner, 1984) and one Apodida (Flammang & Conand, 2004). Roberts & Moore (1997) and Fankboner (1981) used scanning electron microscopy to study the fine external structure of tentacles of four species of Elasipodida and five species of Dendrochirotida without detailing their cellular composition. Based on these studies and behavioural experiments, several authors have proposed mechanisms for the capture of food particles in several

species (Roberts, 1979; Hammond, 1982; Boulard *et al.*, 1982; Cameron & Fankboner, 1984). Holothurians appear to use two methods for food collection: food particle ensnarement (a mechanical process) and food particle adhesion (a chemical process). Food particle ensnarement was proposed in early studies about the functioning of the tentacles, since when adhesion has gained support as the principal mechanism involved in the capture of food particles.

(a) Food particle adhesion

Several authors described a mucus-like secretion on the oral tentacles and suggested a primary role in holothurian food capture (Boulard *et al.*, 1982). Roberts & Bryce (1982) described mucus-secreting cells in the tentacular epidermis of several tropical species and stated that the adhesive material would play a role in collecting food particles. Levin (1982), studying *Apostichopus japonicus* (Selenka), presumed that adhesion was a primary function of the tentacle during feeding. Similarly, Hammond (1982) observed mucus coating the tentacular surface and stated that adhesion is a significant factor in food collection for Synallactida, Persiculida, and Holothuriida, a statement reiterated by other

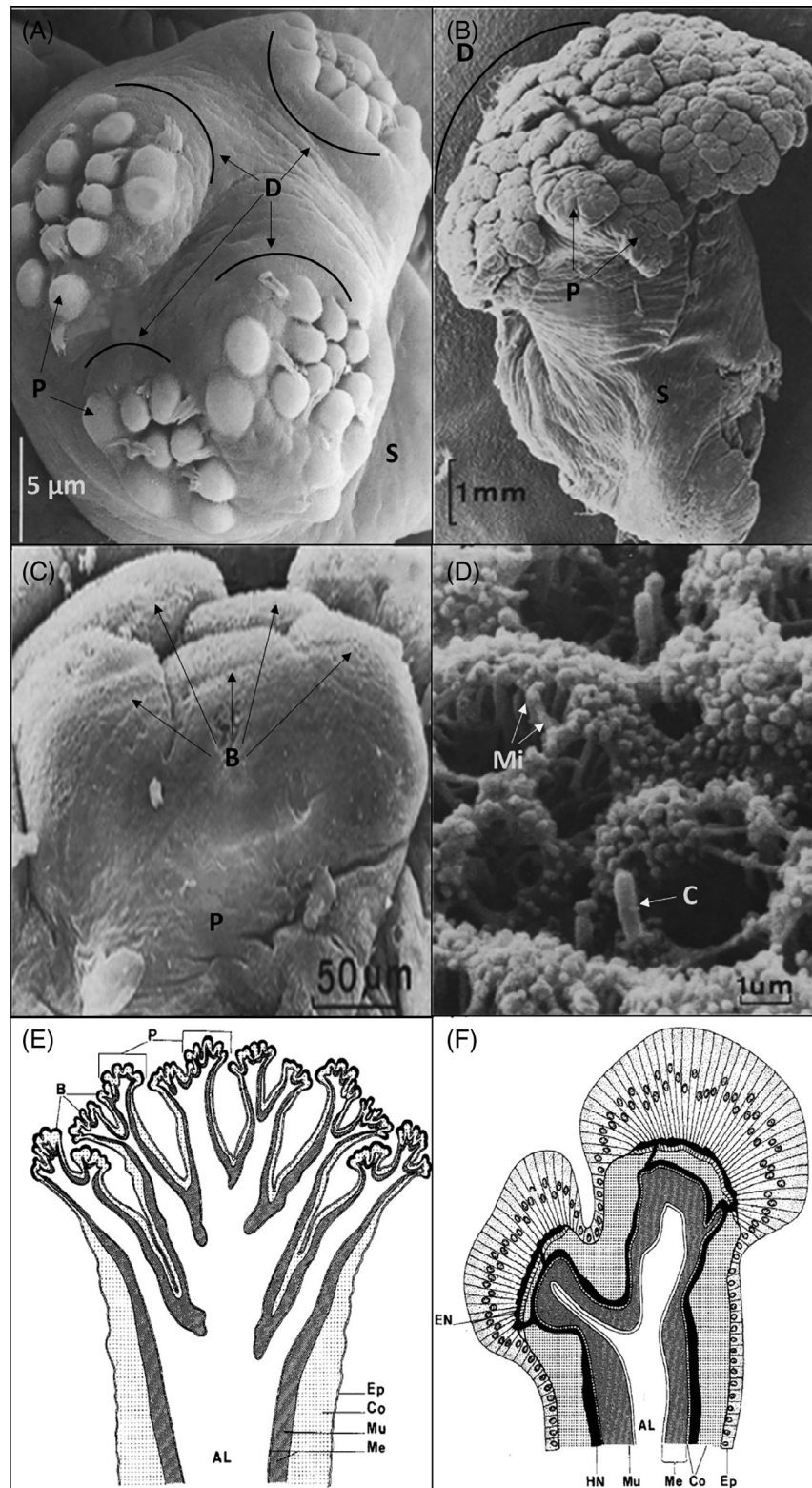


Fig 3. Fine ultrastructure of the holothurian tentacle. Digitate tentacle of (A) *Thyonidium* sp. and (B) *Holothuria forskali*. D, discs; P, Papillae; S, shaft. (C) Profile view of a five-bud papilla. B, buds. (D) Arrangement of cilia and microvilli on bud surface. C, cilium; Mi, microvilli. (E, F) Diagrammatic sections through a whole tentacle (E) and through a two-bud papilla (F). AL, ambulacral lumen; Co, connective tissue layer; EN, epineural nerve plexus; Ep, epidermis; HN, hyponeural nerve plexus; Me, mesothelium; Mu, muscles. Modified from Bouland *et al.* (1982) and McKenzie (1987).

authors working on temperate species (Smith, 1983; Costelloe & Keegan, 1984; Holtz & MacDonald, 2009).

Transmission electron microscopy studies have shown buds to be the sites of secretion. The bud epidermis can include six types of cells in the species investigated to date, with 3–5 of these cell types usually being present (Table 1). The functions of these cells have been deduced by drawing parallels with the roles of cells in the adhesive disc of podia of echinoderms. (i) ‘Support cells’ have been described in some Dendrochiro-tida (McKenzie, 1987) and in the only species of Apodida investigated (Flammang & Conand, 2004) (Table 1). (ii) A ‘vesicular cell’ type has only been described in Apodida (Flammang & Conand, 2004) with unknown functions. (iii) Ciliated cells, also named ‘unciliated cells’, ‘ciliated cells’ or ‘unciliated sensory cells’ have been observed in all species investigated (Table 1). Two roles have been suggested for this cell type: they could mechanically disengage particles (Fankboner, 1978) or they could be sensory (Flammang & Conand, 2004). Bouland *et al.* (1982) associated the bud structure of *Holothuria* (*Panningothuria*) *forskali* Delle Chiaje with Laverack’s (1974) description of chemosensory organs in marine invertebrates, where cilia are proposed to function as olfactory receptors, while microvilli are gustatory. Cilia are generally short and non-motile (Dorsett & Hyde, 1969; Schulte & Riehl, 1976), like those depicted by Bouland *et al.* (1982). The apical elements of the tentacles are the first structures to contact substrates while the tentacles actively forage for food, strongly suggesting that the buds are likely to be chemosensory (Bouland *et al.*, 1982). The cells of the epidermis of each bud are adjacent to the epineural nervous plate, which is connected to the hyponeural nerve plexus of the tentacles (Fig. 3E, F). Mesothelial muscles are present in each buccal tentacle (Fig. 3E; Bouland *et al.*, 1982; McKenzie, 1987). (iv) ‘Mucous cells’, also called ‘mucocytes’ were observed in the tentacles of Dendrochiro-tida, Holothuriida and Apodida (Table 1). As similar cells are also observed outside the tentacles, their mucus-secreting role is probably to ensure the presence of a physical barrier on the epidermis. (v) Type-1 and (vi) type-2 secretory cells are thought to play a direct role in the capture of food particles. Type-1 secretory cells, also referred to as ‘granular cells’, ‘glandular vesicular cells’ or ‘papillate cells’ (Table 1) are characterised by numerous dense-cored vesicles of 200–700 nm, with their diameter varying among species. These vesicles are found in many microvilli and are thought to be secreted into the cuticle. They may be homologues of the adhesive cells found in the podia of echinoderms. Type-2 secretory cells, also named ‘type-2 neurosecretory cells’ or ‘granular cells’, have been observed in Dendrochiro-tida and Apodida (Table 1). This cell type possesses numerous dense-cored vesicles of 60–130 nm, again with their diameter varying among species. These vesicles are found in the cell apex and also are thought to be secreted into the cuticle. They are thought to be homologues of de-adhesive cells found in the podia of echinoderms. Thus, the chemical capture of food particles could be performed by the secretion of type-1 secretory cells and, when the tentacles are placed in the oral cavity, adhesion could be removed by the secretion of type-2 secretory cells.

This hypothesis is currently only theoretical and future functional morphology studies are needed, together with transcriptomic data to characterise the proteins expressed at the tentacle buds.

(b) Food particle ensnarement

Authors have long debated the mechanisms of food capture by which particles are trapped within bud interstices. Roberts (1979) suggested that expansion of the peltate tentacles caused inter-bud spaces to open on the tentacles, which could then mechanically trap particles when the tentacles retract. Cameron & Fankboner (1984) reported mechanical ensnarement to be of relatively little importance for *Parastichopus californicus* [now *Apostichopus californicus* (Stimpson)] since there is no reverse process of tentacle expansion/relaxation while the tentacle is in the pharyngeal cavity. Food detection is likely to be related to the ciliated cells as described in Section II.2, as the tentacles spread onto the substratum or in the water column.

(3) Digestive and assimilation processes in holothurians

(a) Digestive tract morphology

The nomenclature used for parts of the digestive tract varies among authors. Trefz (1958) distinguished between the mouth, pharynx, oesophagus, foregut, midgut, hindgut, cloaca and cloacal opening when studying the physiology of *Holothuria* (*Halodeima*) *atra* (Jaeger) (Fig. 4A). Massin (1978), in his study on holothurian nutrition, focused mainly on *Holothuria* (*Holothuria*) *tubulosa* Gmelin (Fig. 4B), differentiating the pharyngeal bulb surrounded by the calcareous ring, followed by the three portions of the digestive tract (foregut, midgut and hindgut), and finally the cloaca. Dividing the digestive tract between mouth and cloacal opening into three sections – foregut, midgut and hindgut – gives a broader view, and many studies describing holothurian feeding use this partitioning (e.g. Trefz, 1958; Ward-Rainey, Rainey & Stackebrandt, 1996; Taddéi, 2006; Plotieau, 2012; Amaro *et al.*, 2012).

The foregut includes the pharyngeal bulb, oesophagus, and the first descending loop of the intestine. The oesophagus is divided into two parts of equivalent lengths, both ending in a sphincter (sphincters 1 and 2) (Fig. 4B). The midgut begins immediately beyond the first intestinal loop where the proximal tubules of the *rete mirabile* are attached and consists of the ascending anterior loop of the intestine. A third sphincter separates the midgut from the hindgut. The hindgut, which represents approximately 70% of the total length of the gut, consists of the final posterior descending loop of the intestine and the cloaca from which respiratory trees emanate (Dolmatov & Ginanova, 2009). Only species from Pneumophora have respiratory trees (Dendrochiro-tida, Holothuriida, Molpadida, Persiculida and Synallactida; Miller *et al.*, 2017). The descending loop of the foregut is paralleled by the dorsal haemal plexus, which extends posteriorly to the

Table 1. Ultrastructural composition of the tentacle buds of holothuroids with the nomenclatural terms used in analyses by various authors. The first column describes the corresponding cells observed in podia of echinoderms where a duo glandular adhesive system (adhesion and de-adhesion) has been identified

| Authors | Fankboner (1978) | Boulard <i>et al.</i> (1982) | Smith (1983) | Cameron & Fankboner (1984) | McKenzie (1987) | Flammang & Conand (2004) |
|--|---|---------------------------------|---|--|--|--|
| Analysed taxon | Dendrochirotida | Holothuriida | Dendrochirotida | Synallactida | Dendrochirotida | Apodida |
| Number of species studied | 1 | 1 | 1 | 1 | 11 | 1 |
| Cell with no particular development of intracellular component | – | – | – | – | Support cell | Support cell (T-shaped cell) with 400–600 nm vesicles |
| Cell with large vesicle containing one spherule | – | – | – | – | – | Vesicular cell with numerous 4 µm vesicles |
| Cell with small apical cilia [presumed to be sensory cell, or cilia suggested to disengage food particles by Fankboner, 1978] | Uniciliated cell | Uniciliated cell | Ciliated cell | Uniciliated cell | Uniciliated cell | Uniciliated sensory cell |
| Cell filled with large clear vesicle (presumed to participate to the protection of the external surface) | Mucous cell | Mucous cell | – | – | Mucous cell (two different types) | Mucocyte (goblet-shaped cell) with 2 µm dense-cored vesicles |
| Cell with large dense-cored vesicles secreted into the cuticle (presumed to be similar to the adhesive cells identified in podia of echinoderms) | Papillary cell with 500 nm dense-cored vesicles released into the cuticle | Glandular vesicular cell | Papillate cell with 300–600 nm dense-cored vesicles released into the cuticle | Granular cell with 600–700 nm dense-cored vesicles released into the cuticle | Type-1 secretory cell with 200–400 nm dense-cored vesicles released into the cuticle | Type-1 secretory cell (spherical) with 250 nm dense-cored vesicles |
| Cell with small dense-cored vesicles (presumed to be similar to the de-adhesive cells identified in podia of echinoderms) | – | – | Granular cell with 130 nm dense-cored vesicles | – | Type-2 presumed neurosecretory cell (due to low numbers) with 60–100 nm dense-cored vesicles | Type-2 secretory cell (spherical) with 90 nm dense-cored vesicles |

anal end of the animal where it loops back. The dorsal haemal plexus is a complex of pulsating channels attached to the dorsal midline of the foregut. Longitudinally along the ventral side of this region runs the ventral sinus. The *rete mirabile* is a complex of tiny tubules, entangled within the respiratory tree, and attached to the ascending loop of the midgut. These tubules terminate into the main channel of the dorsal haemal plexus.

Although the foregut, midgut, and hindgut have three different functions (accumulation, digestion, and assimilation), most species of Synallactida, Persiculida, and Holothuriida

are able to assimilate food from these three parts (Massin, 1978).

(b) Cell and enzymatic activities

The epithelia of the pharynx, oesophagus, and cloaca all contain T-shaped cells that are involved in the uptake of dissolved organic matter. The gut epithelium is composed of enterocytes with a filamentous coat for intracellular digestion (Féral & Massin, 1982). A broad range of hydrolytic gut enzymes has evolved in association with the diet

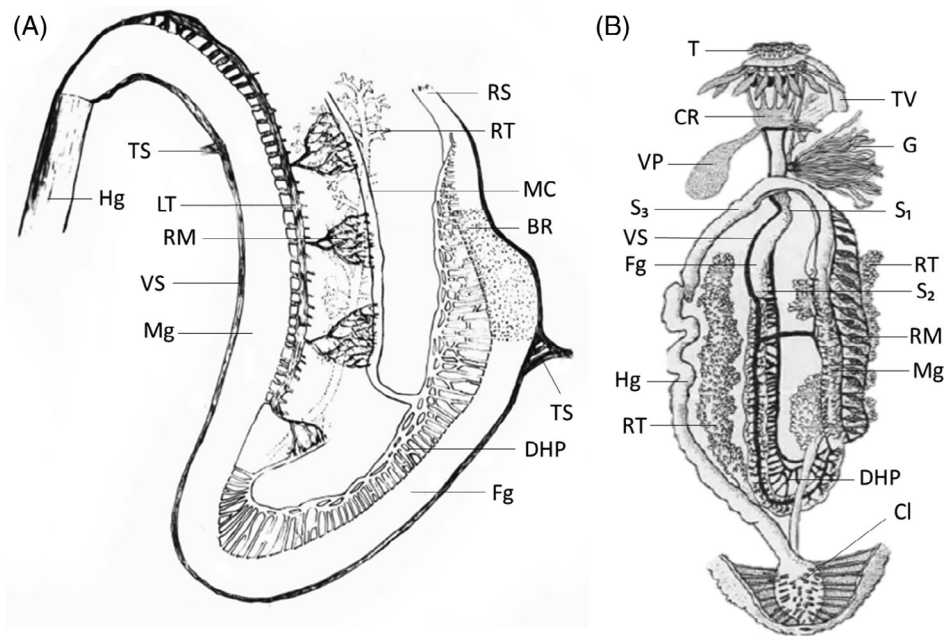


Fig 4. Digestive tract morphology of (A) *Holothuria atra* and (B) *H. tubulosa*. BR, brown region of the anterior foregut; Cl, cloaca; CR, calcareous ring; DHP, dorsal haemal plexus; Fg, foregut; G, gonads; Hg, hindgut; LT, longitudinal tube of rete mirabile; MC, main channel of dorsal haemal plexus; Mg, midgut; RM, rete mirabile; RS, region of severance; RT, respiratory tree; S₁, sphincter 1; S₂, sphincter 2; S₃, sphincter 3; T, tentacles; TS, transverse sinuses; TV, tentacle vesicles; VP, vesicle of Poli; VS, ventral sinus. Modified from Trefz (1958) and Massin (1978, inspired by Cuénot, 1948).

(Féral, 1989). Hydrolytic enzyme activity has been reported within the digestive tract, on cell membranes (Féral, 1989), and inside cells (Lawrence, 1982). Roberts *et al.* (2000) reviewed the enzymes found in the gut of deposit-feeding holothurians (shallow-water species in Holothuriida, Synallactida, Apodida and Dendrochirotida, and deep-sea species of Holothuriida, Persiculida, Synallactida and Elasipodida), finding broad similarities in their hydrolytic enzymes. Gut enzymes include: (i) esterases that hydrolyse short-chain fatty acids, with strong activity throughout the gut (Fish, 1967; Féral, 1989), (ii) lipases that hydrolyse long-chain fatty acids, with lowest activity (Clifford *et al.*, 1982; Féral, 1989; Manship, 1995), (iii) proteases, (iv) peptidases, which hydrolyse peptide bonds and may be important in the initial stages of protein digestion (Massin, 1984; Féral, 1989; Roberts *et al.*, 2000), (v) saccharidases, and (vi) phosphatases, present at high concentrations in the gut tissue (Féral, 1989; Boetius & Felbeck, 1995).

Trefz (1958) observed phagocytic activity of abundant cells in the holothurian gut, such as round phagocytic amoebocyte cells that ingest and digest bacteria (*Bacillus subtilis*), but not indigestible particles.

(c) Obtaining nutrients through cloacal water retention

The major functions of cloacal ventilation are commonly described as respiration, excretion, and salt balance. Holothurians with respiratory trees (absent in Apodida and Elasipodida) have the ability to pump large volumes of water into

and out of the cloacal opening. A similar pumping mechanism is reported for echinoderms such as holothuroids (Newell & Courtney, 1965; Brown & Shick, 1979) and edrioasteroids (Bell, 1977), annelids (Wolcott, 1981), and crustaceans (Fox, 1952).

Innovative work, using radiography and stable isotope enrichment, demonstrated that the epithelium of the respiratory tree can assimilate dissolved organic matter such as monosaccharides and amino acids from the water column during cloacal water retention (Fontaine & Chia, 1968; Brothers, Lee & Nestler, 2015). The respiratory tree of *Cucumaria lactea* [now *Ocnus lacteus* (Forbes & Goodsir)] can assimilate dissolved ³H-labelled glycine and ³H-labelled glucose (Fontaine & Chia, 1968). The respiratory tree of *A. californicus* assimilated ¹⁵N-labelled amino acids and peptides (Brothers *et al.*, 2015) and ¹⁴C-labelled unicellular algae, suggesting the transfer of nutrients from the respiratory tree into the haemal system (Jaekle & Strathmann, 2013). Thus, it is possible that holothurians can obtain nutrients through their respiratory trees as a result of cloacal pumping, but additional studies are needed to clarify its importance to these animals.

III. HOLOTHURIAN FEEDING BEHAVIOUR

As for all organisms, the movements and behaviour of holothurians will be driven by physiological requirements such

as feeding, reproduction, or avoidance of stress factors (Mercier, Battaglene & Hamel, 1999; Pitt & Duy, 2004; Meng *et al.*, 2011; Hamel *et al.*, 2019). A possible role of pheromones in intra- or inter-specific control of reproduction was recently hypothesised for holothurians (Marquet *et al.*, 2018).

(1) Deposit- or suspension-feeding

The first larval stage in holothurians, the auricularia, feeds on phytoplankton (planktotrophic) by means of cilia located on the epidermis. The second larval stage, the doliolaria, is non-feeding, and the final stage, the pentactula, develops the tentacles that are present in juveniles and adults. The first and second stages can be absent in holothurians with direct development (McEdward & Miner, 2001; Raff & Byrne, 2006). In general, juveniles and adults are particle-feeders, which usually exhibit either suspension- (in Dendrochirotida) or deposit-feeding (most other taxa) behaviour (Roberts *et al.*, 2000; Pawson, 1970; Bakus, 1973). Suspension-feeding holothurians tend to live in high-energy environments to allow them to intercept food particles (Taghon & Jumars, 1984). Deposit-feeding behaviour can be subdivided into three groups related to feeding depth (Roberts *et al.*, 2000). (i) Epibenthic deposit-feeders feed at the interface between the water column and sediment, or on the surface of seagrasses (Cuvillier, 2016), the sand scattered over the tegument of other specimens (e.g. *H. atra*; J.P., personal observations), boulders and corals (Roberts *et al.*, 2000; J.P., personal observations), or sponges (Hammond & Wilkinson, 1985). (ii) Subsurface deposit-feeders feed below the sediment surface. (iii) Funnel deposit-feeders create funnel-shaped depressions in sediments and feed on particles trapped in these funnels (Jumars, 1993).

Feeding mode may vary at the individual or population scale. According to Cadée (1984), many benthic individuals are opportunistic in their feeding mode. Some individuals of species in Dendrochirotida (Roberts *et al.*, 2000), Synallactida, Persiculida, and Holothuriida (Da Silva, Cameron & Fankboner, 1986) can shift between deposit-feeding and suspension-feeding. This may be possible if the individual is positioned in a negative geotropic orientation (Da Silva *et al.*, 1986). *Leptopentacta elongata* (Düben & Koren) and other shallow-water species (Fankboner, 1981; Levin, 1989) are known to shift opportunistically from suspension- to deposit-feeding when the concentration of suspended particulate organic matter is low. This opportunistic behaviour is driven by environmental factors such as food pulses in shallow-water (tidal or seasonal fluctuations) (Cushing, 1959) and deep-sea ecosystems (Jumars, Self & Nowell, 1982; Billett *et al.*, 1983).

At the population scale, the feeding modes of some holothurian species can vary depending on their geographic location. *Holothuria (Thymiosyscia) arenicola* Semper is an epibenthic feeder, feeding on surface layers of sand under coral debris in the Indo-West Pacific region and Cuba (Levin, 1989), whereas in the Bahamas, it feeds on particles in sediments

(Mosher, 1980) as a subsurface deposit-feeder. The underlying reasons for these intraspecific differences at a population scale, and their drivers, are poorly studied.

(2) Daily burrowing cycle

A study on the burrowing behaviour of shallow-water holothurians in Palao Island classified holothurians into two groups according to their feeding habits (Yamanouchi, 1939): (i) species that do not burrow into sediments or other substrates and feed continuously, such as *Holothuria (Halodeima) edulis* Lesson, *H. (Semperothuria) flavomaculata* Semper or *H. atra* (Yamanouchi, 1939; Trefz, 1958; Uthicke, 1994) and (ii) species that show a daily cycle of burrowing and feeding (Yamanouchi, 1939, 1956; Mercier *et al.*, 1999; Lavitra *et al.*, 2009). The factors regulating burrowing and feeding cycles appear to be complex (Yamanouchi, 1939, 1956), with differences among species increasing the complexity of cues that drive such behaviour and impeding generalisation.

According to Yamanouchi (1956), most adults prefer to feed during the day, particularly in the early afternoon when the water temperature is highest (Mercier *et al.*, 1999) and productivity of the marine ecosystems is maximal (Heil *et al.*, 2004). Some species, such as *Stichopus chloronotus* Brandt, move under corals from 00:00 to 10:00 and then return to sediments to feed during the day (Yamanouchi, 1956). Similarly, *Stichopus variegatus* (now *Stichopus hermanni* Semper) hides under seagrasses between 20:00 and 10:00 (Yamanouchi, 1956). Coulon & Jangoux (1993) argued that it may be more energetically advantageous for juveniles of *H. tubulosa* to stop feeding during the coldest hours of the day. Individuals exposed to abnormal water temperatures exhibit unusual behaviour and feeding activity (Kato & Hirata, 1990). The temperate species *A. japonicus* aestivates during the warmest months when water temperatures are between 20 and 24.5°C (Choe, 1963), especially large and mature individuals (Liu *et al.*, 1996). Aestivating individuals hide under structures and enter a state of dormancy (Liu *et al.*, 1996; Yang *et al.*, 2005) in which feeding activity ceases (Yuan *et al.*, 2007) until water temperature becomes more favourable. During aestivation, *A. japonicus* individuals undergo a series of physiological and morphological changes (Wang *et al.*, 2008), losing 30–50% of their body mass (Liu *et al.*, 1996). The digestive tract degenerates to half its pre-aestivation size (Li *et al.*, 1996; Liu *et al.*, 1996). Physiological responses associated with digestion are reduced during aestivation, with lower activity of digestive enzymes in the principal portions of the gut (Cui, Dong & Lu, 2000). Similarly, under high temperatures (austral summer), *Holothuria (Metriatyla) scabra* Jaeger, changes its burrowing behaviour, remaining on the sediment surface and feeding frequently, whereas at low water temperatures (austral winter) it remains beneath the sediment surface for most of the day (Mercier, Battaglene & Hamel, 2000). Thus, seasonal patterns will play a major role in the regulation of the daily burrowing cycle of this species.

The effects of salinity on the foraging activities of holothurians are unclear. Mercier *et al.* (1999, 2000) found that *H. scabra* cease feeding when water salinity is lowest, and burrow into the substrate. James & James (1994) reported conflicting results, with juveniles emerging during low tide. Skewes *et al.* (2006) reported that only a third of *H. scabra* adults emerge during high tide.

Some studies show conflicting results concerning daily burrowing cycles of juveniles and adults. On one hand, daily burrowing cycles can be similar for juveniles and adults of the same species. In Palao Island, *H. scabra* adults were observed burrowing in the sediment between 03:30 and 15:00 and feeding between 15:00 and 03:30 (Yamanouchi, 1956). In the Philippines, *H. scabra* juveniles burrow in sediments between 03:00 and 09:00 and remain hidden until they emerge to feed between 15:00 to 03:00 (Altamirano, Recente & Rodriguez Jr., 2017). Despite their different life stages and different geographical regions, both species therefore feed during the same time periods. On the other hand, holothurian size has been reported to affect burrowing and daily cycles. Mercier *et al.* (1999) found the burrowing cycle of *H. scabra* to be driven by light for small juveniles (10–40 mm), which burrow at sunrise and emerge close to sunset, whereas temperature was more influential for intermediate-sized juveniles (40–140 mm), which emerge earlier in the afternoon. Like many echinoderms, the nocturnal feeding and movement habits of smaller holothurians might be an adaptation to avoid predation (Nelson & Vance, 1979; Hammond, 1982).

(3) Selectivity by deposit-feeding holothurians

Feeding selectivity is observed in many marine species of different taxa, such as Polychaeta (Petch, 1986; Shimeta, 1996), Gastropoda (Whitlatch & Obrebski, 1980), or Bivalvia (Hylleberg & Gallucci, 1975). Some other echinoderms show selectivity in feeding, including species in Asteroidea (Mellin *et al.*, 2017) and Echinoidea (Larson, Vadas & Keser, 1980; Boon & Duineveld, 2012).

To determine particle selectivity in deposit-feeding holothuroids, most studies compare the biochemical composition and concentration of compounds present within sediments around and beneath foraging individuals and in their foregut (Moriarty, 1982; Hammond, 1983; Amaro *et al.*, 2010). Examining particle selectivity is subject to the difficulty of investigating sediment ingestion at the appropriate scale (Lopez & Levinton, 1987). Particle selection is generally based on size, surface texture, specific gravity, and the presence of an organic coating (Taghon, 1989), but it can also be explained by mechanical processes (Jumars *et al.*, 1982). Discussion of particle selectivity by deposit-feeding holothurians usually refers to two different modes: the selection of particles with a specific grain size or selection of those with higher organic content. We found 29 studies describing both aspects of particle selectivity for shallow-water (23 studies) and deep-sea (six studies) holothurian species (Table 2). A total of 50 species have been studied, with

only one study focused on suspension-feeding species (Hamel & Mercier, 1998).

(a) Size selection of particles

Whether deposit-feeding holothurians exhibit a preference for substrates with a specific grain size is still debated (Mercier *et al.*, 2000), with conflicting results available (Table 2) even within a single species. For example, Mezali & Soualili (2013) stated that *H. tubulosa* and *Holothuria (Roweothuria) poli* Dell Chiaje select a grain size ranging between 200 and 600 μm , whereas Massin & Jangoux (1976) found no size preference for these same species. Some authors conclude that holothurians are unselective regarding the size of ingested particles because ingested sediment particles tend to be very similar in size to those of the bottom sediment (Yingst, 1976; Sloan & von Bodungen, 1980; Hammond, 1982). Trefz (1958) noted that diverse holothurian species forage and subsist on different substrates, suggesting that they may focus their feeding on specific substrates. Of the 29 shallow-water species evaluated to date, 20 show grain size selectivity (Table 2); among the four deep-sea species studied, only *Molpadia blakei* (Th  el) is unselective (Table 2; Khripounoff & Sibuet, 1980).

Particle-size selection seems to be species dependent: preferred grain size is 200–600 μm for *H. tubulosa* (Mezali & Soualili, 2013), 125–250 μm for *H. scabra* (Baskar, 1994), and 2000–3500 μm for *H. nobilis* (Roberts & Bryce, 1982). Deep-sea species seem to prefer finer particles than shallow-water species, with a median of 7–14 μm for *Benthogone rosea* Koehler and 8–54 μm for *Paroriza pallens* (Koehler) (Khripounoff & Sibuet, 1980). Particle-size selection may change within a species depending on the season: *Holothuria (Stauropora) hawaiiensis* Fisher and *Bohadschia vitiensis* (Semper) prefer finer particles during the spawning period and gravel and coarse particles during the rest of the year (Dar & Ahmad, 2006).

More information is required to assess the benefits of feeding on small-grain sediments for holothurians and the reasons why only some species are selective. Although the mechanisms by which a preferred substrate is selected are not understood, Roberts (1979) proposed that this could be associated with the morphology of the oral tentacles.

(b) Selection of organically rich particles

A wide range of holothurian species from tropical, shallow-water regions (Moriarty, 1982; Hammond, 1983; Uthicke, 1999; Uthicke & Karez, 1999), temperate seas (Hauksson, 1979; Amon & Herndl, 1991), and deep seas (Miller *et al.*, 2000; Wigham *et al.*, 2003, 2008) are known to select organically rich particles from the sediment (Table 2). These species seem to be attracted by sediments with high organic content (Yingst, 1982) and can differentiate and capture the preferred particles (Massin & Jangoux, 1976; Moriarty, 1982). Out of the 37 species studied, 29 prefer particles enriched in organic matter (Table 2). Both shallow-

Table 2. Synthesis of studies on selective feeding strategies in shallow-water and deep-sea holothurians

| Taxon | Size particle selectivity | Organic matter selectivity | Location | Authors |
|--------------------------------|--|----------------------------|-------------------------------------|-------------------------------|
| <i>Coastal species</i> | | | | |
| > Apodida | | | | |
| <i>Euapta lappa</i> | No | Not studied | Discovery Bay, Jamaica | Hammond (1982) |
| <i>Leptosynapta tenuis</i> | No | Not studied | North Carolina, USA | Powell (1977) |
| > Holothuriida | | | | |
| <i>Actinopyga agassizi</i> | No | Not studied | Discovery Bay, Jamaica | Hammond (1982) |
| <i>Bohadschia bivittata</i> | Yes, 400 µm | Not studied | Pari Island, Indonesia | Roberts (1979) |
| <i>Bohadschia vitiensis</i> | Yes, gravel and coarse particles (spawning period: fine particles) | Yes | Hurghada, Egypt | Dar & Ahmad (2006) |
| <i>Holothuria arenicola</i> | No | Not studied | Discovery Bay, Jamaica | Hammond (1982) |
| <i>Holothuria atra</i> | Not studied | No | Great Palm Island, Australia | Uthicke & Karez (1999) |
| | Yes, coarser particles | Yes | Red Sea coast, Egypt | Dar (2004) |
| | Yes | Not studied | Waikiki branch, Hawai | Trefz (1958) |
| | Yes, gravel and coarse particles (in spawning period fine particles) | Yes | Hurghada, Red Sea, Egypt | Dar & Ahmad (2006) |
| | Not studied | Yes | Great Barrier Reef, Australia | Moriarty (1982) |
| | Yes, coarser particles | Yes | El Qasr reef, Saudi Arabia | Behairy, Beltagi & Rao (1985) |
| | Yes, 350 µm | Not studied | Pari Island, Indonesia | Roberts (1979) |
| <i>Holothuria cinerascens</i> | Yes, <63–500 µm | Not studied | Beacon Island, Australia | Roberts & Bryce (1982) |
| <i>Holothuria edulis</i> | Not studied | No | Great Palm Island, Australia | Uthicke & Karez (1999) |
| | Yes, 63–125 µm | Not studied | Beacon Island, Australia | Roberts & Bryce (1982) |
| <i>Holothuria forskali</i> | Yes, 60–200 µm | Yes | Algiers and Bou-Ismaïl Bay, Algeria | Mezali & Soualili (2013) |
| | Not studied | Yes | Toulon, France | Massin & Jangoux (1976) |
| <i>Holothuria grisea</i> | No | Not studied | Discovery Bay, Jamaica | Hammond (1982) |
| <i>Holothuria hartmeyeri</i> | Yes, 2000–3500 µm | Not studied | Beacon Island, Australia | Roberts & Bryce (1982) |
| <i>Holothuria hawaiiensis</i> | Yes, gravel and coarse particles (in spawning period fine particles) | Yes | Hurghada, Egypt | Dar & Ahmad (2006) |
| | Yes, <63–500 µm | Not studied | Beacon Island, Australia | Roberts & Bryce (1982) |
| <i>Holothuria impatiens</i> | Yes, coarse particles | Yes | Red Sea coast, Egypt | Dar (2004) |
| <i>Holothuria leucospilota</i> | Yes, coarse particles | Yes | Red Sea coast, Egypt | Dar (2004) |
| <i>Holothuria marmorata</i> | Yes, coarse particles | Yes | Red Sea coast, Egypt | Dar (2004) |
| <i>Holothuria mexicana</i> | No | Not studied | Discovery Bay, Jamaica | Hammond (1982) |
| <i>Holothuria nobilis</i> | Not studied | No | Great Palm Island, Australia | Uthicke & Karez (1999) |
| | Yes, 2000–3500 µm | Not studied | Beacon Island, Australia | Roberts & Bryce (1982) |
| <i>Holothuria cf. povicax</i> | Yes, 2000–3500 µm | Not studied | Beacon Island, Australia | Roberts & Bryce (1982) |
| <i>Holothuria poli</i> | Yes, 200–600 µm | Yes | Algiers and Bou-Ismaïl Bay, Algeria | Mezali & Soualili (2013) |
| | No | Yes | Toulon, France | Massin & Jangoux (1976) |
| <i>Holothuria sanctori</i> | Yes, 60–200 µm | Yes | Algiers and Bou-Ismaïl Bay, Algeria | Mezali & Soualili (2013) |
| | Not studied | Yes | Canary Islands, Spain | Navarro <i>et al.</i> (2013) |
| <i>Holothuria scabra</i> | Yes, 125–250 µm | Not studied | Palk Bay, India | Baskar (1994) |
| <i>Holothuria stellati</i> | Yes, 60–200 µm | No | Algiers and Bou-Ismaïl Bay, Algeria | Mezali & Soualili (2013) |

Table 2. (Cont.)

| Taxon | Size particle selectivity | Organic matter selectivity | Location | Authors |
|-----------------------------------|---------------------------|----------------------------|---|-------------------------------|
| <i>Holothuria tubulosa</i> | Yes, 200–600 µm | Yes | Algiers and Bou-Ismaïl Bay, Algeria | Mezali & Soualili (2013) |
| | Not studied | Yes | Gulf of Naples, Italy | Amon & Herndl (1991) |
| | No | Yes | Toulon, France | Massin & Jangoux (1976) |
| > Molpadida | | | | |
| <i>Molpadia oolitica</i> | Yes, smallest particles | Not studied | Cape Cod Bay, USA | Rhoads & Young (1971) |
| > Synallactida | | | | |
| <i>Australostichopus mollis</i> | No | Yes | Mahurangi Harbour, New Zealand | Slater, Jeffs & Sewell (2011) |
| <i>Isostichopus badionotus</i> | No | Yes | Bermuda | Sloan & von Bodungen (1980) |
| | No | Not studied | Discovery Bay, Jamaica | Hammond (1982) |
| <i>Parastichopus californicus</i> | Not studied | Yes | British Columbia, Canada | Paltzat <i>et al.</i> (2008) |
| <i>Parastichopus parvimensis</i> | No | Yes | Santa Catalina Island, USA | Yingst (1976) |
| <i>Stichopus chloronotus</i> | Not studied | Yes | Great Palm Island, Australia | Uthicke & Karez (1999) |
| | Not studied | Yes | Lizard Island, Australia | Uthicke (1999) |
| | Not studied | Yes | Great Barrier Reef, Australia | Moriarty (1982) |
| <i>Stichopus japonicus</i> | Not studied | Yes | Aquarium experiment, Japan | Michio <i>et al.</i> (2003) |
| <i>Stichopus tremulus</i> | Yes, coarse particles | Yes | Raunefjorden, Norway | Hauksson (1979) |
| <i>Stichopus variegatus</i> | Not studied | Yes | Great Palm Island, Australia | Uthicke & Karez (1999) |
| Deep-sea species | | | | |
| > Apodida | | | | |
| <i>Chiridota</i> sp. | Not studied | No | Santa Catalina Basin and Hawaiian slope | Miller <i>et al.</i> (2000) |
| > Elasiopodida | | | | |
| <i>Amperima rosea</i> | Not studied | Yes | Porcupine Abyssal Plain, NE Atlantic | Wigham <i>et al.</i> (2003) |
| | Not studied | Yes | Porcupine Abyssal Plain, NE Atlantic | Ginger <i>et al.</i> (2001) |
| <i>Benthogone rosea</i> | Yes, 7–14 µm | Yes | Golfe de Gascogne, France | Khripounoff & Sibuet (1980) |
| <i>Ellipinion molle</i> | Not studied | Yes | Porcupine Abyssal Plain, NE Atlantic | Ginger <i>et al.</i> (2001) |
| <i>Pannychia moseleyi</i> | Not studied | Yes | Santa Catalina Basin and Hawaiian slope | Miller <i>et al.</i> (2000) |
| <i>Peniagone vignoni</i> | Not studied | No | West Antarctic Peninsula | Wigham <i>et al.</i> (2008) |
| <i>Protelpidia murrayi</i> | Not studied | No | West Antarctic Peninsula | Wigham <i>et al.</i> (2008) |
| <i>Psychropotes longicauda</i> | Yes, 6.2–44 µm | Yes | Golfe de Gascogne, France | Khripounoff & Sibuet (1980) |
| | Not studied | Yes | Porcupine Abyssal Plain, NE Atlantic | Wigham <i>et al.</i> (2003) |
| <i>Scotoplanes globosa</i> | Not studied | Yes | Santa Catalina Basin and Hawaiian slope | Miller <i>et al.</i> (2000) |
| > Holothuriida | | | | |
| <i>Mesothuria camosa</i> | Not studied | Yes | Santa Catalina Basin and Hawaiian slope | Miller <i>et al.</i> (2000) |
| > Molpadida | | | | |
| <i>Molpadia blakei</i> | Not studied | Yes | Porcupine Abyssal Plain, NE Atlantic | Wigham <i>et al.</i> (2003) |
| | No | Not studied | Golfe de Gascogne, France | Khripounoff & Sibuet (1980) |
| <i>Molpadia musculus</i> | Not studied | No | West Antarctic Peninsula | Wigham <i>et al.</i> (2008) |
| > Persiculida | | | | |
| <i>Paroriza pallens</i> | Yes, 8–54 µm | Yes | Golfe de Gascogne, France | Khripounoff & Sibuet (1980) |

(Continues)

Table 2. (Cont.)

| Taxon | Size particle selectivity | Organic matter selectivity | Location | Authors |
|---------------------------------|---------------------------|----------------------------|---|-------------------------------|
| <i>Pseudostichopus villosus</i> | Not studied | Yes | Porcupine Abyssal Plain, NE Atlantic | Wigham <i>et al.</i> (2003) |
| <i>Pseudostichopus sp.</i> | Not studied | Yes | Porcupine Abyssal Plain, NE Atlantic | Wigham <i>et al.</i> (2003) |
| | Not studied | No | West Antarctic Peninsula | Wigham <i>et al.</i> (2008) |
| > Synallactida | | | | |
| <i>Paelopatides retifer</i> | Not studied | No | Santa Catalina Basin and Hawaiian slope | Miller <i>et al.</i> (2000) |
| <i>Oneirophanta mutabilis</i> | Not studied | Yes | Porcupine Abyssal Plain, NE Atlantic | Witbaard <i>et al.</i> (2001) |
| | Not studied | Yes | Porcupine Abyssal Plain, NE Atlantic | Wigham <i>et al.</i> (2003) |

water and deep-sea species seem to be highly selective, with 17 out of 20 (85%) and 12 out of 17 species (70%), respectively, showing selectivity for organic matter (Table 2).

Generally, the organic matter content of the ingested sediment is much higher than in the sediment surrounding the animal (Hauksson, 1979). Khrpounoff & Sibuet (1980), studying the selective feeding of four abyssal species (*Psychropotes longicauda* Théel, *P. pallens*, *B. rosea* and *M. blakei*), found that with concentrations of organic carbon and nitrogen in the foregut were four and six times greater, respectively, than concentrations in the local sediment.

Moriarty (1982) suggested that particle size selection in holothurian deposit-feeders might be explained by a non-uniform distribution of organic matter in sediments. Uthicke (1999) supported this hypothesis, arguing that a higher organic content is usually associated with smaller sediment grains with low specific gravity, which a richer host microflora (bacteria, microalgae, etc.), due to their higher surface area to volume ratio (Johnstone, Koop & Larkum, 1990). All studies, with one exception [for *Holothuria (Holothuria) stellati* Delle Chiaje (Mezali & Soualili, 2013)], that showed a preference for a particular grain size also found a preference for organically rich particles where this was investigated (Table 2). A relationship between the organic load and particle size could therefore explain the confusion between these two selective strategies of deposit-feeders.

(c) Patch selectivity

Patch selectivity describes the preference of a mobile organism to feed on patches in a heterogeneous environment (Uthicke & Karez, 1999). A patchy distribution of shallow-water holothurians, thought to be related to their feeding habits (Uthicke & Karez, 1999), has been described in ecosystems including Milne Bay Province and Torres Strait islands (Skewes *et al.*, 2002, 2006), the Gulf of Mannar in India (Asha *et al.*, 2015), New Caledonia (Purcell *et al.*, 2009), and most parts of the western Indian Ocean (Conand, 2008). For holothurians, the energetic cost of processing poor sediment is higher than the cost of moving to a more suitable feeding substrate (Mercier

et al., 1999). Mercier *et al.* (1999) found that the locomotive speed of *H. scabra* juveniles was higher on substrates with low organic matter content ($71\text{--}331\text{ cm day}^{-1}$) than on substrates of better quality ($150\text{--}215\text{ cm day}^{-1}$), reflecting an active search for organically rich sediments. Similar results were reported for *H. atra* and *S. variegatus* (now *S. hermanni*), which move 0–52 m per day and cover greater distances on poorer feeding areas (Yamanouchi, 1939). Other studies on *Cucumaria frondosa* (Gunnerus) (Hamel & Mercier, 1998), *P. californicus* (now *A. californicus*) (Cameron & Fankboner, 1984), *Parastichopus chitonoides* (Young & Chia, 1982), *Actinopyga echinites* (Jaeger) (Wiedemeyer, 1994), and *H. scabra* (James & James, 1994; Altamirano *et al.*, 2017) generally describe the quality of the substrate where holothurians were observed but do not detail optimal substrates in terms of nutritional or ecological features (e.g. optimal grain size, organic matter content, origin of sediments, presence of conspecifics or optimal light regime).

A patchy distribution also has been observed in deep-sea ecosystems for several species (Billett, Llewellyn & Watson, 1988; Ruhl & Smith, 2004) that feed preferentially on nutritionally rich food patches (Hauksson, 1979; Hudson *et al.*, 2005; Jamieson *et al.*, 2011). Particulate organic carbon deposited on the seabed is considered an important factor controlling the local abundance and composition of macrofauna and megafauna (Sibuet, 1985; De Leo *et al.*, 2010). Significant correlations have been found between food availability and megafaunal abundance, particularly for holothurians (Billett *et al.*, 2001; Ruhl & Smith, 2004). Preferences for selective feeding amongst holothurian species may therefore function in niche partitioning (Roberts, 1979; Sloan & von Bodungen, 1980; Massin & Doumen, 1986), with reduced intra- and inter-specific competition as a result (Jamieson *et al.*, 2011).

IV. THE SEDIMENT INGESTED BY HOLOTHURIANS: A COMPLEX FOOD SOURCE

Despite much published work on holothurians, very few studies have focused on the quality of their food. Table 3 provides

a list of known food sources for holothurians, of both organic and inorganic origin. We divide these food sources into two categories: living and non-living fractions. Living fractions include organisms associated with sediments such as bacteria, photosynthetic organisms or meiofauna. Non-living fractions include organically derived detrital matter such as phytodetritus, dead and decaying animals, and faecal pellets and inorganic compounds such as coral scraps, shell remains, coralline algae, foraminiferal tests, and silicates. Sadly, holothurians have not escaped the increasing pollution of the ocean, with ingestion of plastic particles or microplastics recorded (Graham & Thompson, 2009; Renzi *et al.*, 2018).

(1) The living fractions

(a) Bacteria

The total biomass of bacteria is relatively high in shallow-water and deep-sea sediments (Zobell & Morita, 1959; Danovaro, Fabiano & Della Croce, 1993; Danovaro *et al.*, 1998; Rex *et al.*, 2006). Most benthic bacteria are not suspended in interstitial water but are attached to mineral or organic sediment particles (Dale, 1974) in aggregates and colonies (Taddéi, 2006). Thus, bacteria could represent a major food source for deposit-feeders (Sorokin, 1972; Massin, 1982) such as holothurians. We found only 10 articles that studied bacteria in the diet of deposit-feeding holothurians, of which four focused on the diversity of ingested bacteria, three on their abundance, and three on both. To our knowledge, the abundance and diversity of bacteria in the gut of suspension-feeding holothurians has not been assessed.

Most studies found a similar distribution pattern of bacteria in the digestive tract for coastal (Taddéi, 2006; Plotieau *et al.*, 2013) and deep-sea holothurian species (Deming & Colwell, 1982; Roberts *et al.*, 2001; Amaro *et al.*, 2012): an increase in bacterial abundance between the sediment and the foregut contents, then a decrease between the foregut and the hindgut. For example, Taddéi (2006), studying two coastal species *H. atra* and *Holothuria (Mertensiothuria) leucospilota* (Brandt), found an abundance of 1.50×10^7 bacteria g^{-1} in coral reef shallow-water sediments of La Réunion. This increased to 3.66×10^7 bacteria g^{-1} in the foregut of *H. atra* and 4.46×10^7 bacteria g^{-1} in the foregut of *Holothuria leucospilota*, which decreased significantly to 1.10×10^7 and 1.01×10^7 bacteria g^{-1} , respectively, in the hindgut. Bacterial abundance was similar in the faeces (1.91×10^7 and 0.87×10^7 bacteria g^{-1}). Taddéi (2006) reported that about 53% of bacteria are digested by *H. atra*. Plotieau *et al.* (2013) reported a higher abundance of 11×10^9 bacteria g^{-1} in the foregut of the coastal species *H. scabra*, which again decreased in the midgut (4×10^9 bacteria g^{-1}) and remained stable in the faeces. They estimated that *H. scabra* digested up to 59% of the ingested bacteria. Amaro *et al.* (2012) reported a value of 80% for *Molpadia musculus* Risso.

Ward-Rainey *et al.* (1996) found the opposite pattern for the two specimens of *H. atra* that they studied. The abundance of bacteria decreased from 3×10^4 and 3×10^6 colony

forming units (cfu) to 3.4×10^3 and 6.2×10^4 cfu, respectively, between the sediment and the foregut, and then increased to 3×10^4 and 1.8×10^6 cfu in the hindgut. However, the low number of replicates means that these results should be considered with caution.

All deposit-feeding holothurian species studied to date regarding the abundance of bacteria have peltate tentacles, except *M. musculus*, which has digitate tentacles (Amaro *et al.*, 2012). No quantitative difference in the ingestion of bacteria has been observed between species with these two tentacle types (Fig. 5). No studies to date have focused on species in Apodida and Dendrochirotida (Table 3). Apodida with pinnate (highly branched) tentacles and suspension-feeding Dendrochirotida with dendritic (ultra-branched) tentacles are likely to show different ingestion rates of bacteria due to their higher tentacle surface area or different feeding behaviour.

Our understanding of the profiles of bacterial abundance in holothurian guts is based on only nine species, and 131 individuals, and additional data on the ingestion of bacteria by holothurians are needed.

Deming & Colwell (1982) suggested that holothurians can enhance the value of ingested bacteria by cultivating them in the foregut. Some bacteria may inhabit internal pouches or be attached to the gut epithelium (Harris, 1993). These enteric bacterial strains could help holothurians to digest large molecules through their secretion of hydrolytic ectoenzymes (Roberts *et al.*, 1991). Amaro *et al.* (2012) studied bacterial diversity through the digestive tract of *M. musculus*. They found that bacterial diversity showed the same pattern as bacterial biomass: there was a higher number of operational taxonomic units (OTUs) in the foregut (< 100 OTUs) than in sediment (28–71 OTUs), and bacterial diversity decreased towards the hindgut (< 70 OTUs). These observations may highlight the ability of holothurians to cultivate particular strains of bacteria in their foregut. As some endosymbiotic bacterial production could be involved, this requires further investigation.

Only a few studies have explored bacterial diversity in the holothurian gut. Five species have been investigated to date, including one deep-sea species. All these studies found Proteobacteria to be the most abundant taxon (Fig. 5), representing 43% of bacteria in the diet of *H. leucospilota* (Zhang *et al.*, 2012), 86% in *H. scabra* (Plotieau *et al.*, 2013), and 55% (Sha *et al.*, 2016) and 89.6% (Gao *et al.*, 2017) in *A. japonicus*. The forms ingested most were γ -proteobacteria and α -proteobacteria for all holothurian species studied. β -proteobacteria, δ -proteobacteria and ε -proteobacteria were also present in some species.

In coastal species, 15 taxa of bacteria were identified, whereas only seven taxa were found in deep-sea species. Five of these taxa (γ -proteobacteria, α -proteobacteria, CFB lineage, Bacteroidetes and Spirochaetes) were shared between coastal and deep-sea species, perhaps because the diversity of deep-sea bacteria is lower than the diversity of coastal bacteria.

Knowledge on the abundance and diversity of bacteria is based on the study of less than 500 specimens from only

Table 3. Food sources recorded as ingested by coastal and deep-sea holothurians

| Food sources ingested | Indicator studied | Authors | Order | Species studied | Habitat | Number of individuals studied |
|--------------------------|-------------------------|----------------------------------|-----------------|--------------------------------|----------|-------------------------------------|
| Living fractions | | | | | | |
| Bacteria | Abundance | Deming & Colwell (1982) | Synallactida | <i>Deima</i> sp. | Deep sea | 2 |
| | | | Persiculida | <i>Pseudostichopus</i> sp. | Deep sea | 3 |
| | | Roberts <i>et al.</i> (2001) | Persiculida | <i>Molpadiodemas villosus</i> | Deep sea | Up to 25 |
| | | | Synallactida | <i>Oneirophanta mutabilis</i> | Deep sea | Up to 25 |
| | | | Elasipodida | <i>Psychropotes longicauda</i> | Deep sea | Up to 25 |
| | | Taddéi (2006) | Holothuriida | <i>Holothuria atra</i> | Coastal | 18 |
| | | | | <i>Holothuria leucospilota</i> | Coastal | 7 |
| | Abundance and diversity | Amaro <i>et al.</i> (2012) | Molpadida | <i>Molpadia musculus</i> | Deep sea | 20 |
| | | Plotieau <i>et al.</i> (2013) | Holothuriida | <i>Holothuria scabra</i> | Coastal | 4 for abundance 30 for diversity |
| | | Ward-Rainey <i>et al.</i> (1996) | Holothuriida | <i>H. atra</i> | Coastal | 2 |
| | | Amaro <i>et al.</i> (2009) | Molpadida | <i>M. musculus</i> | Deep sea | 15 |
| | Diversity | Gao <i>et al.</i> (2017) | Synallactida | <i>Apostichopus japonicus</i> | Coastal | 240 |
| | | Sha <i>et al.</i> (2016) | Synallactida | <i>A. japonicus</i> | Coastal | 30 |
| | | Zhang <i>et al.</i> (2012) | Holothuriida | <i>H. leucospilota</i> | Coastal | 2 |
| | | Taddéi (2006) | Holothuriida | <i>H. atra</i> | Coastal | 18 |
| | | | | <i>H. leucospilota</i> | Coastal | 6 |
| | Abundance and diversity | Belbachir & Mezali (2018) | Holothuriida | <i>Holothuria forskali</i> | Coastal | 10 |
| | | | | <i>Holothuria poli</i> | Coastal | 10 |
| | | | | <i>Holothuria sanctori</i> | Coastal | 10 |
| | | | | <i>Holothuria tubulosa</i> | Coastal | 10 |
| | | | | <i>Cucumaria frondosa</i> | Coastal | 20 |
| | | | Dendrochirotida | <i>Psolus fabricii</i> | Coastal | 30 |
| | | | Holothuriida | <i>H. atra</i> | Coastal | NA |
| | | | | <i>Benthogone rosea</i> | Deep sea | NA |
| | | | Elasipodida | <i>P. longicauda</i> | Deep sea | NA |
| | | | | <i>Molpadia blakei</i> | Deep sea | NA |
| Photosynthetic organisms | Abundance | Hamel & Mercier (1998) | Dendrochirotida | <i>Paroriza pallens</i> | Deep sea | NA |
| | | | | <i>Holothuria theeli</i> | Coastal | 200 |
| | | Sonnenholzner (2003) | Persiculida | <i>P. pallens</i> | Deep sea | 52 |
| | | | | <i>H. atra</i> | Coastal | 6 |
| | | Tyler <i>et al.</i> (1992) | Holothuriida | <i>Stichopus chloronotus</i> | Coastal | 6 |
| | | | | <i>H. forskali</i> | Coastal | 10 |
| | | Uthicke (1999) | Synallactida | <i>H. poli</i> | Coastal | 10 |
| | | | | <i>H. sanctori</i> | Coastal | 10 |
| | Abundance and diversity | Belbachir & Mezali (2018) | Holothuriida | <i>H. tubulosa</i> | Coastal | 10 |
| | | | | <i>C. frondosa</i> | Coastal | 20 |
| | | | | <i>H. forskali</i> | Coastal | 10 |
| | | | | <i>H. poli</i> | Coastal | 10 |
| | | Hamel & Mercier (1998) | Dendrochirotida | <i>H. sanctori</i> | Coastal | 10 |
| | | | | <i>H. tubulosa</i> | Coastal | 10 |
| | | | | <i>C. frondosa</i> | Coastal | 20 |
| | | | | <i>H. forskali</i> | Coastal | 10 |

Table 3. (Cont.)

| Food sources ingested | Indicator studied | Authors | Order | Species studied | Habitat | Number of individuals studied |
|---|--------------------------------------|---------------------------------|------------------|--------------------------------|----------|-------------------------------|
| Non-living fractions Detrital matter | Abundance | Kang <i>et al.</i> (2008) | Holothuriida | <i>H. atra</i> | Coastal | NA |
| | | Khripounoff & Sibuet (1980) | Elasipodida | <i>B. rosea</i> | Deep sea | NA |
| | | | | <i>P. longicauda</i> | Deep sea | NA |
| | | | Molpadida | <i>M. blakei</i> | Deep sea | NA |
| | | | Persiculida | <i>P. pallens</i> | Deep sea | NA |
| | | Sonnenholzner (2003) | Holothuriida | <i>H. theeli</i> | Coastal | 200 |
| | | Tyler <i>et al.</i> (1992) | Persiculida | <i>P. pallens</i> | Deep sea | 52 |
| | | Uthicke (1999) | Holothuriida | <i>H. atra</i> | Coastal | 6 |
| | | | Synallactida | <i>S. chloronotus</i> | Coastal | 6 |
| | | Suchanek <i>et al.</i> (1985) | Holothuriida | <i>Mesothuria verrilli</i> | Deep sea | 5 |
| | | | Elasipodida | <i>Benthodytes lingua</i> | Deep sea | 2 |
| | | | | <i>Psychropotes semperiana</i> | Deep sea | 1 |
| | | Belbachir & Mezali (2018) | Holothuriida | <i>H. forskali</i> | Coastal | 10 |
| | | | | <i>H. poli</i> | Coastal | 10 |
| | | | | <i>H. sanctori</i> | Coastal | 10 |
| Minerals | Abundance and diversity | Costa, Mazzola & Vizzini (2014) | Holothuriida | <i>H. tubulosa</i> | Coastal | 10 |
| | | Khripounoff & Sibuet (1980) | Elasipodida | <i>H. tubulosa</i> | Coastal | 3 |
| | | | | <i>B. rosea</i> | Deep sea | NA |
| | | | | <i>P. longicauda</i> | Deep sea | NA |
| | | | Molpadida | <i>M. blakei</i> | Deep sea | NA |
| | | | Persiculida | <i>P. pallens</i> | Deep sea | NA |
| | | Hammond & Wilkinson (1985) | Apodida | <i>Synaptula lamperti</i> | Coastal | 40 |
| | | Belbachir & Mezali (2018) | Holothuriida | <i>H. forskali</i> | Coastal | 10 |
| | | | | <i>H. poli</i> | Coastal | 10 |
| | | | | <i>H. sanctori</i> | Coastal | 10 |
| | | | | <i>H. tubulosa</i> | Coastal | 10 |
| | | Khripounoff & Sibuet (1980) | Elasipodida | <i>B. rosea</i> | Deep sea | NA |
| | | | | <i>P. longicauda</i> | Deep sea | NA |
| | | | Molpadida | <i>M. blakei</i> | Deep sea | NA |
| | | | Persiculida | <i>P. pallens</i> | Deep sea | NA |
| Plastic particles | Diversity Abundance and diversity | Tyler <i>et al.</i> (1992) | Persiculida | <i>P. pallens</i> | Deep sea | 52 |
| | | Plotieau (2012) | Holothuriida | <i>H. scabra</i> | Coastal | 4 |
| | | Graham & Thompson (2009) | Dendrochirotrida | <i>C. frondosa</i> | Coastal | 42 |
| | | | | <i>Thyonella gemmata</i> | Coastal | 30 |
| | | | Holothuriida | <i>Holothuria grisea</i> | Coastal | 46 |

(Continues)

Table 3. (Cont.)

| Food sources ingested | Indicator studied | Authors | Order | Species studied | Habitat | Number of individuals studied |
|-----------------------|-------------------|---|--------------|-------------------------------|---------|-------------------------------|
| | | | | <i>Holothuria floridana</i> | Coastal | 53 |
| | | Iwalaye, Moodley & Robertson-Andersson (2020) | Holothuriida | <i>Holothuria cinerascens</i> | Coastal | 20 |
| | | Mohsen <i>et al.</i> (2019) | Synallactida | <i>A. japonicus</i> | Coastal | 65 |
| | | Renzi <i>et al.</i> (2018) | Holothuriida | <i>H. tubulosa</i> | Coastal | 30 |

10 species: four from coastal waters and six from the deep sea belonging to Elasipodida, Persiculida, Holothuriida, Molpadida and Synallactida (Table 3). To the extent that these limited findings allow, it can be concluded that holothurians ingest mainly proteobacteria and show a similar distribution of bacteria through the gut across holothurian species, habitat and tentacle type (Fig. 5).

(b) Photosynthetic organisms

Holothurians also consume photosynthetic organisms from both the microphytobenthos (including benthic diatoms, cyanophytes, dinoflagellates, etc.) and macrophytes (macroalgae and phanerogams) for deposit-feeding holothurians, and

phytoplankton (including pelagic diatoms, cyanobacteria, dinoflagellates, etc.) for suspension-feeding holothurians.

The microphytobenthos in shallow-water sediments is mainly dominated by diatoms (Uthicke & Klumpp, 1998; Suzumura *et al.*, 2002; Heil *et al.*, 2004). Taddéi (2006) recorded a mean abundance of 933 diatom cells g^{-1} in the sediment of Reunion Island. In deep-sea sediments, diatoms are also at surprisingly high concentrations (Wood, 1956; Van Iperen *et al.*, 1987; Thiel *et al.*, 1989), with high abundances of cyanophytes and dinoflagellates associated with deposited phytodetritus (Lochte & Turley, 1988; Thiel *et al.*, 1989).

Nine studies have analysed the importance of photosynthetic organisms in the diet of holothurians (Table 3), of which seven

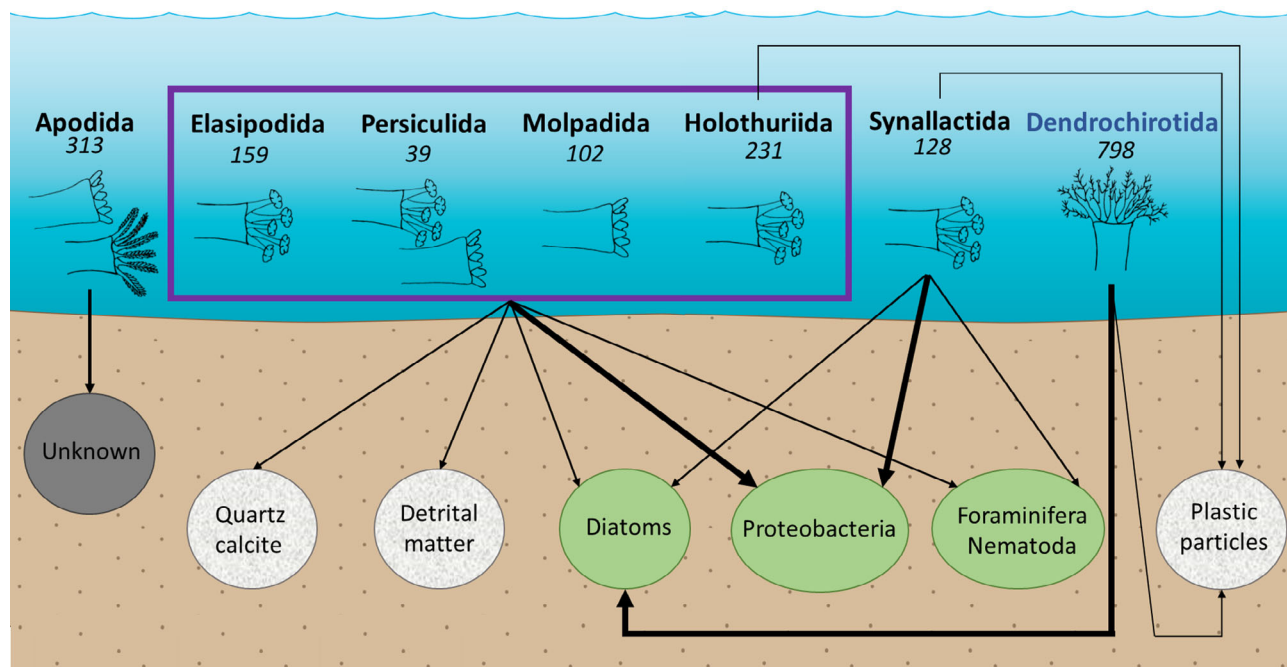


Fig 5. Synthesis of food sources for the seven orders of holothurians. Order colours represent different trophic modes: blue, suspension-feeders (Dendrochirotida); black, deposit-feeders. The number of species per taxon is shown below the order name. The width of arrows corresponds to the proportion of food ingested per source. Colours of food sources vary with food type: green, living fraction; grey, non-living fraction. Tentacle type for each taxon is represented by the drawings (see Fig. 2): digitate (unbranched); peltate (slightly branched); pinnate (highly branched); dendritic (ultra-branched).

focused on 10 coastal species (Hamel *et al.*, 1993; Hamel & Mercier, 1998; Uthicke, 1999; Sonnenholzner, 2003; Taddéi, 2006; Kang *et al.*, 2008; Belbachir & Mezali, 2018) and two studies on four deep-sea species (Khrpounoff & Sibuet, 1980; Tyler *et al.*, 1992).

The most abundant photosynthetic taxa in the diet of holothurians seems to be diatoms (Fig. 5). Hamel *et al.* (1993) analysed the gut contents of *Posolus fabricii* (Düben & Koren), a suspension-feeding species, and found fewer than 6000 pelagic diatom cells in the first centimeter of the foregut. Diatoms were recorded at much higher concentrations in the gut of *Holothuria* (*Selenkothuria*) *theeli* Deichmann (up to 16,500 organisms ml^{-1} ; Sonnenholzner, 2003). For the deep-sea species investigated, the sighting frequency of diatoms in *P. longicauda*, *P. pallens*, *B. rosea*, and *M. blakei* gut ranges between 51 and 84% (Khrpounoff & Sibuet, 1980). Tyler *et al.* (1992) also found a high abundance of diatoms in the gut of *P. pallens*. However, these studies provide no information on whether the diatoms were alive or whether only the siliceous outer skeleton was present.

Photosynthetic organisms may be significant in the diet of holothurians. Fresh organic matter provided by cyanophytes, diatoms, macroalgae and live seagrass leaves represents more than 50% of the diet for *Holothuria poli*, *H. tubulosa*, and *H. forskali* in *Posidonia oceanica* meadows in Algeria (Belbachir & Mezali, 2018). In shallow waters off Ecuador, 35% of the gut content of *H. theeli* consisted of microphytobenthos, with the most dominant taxa being diatoms (91%), followed by cyanophytes and dinoflagellates (Sonnenholzner, 2003). However, these studies did not assess all the possible dietary fractions for holothurians, such as bacteria or minerals, therefore these high proportions may be significant overestimates.

Hamel *et al.* (1993) estimated that phytoplanktonic cells represent between 20 and 50% of the gut content of *P. fabricii* in autumn and winter, increasing during spring to reach 100% in summer in the St. Lawrence Estuary (Canada). They conclude that the ingestion of phytoplanktonic cells, especially pelagic diatoms ingested by this dendrochirotidan holothurian (suspension feeder), depends on seasonal blooms. Similar results were observed for *C. frondosa*, with the proportion of phytoplanktonic cells ingested reaching a maximum in summer (Hamel & Mercier, 1998). Sonnenholzner (2003) investigated the ingestion of photosynthetic organisms in a coastal deposit-feeding species from Holothuriida, finding that the relative abundance of microphytobenthic species in the *H. theeli* gut was significantly higher during the dry season (45.5%) than in the rainy season (20.2%) in the Gulf of Guayaquil. It would be interesting to extend these findings to all fractions present in the holothurian diet to understand which components replace photosynthetic organisms during the cold seasons.

Taddéi (2006) used the concentration of chlorophyll *a* as proxy for photosynthetic organisms in the gut for two coastal deposit-feeders, *H. atra* and *H. leucospilota*. The profile observed was the same as for bacterial abundance: chlorophyll *a* concentration increased from the local sediment ($2.87 \pm 1.17 \mu\text{g g}^{-1}$) to the foregut ($18.74 \pm 7.65 \mu\text{g g}^{-1}$),

decreased in the hindgut ($11.28 \pm 4.61 \mu\text{g g}^{-1}$), and remained at this level in the faeces ($11.34 \pm 4.63 \mu\text{g g}^{-1}$). These observations may indicate that photosynthetic organisms are actively selected for ingestion by these deposit-feeding holothurians.

These studies combined used 378 individuals from six orders of holothurians: Dendrochirotida, Holothuriida, Synallactida, Molpadida, Elasipodida and Persiculida (Table 3). Benthic and pelagic diatoms are a significant dietary component for deposit-feeding and suspension-feeding holothurians respectively, independent of habitat, taxon and tentacle type (Fig. 5). However, the importance of photosynthetic organisms remains unclear without further studies because their biomass is subject to seasonal variations.

(c) Meiofauna

Meiofauna densities in shallow-water sediments range from 3.2 to 1,020.6 individuals 10 cm^{-2} (Guzman, Obando & Cortés, 1987; Armenteros, Creagh & González-Sansón, 2009). In deep-sea sediments, meiofauna densities are lower than in shallow waters, ranging from 15 to 315 individuals 10 cm^{-2} (Coull *et al.*, 1977; Pfannkuche, 1985). In both ecosystems, nematodes and foraminifera are the dominant taxa (Coull *et al.*, 1977; Pfannkuche, 1985; Guzman *et al.*, 1987; Armenteros *et al.*, 2009).

Only seven studies have considered the ingestion of meiofauna by holothurians (Table 3). Nematoda, Copepoda and Foraminifera seem to be the taxa ingested most frequently by holothurians although there are interspecific differences. *Holothuria atra* ingests up to 79% of the meiofauna present in the sediment, with polychaetes the dominant group (Kang *et al.*, 2008). Dissection of *H. theeli* individuals showed that 65% of the gut contents consisted of meiofauna from eight taxa, with a dominance of crustaceans (46%) and foraminifera (35.5%) (Sonnenholzner, 2003). Belbachir & Mezali (2018) demonstrated that the percentage contribution of meiofauna in the holothurian diet can differ among locations and species. They studied the diet of four species *H. poli*, *H. tubulosa*, *H. forskali*, and *Holothuria* (*Platyperona*) *sanctori* Dell Chiaje, at two sites (Stidia and Salamandre in Algeria; separated by less than 20 km). At Salamandre, 20% of the diet of *Holothuria sanctori* consisted of crustaceans, whereas in Stidia crustaceans comprised less than 3%. For the three other species, crustaceans represented only 6% of the diet at Salamandre. The contribution of foraminifera ranged between 3.33% for *H. sanctori* and 15% for *H. forskali* at the same site.

For the deep-sea species *P. longicauda*, *P. pallens*, and *B. rosea*, the sighting frequency of benthic foraminifera reached 100% of the gut content and was 89% for *M. blakei* (Khrpounoff & Sibuet, 1980). The sighting frequency of Nematoda and Copepoda was also high for these four species (96–100% and 0–87%, respectively). Tyler *et al.* (1992) recorded abundant foraminifera in the gut of *P. pallens* but no Nematoda nor Copepoda. Note that the sighting frequency is a poor indicator of the actual contribution of meiofauna to the diet.

Uthicke (1999) compared the abundances of Nematoda, Polychaeta and Harpacticoida between the sediment and the midgut content of *S. chloronotus* and *H. atra*. Nematoda were the dominant taxon in the sediments (31.33 ± 10.90 individuals 2 ml^{-1}), but they were sparsely represented in the midgut of these two species (between 0.50 ± 0.84 and 0.66 ± 1.03 individuals 2 ml^{-1}). Polychaeta were more uncommon in the sediment (9.66 ± 5.31 individuals 2 ml^{-1}) and in the midgut of *S. chloronotus* and *H. atra* (0.17 ± 0.41 and 0.01 ± 0.00 individuals 2 ml^{-1} respectively). Uthicke (1999) concluded that meiofauna play a negligible role in the nutrition of these two species due to the extremely low abundance recorded in the midgut. The ingestion of these two meiofaunal taxa appears to be in proportion to their presence in the sediment, suggesting that meiofauna are probably ingested accidentally with detrital matter (Khrapounoff & Sibuet, 1980; Billett *et al.*, 1988).

One study focused on the ingestion of larval planktonic stages of meiofauna by a suspension-feeding holothurian. Meiofaunal eggs and embryos accounted for less than 15% of the intestinal content of *C. frondosa* in autumn and winter (Hamel & Mercier, 1998). Maximum zooplankton presence peaked just after breeding of broadcast-spawning species. The actual contribution of meiofauna to the diet of suspension-feeding holothurian thus remains largely unknown. As for photosynthetic organisms, the availability of larval planktonic stages of meiofauna varies seasonally and suspension-feeding holothurians that ingest larval planktonic stages of meiofauna in proportion to their presence in the water column must utilise other resources in the autumn and winter in cold or temperate waters. This implies that meiofauna are not an essential dietary component for holothurians.

The ingestion of meiofauna has been investigated in all holothurian orders except Apodida (Table 3). The meiofauna ingested differs between deposit-feeding and suspension-feeding holothurians (Dendrochirotrida), because the latter ingest larval planktonic stages of meiofauna. However, for the other orders Foraminifera, Nematoda, Polychaeta and Crustacea are all ingested at a similarly low rate, independent of tentacle or habitat type (Fig. 5).

(2) Non-living fractions

(a) Detrital matter

Very few studies have considered the detrital matter (Table 3) ingested by coastal and deep-sea holothurians. Shallow-water species can feed on detrital matter from seagrasses or algae. Massin & Jangoux (1976) recorded *H. tubulosa* feeding on detrital seagrass leaves and Costa *et al.* (2014) reported a population of *H. tubulosa* to ingest 30–100% of the detritus of *P. oceanica* meadows, depending on holothurian density. Dead *Posidonia* leaves comprise 0.66–14% of the gut content of *H. tubulosa* and *H. forskali*, respectively (Belbachir & Mezali, 2018). Deep-sea ecosystems also receive phytodetritus, such as dead leaves of seagrasses or algae (Inman & Frautschy, 1965). Using stable

isotope analyses, Suchanek *et al.* (1985) reported that at least two deep-sea species, *Mesothuria verrilli* (Th  el) and *Benthodytes linqua* Perrier R., feed on sediments enriched by decaying seagrasses and consume and metabolise seagrass detritus. Khrapounoff & Sibuet (1980) estimated that *P. pallens*, *M. blackei*, *B. rosea* and *P. longicauda* feed on macrophytic detritus, although with a low sighting frequency of 13%. Together these observations indicate that detrital matter from macrophytes, in both coastal and deep-sea species is ingested relatively rarely (Fig. 5).

In the deep sea, only a small fraction of macroaggregates originating from the euphotic zone reaches the seabed to form detrital matter, with a low increment of 100–150 m day^{-1} (Gooday & Turley, 1990), limiting their accessibility for holothurians (Thurston *et al.*, 1994; Thurston, Rice & Bett, 1998). Most of this fraction is faecal matter, which represents 95% of vertical particle flow (Wiebe, Boyd & Winget, 1976; Honjo, 1978) and constitutes a key component of the abyssal food web (Frankenberg & Smith, 1967). Holothurians are known to be coprophagous (Bakus, 1973; Hauksson, 1979). However, it remains unclear whether coprophagous holothurians feed on faecal pellets intentionally or simply take advantage of a proximate source of concentrated nutrients. Faecal pellets were found in the gut of all specimens of *P. longicauda*, *P. pallens*, *B. rosea*, and *M. blakei* examined (Khrapounoff & Sibuet, 1980). These faecal pellets were from bivalves, pelagic crustaceans, and unidentified sources. Faecal pellets have not been recorded in the diet of coastal species.

Several studies have suggested that detrital matter is a major dietary component for both shallow-water and deep-sea holothurian species (Bordovskiy *et al.*, 1974; Yingst, 1976; Massin, 1982; Moriarty, 1982; Jeffreys *et al.*, 2011). Plotieau (2012) focused on the assimilation of organic compounds from seagrass phytodetritus by *H. scabra*. Experiments showed that *H. scabra* assimilated organic compounds from seagrass leaves, however, this assimilation appeared insufficient to support juvenile growth. Plotieau (2012) therefore suggested that the ingestion of seagrass detritus could be related to the heterotrophic bacteria and microautotrophs attached to them. Indeed, detrital matter, such as phytodetritus, is mainly degraded by bacteria, which colonise it rapidly in both coastal and deep-sea ecosystems (Fenchel & Jorgensen, 1977; Lochte & Turley, 1988; Thiel *et al.*, 1989; Kaiser & Benner, 2008). If correct, this suggests that ingestion of detrital material by holothurians may represent an opportunistic behaviour, to access the bacteria attached to it, rather than detritivorous feeding *per se*.

The ingestion of detrital matter has been relatively poorly studied, with fewer than 100 individuals analysed (Table 3). No studies have focused on Dendrochirotrida (suspension feeders) nor Synallactida. Deep-sea species ingest faecal pellets while coastal species do not. Overall, detrital matter may not be a significant fraction in the diet of holothurians, with the small quantities ingested probably more linked to the presence of bacteria.

(b) Minerals

The non-organic fractions ingested by holothurians are receiving increasing attention, with innovative work suggesting an important role in feeding efficiency and nutritional benefits (Plotieau, 2012).

The sediments ingested by deposit-feeding holothurians contain insoluble clastic products originating from physical and biological breakdown. An analysis of minerals in the sediment ingested by tropical shallow-water holothurians identified both primary and secondary minerals (Plotieau, 2012). The primary minerals were (i) quartz made up of a continuous framework of SiO_4 (identified as the main mineral holothurian gut component in most studies), (ii) calcite, the most stable polymorph of calcium carbonate (CaCO_3), (iii) aragonite, a thermodynamically unstable form of CaCO_3 at standard temperature and pressure, and (iv) magnesian calcite (Ca, MgCO_3 , a variety of calcite containing randomly substituted magnesium carbonate in a disordered calcite lattice that is present in echinoderm skeletons. The secondary minerals were (i) bioclasts, skeletal fragments of marine or land organisms found in sedimentary rocks (mainly composed of aragonite but also magnesian calcite and calcite), (ii) feldspars (KAlSi_3O_8 , $\text{NaAlSi}_3\text{O}_8$, $\text{CaAl}_2\text{Si}_2\text{O}_8$), a group of rock-forming tectosilicate minerals, (iii) hornblende or dark amphibole, an isomorphous mixture of calcium–iron–magnesium silicate, aluminium–iron–magnesium silicate, and iron–magnesium silicate), and (iv) other, trace minerals, mainly zircon.

Three studies analysed the gut contents of shallow-water or deep-sea species with respect to minerals (Table 3; Khripounoff & Sibuet, 1980; Tyler *et al.*, 1992; Belbachir & Mezali, 2018). They identified the presence of mollusc, echinoderm, and sponge ossicles along with various pelagic and benthic foraminifera tests. The mineral fraction in the diets of four shallow-water species can exceed 30%, with less than 12% bivalve shells and 12–24% sponge ossicles (Belbachir & Mezali, 2018). A high sighting frequency of 59–79% was reported for spicules and 100% for coccoliths, in the diets of four deep-sea species (*P. longicauda*, *P. pallens*, *B. rosea*, and *M. blakei*; Khripounoff & Sibuet, 1980). Tyler *et al.* (1992) only rarely recorded spicules or ossicles in the gut of *P. pallens* although coccoliths were abundant.

(c) Plastic particles

Recent decades have revealed the impact of increasing plastic input into the ocean on marine taxa, such as fishes, seabirds, turtles, and cetaceans (Cole *et al.*, 2011), and how plastic particles penetrate the marine trophic web (Ivar do Sul & Costa, 2014). Species belonging to lower trophic levels can ingest high levels of plastic particles because they do not differentiate between these and their preferred food (Renzi *et al.*, 2018).

Several studies have recorded the ingestion of plastic particles (Table 3) in shallow-water deposit-feeding species (Graham & Thompson, 2009; Renzi *et al.*, 2018; Mohsen

et al., 2019) and suspension-feeding species (Graham & Thompson, 2009; Iwalaye *et al.*, 2020). *C. frondosa*, *Holothuria (Semperothuria) cinerascens* (Brandt), *Holothuria grisea* Selenka, *Holothuria floridana* (Pourtales) and *H. tubulosa* were all shown to ingest plastic particles. Mohsen *et al.* (2019) found a lower abundance of plastic in the gut of farmed *A. japonicus* (0–30 particles per individual) than in the local sediments (20–1040 particles kg^{-1}). It is likely that most shallow-water species will ingest plastics if they are present in the sediment (Graham & Thompson, 2009). Recent studies show that, due to the vertical transport of particles from the sea surface to the sea floor, plastic particles are sequestered in deep-sea sediments from the Atlantic, Pacific, and Indian Oceans and the Mediterranean Sea (Van Cauwenberghe *et al.*, 2013; Woodall *et al.*, 2014; Fischer *et al.*, 2015). Plastic particles have been found in the gut of other deep-sea benthic invertebrates including Cnidaria, Echinodermata, Arthropoda (Taylor *et al.*, 2016), and Mollusca (Courteney-Jones *et al.*, 2017). Thus, deep-sea holothurian species may also be exposed to and ingest plastics.

Mohsen *et al.* (2019) also found plastic particles in the coelomic fluid, ranging from 0 to 19 particles per individual. Iwalaye *et al.* (2020) found a similar result in tank experiments with *H. cinerascens* (32–227 microfibrils per individual). In more than half of the individuals studied (57.8%), plastic particles were present in the respiratory tree (0–12 microfibrils per individual). The biological impacts of the transfer of microplastic particles from the holothurian gut to the rest of the body were not analysed further.

Grossmann (2014) and Assidqi (2015) assessed the impact of the ingestion of plastic particles on *H. sanctori* and *H. leucospilota*, respectively. They exposed these holothurians to plastic particles followed by hypoxic conditions to investigate the effects of plastic particle ingestion on resistance to environmental stress. Ingestion of plastic particles did not affect faeces production, evisceration, respiration rate, survival, or behavioural responses in these two species. However, Assidqi (2015) reported a higher susceptibility of *H. leucospilota* to oxygen depletion 60 days after plastic particle ingestion.

Experiments in mesocosms revealed that holothurians appear preferentially to select plastic particles from the sediment (Graham & Thompson, 2009). The authors hypothesised that the larger surface area of the plastic particles reduces the need to shovel or rake for other particles leading to their selection in preference to sand grains. Similar results were found in a natural environment: analyses of sediment sieved through 63–4,000 μm sieves revealed that *H. tubulosa* selects 100–2,000 μm plastic fragments for ingestion (Renzi *et al.*, 2018).

When they reach shallow-water sediments, plastic particles can rapidly be colonised by bacteria (Harrison *et al.*, 2014). Dussud *et al.* (2018) demonstrated that bacterial abundance and diversity is higher on plastic particles than on organic particles. Bacterial communities found on the surface of plastic particles include Proteobacteria, Bacteroidetes (Oberbeckmann *et al.*, 2014; Curren & Leong, 2019), and

Cyanobacteria (Oberbeckmann *et al.*, 2014; Dussud *et al.*, 2018). Other organisms, such as diatoms (Carson *et al.*, 2013; Eich *et al.*, 2015) and dinoflagellates (Masó *et al.*, 2003) can also be attached to plastic particles. All of these organisms are found in the holothurian diet. Finally, because holothurians select more bacteria-rich sediments, they could easily preferentially select plastic particles with bacteria attached, although this requires further investigation.

V. CONCLUSIONS

- (1) Cells on the holothurian tentacle bud epidermis are important for food capture. Ciliated cells are likely to be sensory cells that recognise food particles. Mucous cells may be responsible for the production of a physical barrier on the epidermis. The adhesion of food particles is suggested to be performed by the secretion of type-1 secretory cells; when the tentacles are placed in the oral cavity, adhesion may be removed by the secretion of type-2 secretory cells. Further studies are needed to confirm these hypothesised roles in holothurian taxa.
- (2) Holothurian nutrition appears to involve two main feeding modes: some species are suspension-feeders whereas others are deposit-feeders, with a few species able to shift between suspension- and deposit-feeding. A third feeding pathway may exist, which relies on nutrient transfer from cloacal water retained by the animal to the haemal system.
- (3) Synallactida, Molpadida, Persiculida, Holothuriida and Elaspodida, despite their different tentacle types and the different habitats they colonise, ingest similar proportions of different food types. Their diet seems to be composed mainly of proteobacteria. Diatom ingestion may be high, but its overall contribution remains uncertain as its availability varies seasonally. Other smaller fractions, such as detrital matter, minerals and plastic particles, may be ingested only because they support a high bacterial load.
- (4) Dendrochirotida is the most recent order of holothurians. The species belonging to this order have dendritic tentacles that allow them to act as suspension-feeders. Most species in Dendrochirotida are coastal (more than 98%). Their trophic specialisation presumably reduces interspecific competition for food, and they ingest large quantities of pelagic diatoms. The ingestion of other food fractions has not yet been studied for Dendrochirotida.
- (5) There is very little information on the ingestion of food by Apodida. This is a considerable knowledge gap because Apodida represent 17% of holothurian species. More than half of Apodida species have pinnate tentacles that are not present in other orders.

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