We are IntechOpen, the world's leading publisher of Open Access books Built by scientists, for scientists



122,000





Our authors are among the

TOP 1%





WEB OF SCIENCE

Selection of our books indexed in the Book Citation Index in Web of Science™ Core Collection (BKCI)

Interested in publishing with us? Contact book.department@intechopen.com

Numbers displayed above are based on latest data collected. For more information visit www.intechopen.com



The Thalassinidean Mud Shrimp *Upogebia vasquezi*: Life Cycle and Reproductive Traits on the Amazonian Coast, Brazil

Danielly Brito de Oliveira, Fernando Araújo Abrunhosa and Jussara Moretto Martinelli-Lemos

Additional information is available at the end of the chapter

http://dx.doi.org/10.5772/intechopen.68934

Abstract

The thalassinideans comprise the infraorders Axiidea and Gebiidea, two distinct groups of decapods that have converged morphologically and ecologically as burrowing forms, commonly known as mud lobsters and mud or ghost shrimps. These groups are an important component of the macroinfauna of intertidal and subtidal environments and are distributed throughout the world, with species diversity increasing from high latitudes toward the equator. These species are burrowing benthic decapods, with more than 95% of species inhabiting shallow waters in marine and estuarine environments, exerting considerable influence over the structure of benthic communities through their ability to bioturbate the sediments, with effects on the infauna and seagrasses in coastal environments. Upogebia vasquezi has an ample geographic distribution, it is typically found in rocky outcrops near mangroves. This species reproduces year round, which is subjected to strong seasonal fluctuations in salinity due to the local precipitation regime. The Amazon Macrotidal Mangrove Coast, representing 10% of the Brazilian coastline and encompassing more than 56% of the country's mangrove forests, is a high priority area for conservation. This chapter aims to elucidate the reproductive traits of U. vasquezi with a revision about the known ecological information available for thalassinidean species all over the world.

Keywords: Decapoda, Gebiidea, larval biology, reproduction



© 2017 The Author(s). Licensee InTech. This chapter is distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. (cc) BY

1. Introduction

The Thalassinidea (infraorders Axiidea de Saint Laurent 1979 and Gebiidea de Saint Laurent, 1979) designates a group of decapods popularly known as "mud shrimps," "mud lobsters," "ghost shrimps," or "corruptos" in Portuguese [1–3]. They are among the most common burrowing shrimps frequently found in high densities in coastal and sublittoral sediments, from brackish to euryhaline environments [4]. Most species are marine or estuarine and use sheltered sites as habitats, preferably estuaries, bays, lagoons, beaches, and seas, both in tropical countries and in temperate regions worldwide, and their distribution ranges from shallow mid- and infralittoral to deeper zones [5–7].

These species are also very sensitive to any type of disturbance in their environments, thus serving as effective bioindicators [1]. The potential to accumulate pollutants in their tissues is higher than that of other crustaceans such as crabs and sand crabs, which is useful in the assessment of coastal environments polluted by domestic sewage and industrial waste [1, 8]. On the Amazon Coast, for instance, thalassinidean species have not been found on coastal sites that directly receive untreated domestic effluents, and have only appeared in less anthropized, more preserved regions [9].

Some species also have indirect economic value, as they are used as bait in artisanal and recreational fishing [6, 10–15]. Its capture is usually performed using "prawn pumps," with consequent trampling and digging in several locations, resulting in relevant impacts to the ecosystems where they dwell [16], as well as on the associated biota [17]. Furthermore, thalassinidean populations might occasionally suffer a sharp decrease themselves or even be at risk of extinction [14]. They have been reportedly used as food in some eastern countries, e.g., *Austinogebia edulis* (Ngoc-Ho and Chan, 1992), which is commercialized in Taiwan [18]. On the other hand, they might also cause harm in oyster farming, provoking sediment instability, that impacts on the growth of cultivated organisms or even cause their mortality [19–21].

Despite the ecological relevance of this species in benthic communities of the mid-littoral, very little is known about its biology, especially regarding reproduction and larval development [10, 22–24], mostly due to its cryptic habit and difficulties in capturing specimens [25].

The aim of this chapter is to provide a brief revision on the biology of Axiidea and Gebiidea crustaceans and characterize the Amazon coastal habitats where these organisms are found, with emphasis on *Upogebia vasquezi* (Ngoc-Ho, 1989), one of the most abundant species of this group in the region.

2. Systematics and morphology

The evolutionary position of thalassinidean shrimps inside decapods is still quite controversial, and this is reflected in frequent systematic revisions. These species have already comprised the Infraorder Anomura MacLeay, 1838, together with hermit crabs, porcellanids, and galatids, among other different representatives of this taxon [26–28]. Although this classification was based on the morphological characteristics of adults, some similarities concerning larval morphology were also observed, e.g., the reduction of a pair of marginal bristles of the telson in the zoeal stage [27], which reinforced indications of a relationship between hermit crab species and this group.

Notwithstanding, important distinctions have also been observed, which indicate a strong association between Callianassidae Dana, 1852 and Axiidae Huxley, 1879 and between Upogebiidae Borradaile, 1903 and Laomediidae Borradaile, 1903 with other Anomurans (**Figure 1**), suggesting a inhomogeneous group [29], which would later be called "nephropoidean" and "anomuran" larvae, respectively [30].

Thalassinideans were later considered a distinct group, at the same hierarchical level as anomurans [31], until they were pointed out as a monophyletic taxon, comprising the infraorder Thalassinidea (Latreille, 1831) [32]. The definition of this infraorder was based on some characteristics shared by the species that comprise it, namely, the complex burrow systems they built and the presence of thick feathery bristles on their second pair of pereiopods [32–34].

However, differences observed between two of the main families that comprise Thalassinidea (Callianassidae and Upogebiidae) suggested that they could have distant phylogenetic origins [35]. Thus, the similar habits between these two taxa (reclusive habits, burrows) would have converged throughout their evolutionary history [35]. This assumption corroborated the indications [28] of the existence of two different groups based on larval morphology. This morphological evidence was supported by molecular phylogeny analyses and resulted in the suggestion of dividing this taxon into two infraorders: Axiidea De Saint Laurent, 1979 and Gebiidea De Saint Laurent, 1979 [36, 37], which has been adopted by several authors [38–42]. Another nomenclature was proposed [43] for these taxa: Callianassidea Dana, 1852 and Thalassinidea Latreille, 1831, respectively. However, the names Axiidea and Gebiidea, which were first proposed by Saint Laurent [44], are the most widely accepted and consistently used to designate the two infraorders, which recognizably comprise thalassinidean decapods [42].

According to the most recent classification [40], the following families are included in the infraorder Axiidea: Axiidae Huxley, 1879; Callianassidae Dana, 1852; Ctenochelidae Manning and Felder, 1991; Micheleidae Sakai, 1992; and Strahlaxiidae Poore, 1994; whereas the infraorder Gebiidea is comprised of: Axianassidae Schmitt, 1924; Laomediidae Borradaile, 1903; Thalassinidae Latreille, 1831; and Upogebiidae Borradaile, 1903 [40]. Since 1792, when the first thalassinidean species were described, currently cataloged as *Upogebia pusilla* (Petagna, 1792),



Figure 1. Representative specimens of infraorders Axiidea (*Lepidophthalmus siriboia*) and Gebiidea (*Upogebia vasquezi*) collected in the Amazon coastal region. Scale in millimeters. Photos: Dalila Silva.



Figure 2. *Upogebia vasquezi,* adult female. (A) Dorsal view; (B) detail of the rostrum; (C) detail of part of the abdomen with pleopods and telson, lateral view; (D) embryos adhered to pleopods in the hatching phase, lateral view. Photos: (A) Rory Oliveira; (B)–(D) Danielly Oliveira.

Callianassa tyrrhena (Petagna, 1792), and *Callianassa candida* (Olivi, 1792), information available on this group has increased considerably, mainly over the last 100 years, and Callianassidae, Upogebiidae, and Axiidae are the most extensively studied ones [7].

Thalassinidean decapods encompass a relatively small number of species, with approximately 646 catalogued species [38, 39]. This number has recently increased to approximately 674 species, with 465 Axiidea and 209 Gebiidea [45]. Morphologically, these organisms share characters such as the presence of a fairly calcified carapace, symmetrical and extended, while the abdomen is feebly calcified, ending on a well-developed tail fan (telson + uropods) [46]. Some

species are more similar to lobsters with a highly calcified exoskeleton (e.g., Thalassinidae and Axiidae); while others have a more elongated body and a slightly calcified exoskeleton, better adapted to the "burrowing" life style (e.g., Callianassidae) [45].

Differences between the representatives of the infraorders Axiidea and Gebiidea are mainly the shape of the anterior part of the carapace, the structure of appendages, and larval morphology [30, 47]. Gebiidea have a chelated or subchelated first pair of pereiopods, and the second pair is subchelated or simple (never are both pairs chelated), whereas Infraorder Axiidea has the two first pairs of pereiopods chelated [40]. In addition, Axiidea are frequently heterochelic, as opposed to Gebiidea, whose first pair of pereiopods (chelipods) are of the same size, as can be observed in the two species frequently found on the Amazon coast: *Lepidophthalmus siriboia* (Axiidea) and *U. vasquezi* (Gebiidea) (**Figure 1**).

U. vasquezi has a triangular rostrum, whose lateral edges are nearly straight and longer than the ocular peduncles, with presence of postocular spine [47]. The abdomen is robust, broader than long (**Figure 2A**), and the entire body is adorned with bristles, from the anterior portion, on the cephalic appendages (**Figure 2B**), to the abdominal appendages (pleopods) and telson (**Figure 2C**). Females carry the eggs on the pleopods until hatching, in zoea I stage (**Figure 2D**), with approximately 0.88 mm of carapace length [48].

3. Distribution

Thalassinideans are distributed around the world, with a higher concentration of species in the regions located at low latitudes; e.g., the three major groups Callianassidae, Upogebiidae, and Axiidae occur mainly between latitudes 25°N and 10°N and between 0° and 15°S [7]. The highest percentage of species (36.5%) was recorded in the Western Indian-Pacific, but they are also found in the eastern and western portions of the Atlantic, including the Caribbean Sea and the Gulf of Mexico; as well as in the Mediterranean region [41, 49, 50].

They are mostly marine species, usually found in sheltered habitats, such as estuaries, bays, lagoons, beaches, and seas, both in tropical countries and in temperate regions worldwide, and their distribution ranges from shallow mid- and infralittoral to deeper zones [5–7]. Most species (95%) occur in shallow waters (0–200 m), and few have been found in depths lower than 2000 m [7, 49].

In Brazil, the occurrence of 43 species has been registered [47, 51], and they are popularly known as "corruptos" [1]. Their distribution ranges from Amapá (Northern region) to Rio Grande do Sul (Southern Region) in different habitats, such as bottoms of calcareous waters, coral reefs, rocks, sand, mud, near seaweed meadows, surrounding mangrove vegetation, and in deeper waters on the continental shelf and slope, down to a depth of 820 m [47].

Species of only two families have occurred on the Amazon coast: Callianassidae (Axiidea) and Upogebiidae (Gebiidea) [47], typically found in very shallow waters (down to a depth of 2 m) of estuarine regions with decreased salinity [7, 47]. Only 13 species has been recorded in the State of Pará [47, 52, 53] Brazil (**Table 1**).

Infraorder	Family	Gender	Species	Geographical distribution
Gebiidea (De Saint Laurent, 1979)	Upogebiidae Borradaile, 1903	<i>Upogebia</i> (Leach, 1814)	U. acanthura (Coelho, 1973)	Western Atlantic: Gulf of Mexico and the Bahamas, Antilles, northern South America and Brazil – from Pará to Pernambuco and along Espírito Santo.
	U. brasiliens. (Holthuis, 1 U. marina (C 1973) U. paraffinis (Williams, 1		U. brasiliensis (Holthuis, 1956)	Western Atlantic: Belize, French Guiana, Suriname, and Brazil – from Pará to Santa Catarina.
			<i>U. marina</i> (Coelho, 1973)	Western Atlantic: Venezuela and Brazil – from Pará to São Paulo.
		<i>U. paraffinis</i> (Williams, 1993)	Western Atlantic: Brazil – Pará and from Ceará to Paraná.	
			U. vasquezi (Ngoc- Ho, 1989)	Western Atlantic: south of Florida, Bahamas, Central America, Antilles, Venezuela, and Brazil – from Pará and Maranhão to São Paulo.
Axiidea (De Saint Laurent, 1979)	Callianassidae (Dana, 1852)	<i>Corallianassa</i> Manning, 1987	<i>C. longiventris</i> (A. Milne Edwards, 1870)	Western Atlantic: Florida, Bermuda, Antilles, and Brazil – Rocas, and from Pará to Pernambuco.
		<i>Cheramus</i> Bate, 1888	C. marginatus (Rathbun, 1901)	Western Atlantic: Florida, Antilles, and Brazil – from Amapá to Rio de Janeiro.
		<i>Callichirus</i> (Stimpson, 1866)	C. major (Say, 1818)	Western Atlantic: North Carolina to Florida, Gulf of Mexico, Venezuela and Brazil – Rio Grande do Norte, Pernambuco, and from Bahia to Santa Catarina.
		<i>Lepidophthalmus</i> (Holmes, 1904)	<i>L. siriboia</i> (Felder and Rodrigues, 1993)	Western Atlantic: Florida, Gulf of Mexico, Antilles, and Brazil – from Pará to Bahia.
		Neocallichirus (Sakai, 1988)	N. grandimana (Gibbes, 1850)	Western Atlantic: Florida, Gulf of Mexico, Bermuda, Antilles, and South America, and Brazil – from Pará to Bahia.
		<i>Sergio</i> (Manning and Lemaitre, 1994)	S. guara (Rodrigues, 1971)	Western Atlantic: Brazil – from Pará to São Paulo.
		<i>Marcusiaxius</i> (Rodrigues and Carvalho, 1972)	<i>M. lemoscastroi</i> (Rodrigues and Carvalho, 1972)	Western Atlantic: Central America, Colombia, Venezuela, and Brazil – Amapá, Pará, and Ceará.
		<i>Dawsonius</i> (Manning and Felder, 1991)	D. latispinus (Dawson, 1967)	Western Atlantic: Florida and Brazil – from Amapá to Alagoas.



The distribution of *U. vasquezi* encompasses the Western Atlantic, ranging from Florida and several islands in the Caribbean Sea region (such as Aruba and Bonaire), through Central America, Bahamas, Dominican Republic, Barbuda, Antigua, Barbados, Tobago, Mexico, Panama, Venezuela, to Brazil: from Pará to São Paulo [47, 50, 53, 54] (**Figure 3**). It occurs in shallow waters, mostly down to depths of 2 m [7], dwelling in burrows excavated in the sediment of the intertidal zone [47].



4. Amazon coastal habitats

The coast of Pará accounts for 6.6% of the Brazilian coast, and the extension of mangrove area covers approximately 2176 km² [55] in the northeastern portion of the state alone, corresponding to 1.2% of the global mangrove area (181,000 km²) [55–57]. The region known as "Amazon Macrotidal Mangrove Coast" extends from Marajó Bay (PA) to São José Bay (MA), totalling 650 km of coast [55]. It is characterized by a wide coastal plain and an extensive adjacent continental shelf (~200 km wide), which is irregular and cut through by several estuaries [55]. This region is subjected to a quite dynamic tidal regime and currents, with semidiurnal macrotides ranging from 4 to 7.5 m of amplitude [48, 55, 58].

Several habitats comprise the Amazon coast, e.g., sandy beaches and estuaries, temporary tide pools, muddy coastal plains, and rocky outcrops, where several decapod species are found, including thalassinideans. For instance, *L. siriboia* occurs mostly on beaches with sandy-muddy sediment [9, 59], *Callichirus major* might be found in habitats similar to *L. siriboia*, on the most exposed portion of the beach (Danielly B. Oliveira, personal observation), whereas *U. vasquezi* inhabits burrows sheltered under rocky outcrops with sandy-muddy sediment [48, 53].

5. Burrows

One of the characteristics shared by thalassinidean shrimps is their reclusive lifestyle, with the construction of burrows, which are among the deepest and most complex systems recorded in transitional marine environments [6]. They are built on sandy and muddy surfaces of the coastal zone, serving as shelter and protection against predators, as well as feeding and reproduction sites [1, 6, 25]. Thanks to the fossilization of burrows on these species, paleontologists gathered important indications about ancient coastlines [6].

Thalassinidean burrows are considered unique environments, whose physical-chemical conditions are strongly influenced by the behavior of these species, mostly due to their bioturbation activities, which have effects on nutrient cycling (for example, see [60–62]) and also ensure high availability of dissolved oxygen, aside from providing protection from the direct action of waves [1]. The process of burrow construction increases the inner surface area of the sediment, in the oxygenated water-sediment interface [33, 63], and causes physicochemical changes, thus increasing the metabolic activity in the sediment [64].

In regions with intense thalassinidean aggregations, there is a change in the sediment structure, which becomes more porous and has increased concentration of smaller particles and organic matter [65]. Such conditions influence the structure of the local benthic community [66], creating, changing, and maintaining a mosaic of habitats for a wide range of organisms [67].

Burrow structure is specific for each species, and it is related to their feeding mode, as well as to environmental conditions and the population density of these crustaceans in their habitats [63]. Externally, they might be divided into two main types: burrows with and without sediment heaps around their openings [63]. Regarding shape, they might be built in a single U- or Y-shaped tunnel, or in several sediment layers or branched, deep tunnels [63], which might be interconnected and might shelter at least one specimen [68].

Most *Upogebia* species, for example, live in relatively shallow, U-shaped burrows, e.g., *U. africana* (Ortmann, 1894) [69], *U. stellata* (Montagu, 1808) and *U. deltaura* (Leach, 1815) [70], *U. tipica* (Nardo, 1869) [71] *U. noronhensis* (Fausto-Filho, 1969) [23] *U. major* (De Haan, 1841) [66] and *U. vasquezi* [72]). Some species of this group build Y-shaped burrows in sandy-muddy habitats, like *U. omissa* (Gomes Corrêa, 1968), *U. yokoyai* (Makarov, 1938), and *U. carinicauda* (Stimpson, 1860) [25, 73, 74].

Burrows of *U. vasquezi* are built in predominantly sandy-muddy sediment, with small and abundant rock fragments, located below extensive outcrops comprised of rocks of several sizes. These outcrops are located near mangroves on some Amazon estuarine beaches, and

are submerged during high tide and exposed during low tide (Danielly B. Oliveira, personal observation). Regarding its morphology, the burrows of *U. vasquezi* are relatively shallow and U-shaped [72], and in its natural habitat, it is possible to observe the opening of the burrows excavated in the sediment by just removing some rocks from the outcrop (Danielly B. Oliveira, personal observation).

6. Ecological relationships

Several organisms associated to thalassinideans occur inside their burrows, using them as shelter and also for feeding. Examples are some alpheid shrimp species, as *Leptalpheus axianassae* (Dworschak and Coelho, 1999), the crabs *Pinnixa gracilipes* (Coelho, 1997); *Pinnixa transversalis* (H. MilneEdwards and Lucas, 1842); and *Austinixa aidae* (Righi, 1967), as well as invertebrates phoronids, polychaetes, nemertins, copepods, and gobiidae fish [20, 68, 75–80]. Some species might be parasitic to thalassinideans; e.g., isopods are prevalent ectoparasites of *Upogebia* (Leach, 1814) (for example, see [4, 5, 81–83]). There are also endoparasites of thalassinids, such as trematode cysts, Acanthocephala [4], and copepods infesting gills, pereiopods, and egg masses (e.g., [68]).

In addition to these species, there is a varying fauna that coexists in the sandy and muddy plains inhabited by thalassinideans, not necessarily inside the burrows, which are also influenced by the dynamics of "corruptos" (mud crabs). For example, gastropods, bivalves, echiura, echinodermata, polychaetes, and anemones comprise of an important fraction of the macrozoobenthos biomass in *Upogebia issaeffi* habitats (Balss, 1913) [84]. Stomatopods species, bivalves, and echiura, along with other sympratic thalassinids (e.g., *Upogebia* sp. and *Lepidophthalmus* sp.) [80], are macrofauna also associated to habitats of *Axianassa australis* (Rodrigues and Shimizu, 1992) on the tropical beaches of the Brazilian coast [80].

The invading intertidal fish species *Omobranchus punctatus* (Valenciennes, 1836) (Osteichthyes: Blenniidae) and the shrimp *Alpheus estuariensis* (Christoffersen, 1984) (**Figure 4**) have been



Figure 4. Species inhabiting burrows of *Upogebia vasquezi* in the Amazon estuarine region. (A) *Omobranchus punctatus;* (B) *Alpheus estuariensis.* Photos: Rory Oliveira.

found in burrows of *U. vasquezi* on the Amazon coast (Danielly B. Oliveira, personal observation). Gobbidea fish are common dwellers of burrows of Upogebiidea, feeding mostly on small crustaceans [85].

7. Reproduction and life cycle

7.1. Larval biology

Information available on larval biology of thalassinidean shrimps (infraorders Axiidea and Gebiidea) is relatively scarce, mainly because the development of larval stages of most species have not yet been described, thereby hindering the identification of specimens captured in natural environments. Among the available descriptions, many of them are based on specimens collected in zooplankton samples, which might mislead species identification [86], and in others, the characterization of the different development stages [63] are frequently poorly understood.

Only 12.5% of thalassinidean species and 25% of thalassinidean genera are estimated to have a known larval development, and *Upogebia* is the group with the highest number of species whose larvae have been described [87]. In absolute numbers, approximately 80 species (including unidentified morphotypes of some genera) have had their larval cycle partially or completely described [87]. Some of these species have also had their post-larval stage (or first juvenile stage) morphologically described (e.g., *Upogebia affinis* (Say, 1818): [88]; *U. paraffinis* (Williams, 1993): [89]; *L. siriboia*: [90]).

Regarding the 13 thalassinidean species whose occurrence in the Amazon coastal region has been recorded, only four have already had their larval and/or juvenile development stages partially or completely described: *C. major, L. siriboia, U. paraffinis,* and *U. vasquezi* (**Table 2**).

The larval phase of thalassinideans is predominantly planktonic, and in most species, it is the only life-cycle stage where they remain outside their burrows [6]. The complete suppression of larval stages during development is only known for *Upogebia savignyi* (Strahl, 1862), a sponge commensal [91].

Species	Developmental stages	References			
Callichirus major	ZI–ZV, M	[95]			
Lepidophthalmus siriboia	Prezoea, ZI–ZIII, M, JI	[90]			
Upogebia paraffinis	ZI–ZV, M	[89]			
Upogebia vasquezi	ZI–ZIV, M	[48]			

Note: Z, Zoea; M, Megalopa; J, juvenile. Roman numbers represent the number of developmental stages described.

Table 2. Thalassinidean species with occurrence on the coast of Pará whose larval and/or juvenile development stages have already been partially or completely described.

Thalassinideans have varying developmental patterns, generally one to eight zoeal stages [86, 92]. Among Axiidea, a reduced larval cycle is common, with two to three development stages (e.g., *Callichirus kraussi* (Stebbing, 1900) as *Callianassa kraussi* [93]; *Pestarella tyrrhena* (Petagna, 1792) as *C. tyrrhena* [22]; *Lepidophthalmus sinuensis* (Lemaitre and Rodrigues, 1991), *Lepidophthalmus louisianensis* (Schmitt, 1935) [94], and *L. siriboia* [90]). Some species in this group also have a long planktonic larval development, such as *C. major* and *Callichirus isla-grande* (Schmitt, 1935), which undergo 4–5 zoeal stages [95, 96], or *Boasaxius princeps* (Boas, 1880) and *Nihonotrypaea petalura* (Stimpson, 1860), with 7–8 zoea [97, 98].

Regarding Gebiidea, a long larval development is frequent: *Naushonia crangonoides* (Kingsley, 1897) undergoes six to seven zoeal stages until it reaches the post-larval stage [99]; and *A. australis* (Rodrigues and Shimizu, 1992) shows up to eight zoeal stages [100]. The most common larval development pattern of *Upogebiidea* is the presence of three to four zoeal stages (e.g., *Upogebia kempi* (Shenoy, 1967) [101]; *Upogebia darwinii* (Miers, 1884) [102]; *U. major* [10]; *U. pusilla* [103]; *U. issaeffi* [104]; *U. yokoyai* [63]; *U. vasquezi* [48]).

The life cycle of *U. vasquezi* larvae has four zoeal stages [48]. When immature, the eggs of this species are yellowish (**Figure 5A**), their color start becoming more orange by the end of embryo development, in the hatching stage, when the eyes also become visible (**Figure 5B**). Larvae hatch in Zoea I, going through three other zoeal stages and one megalopa until reaching the first juvenile stage (**Figure 5C–H**).

Only *C. major, L. siriboia,* and *U. vasquezi* larvae have already been found in estuarine zooplankton samples from the Amazon coast [53]. Among the studies conducted with these species in the region, the taxonomic studies stand out, namely the morphological description of larval developmental stages of *L. siriboia* [90], as well as the description of mouth appendages and stomachs of larvae [105], analysis of the lecithotrophic behavior of this species during larval cycle [24], and abundance of larvae in the estuarine zooplankton [53]. With regard to *C. major*, the importance of feeding during larval development has been analyzed (as opposed to the lecithotrophic behavior of *L. siriboia*) [106], as well as the abundance of estuarine planktonic larvae throughout an annual cycle [53].

U. vasquezi was the most studied thalassinidean species in the region regarding larval biology, with description of larval morphology [48, 107], analysis of the effect of salinity on survival and duration of larval stages, its implication on larval migration [108], and occurrence of planktonic larvae along a salinity gradient in the Amazon estuary [53].

7.2. Effects of biotic factors on larval development

Diverse environmental factors influence developmental rates, number of stages, and survival of larvae of marine invertebrates [109]. Temperature and salinity are among the physicochemical factors that have a higher influence on survival and larval development of marine decapods [110]. Temperature might influence the growth of decapods during different life-cycle phases, from larvae and post-larvae to juveniles and adults [111], and trigger the acceleration or decrease of larval developmental rate, and impact metabolism and development, as well as



Figure 5. Developmental stages of *Upogebia vasquezi*. (A) Eggs in the initial developmental stage; (B) eggs in the final developmental stage; (C) Zoea I; (D) Zoea II; (E) Zoea III; (F) Zoea IV; (G) Megalopa (without antennas); (H) Juvenile I. Photos: Danielly Oliveira.

the seasonality of larvae emergence in some plankton species [110]. For instance, temperature mainly influences the duration of decapod larval stages, which are prolonged in stressful situations (for example, see [22, 112, 113]).

Saline concentration is generally constant in open sea, whereas it might seasonally fluctuate in coastal and estuarine zones, both regionally and locally [110]. Hence, salinity is considered an

ecological and physiological factor of extreme importance for species in these environments [110], with impact on the development, survival, feeding, and growth rate, as well as on shedding cycles, metabolic rates, and behavior [113].

The reproductive behavior (life-cycle strategies) of decapods might also be influenced by salinity. Most estuarine species export their larvae to marine coastal zones, where salinity is more stable and, on average, higher than in the parental habitat, whereas others retain their initial larval stages inside the estuarine environment [112, 113]. For instance, some typical estuarine crabs increase their swimming activity in higher salinities to avoid being transported outside the estuary [114].

Studies analyzing the effect of salinity on larval development of decapods are also useful to identify which reproductive strategy is adopted by the species (either retention or exportation) due to the fact that saline limits tolerated by decapod larvae under experimental conditions coincide with their distribution along salinity gradients in the field [113]. In the coastal region of Pará, the effect of salinity on larval development of the crabs *Ucides cordatus* (Linnaeus, 1763), *Uca vocator* (Herbst, 1804), and *Uca rapax* was analyzed in the laboratory, obtaining decreased survival rates under lower salinity conditions, thus indicating a strategy of larval dispersal and exportation [115–118].

7.3. Reproduction, dynamics, and secondary production

Studies on the population dynamics and reproductive biology of thalassinideans have been developed in several locations worldwide, thus contributing to understanding the life cycle of these species (for example, see [14, 23, 80, 119–126]). Most of these studies were conducted in temperate and subtropical regions and few have shown estimates of population dynamic parameters for this group. On the Amazon coast, only the population dynamics of *L. siriboia* has been studied [59].

Secondary production might be defined as the production of biomass carried out by heterotrophic organisms, including animals, fungi, and heterotrophic bacteria; it represents an estimated biomass made available for higher trophic levels [127]. Decapod crustaceans have a crucial contribution to secondary production in the habitats they inhabit. For example, even though their abundance is lower than that of other invertebrates, they account for an important fraction of productivity in coral reef ecosystems [128] and on sandy beaches at different latitudes [129].

Secondary production estimates are still quite scarce, mostly in the equatorial region (between latitudes 5°S and 5°N), with absence of studies on benthic macrofaunal populations of sandy beaches [130]. Only 12 decapod populations have been studied [130] at higher latitudes, on tropical and subtropical beaches, including the thalassinids *U. pusilla* [4] and *C. major* [131, 132]. In Brazil, studies of this type have only been conducted in the Southern and Southeastern regions (for example, see [132–139]).

The capture of mud shrimps (Axiidea and Gebiidea) might cause changes in the target species and habitat and might influence resident communities and cause indirect effects on sediment structure [12, 13]). Excessive fishery efforts might lead to overexploitation of naturally abundant populations or even to the total disappearance of some species [12, 14]. Management plans and

efforts for the conservation of these species and recovery of their habitats must be based on their regional population and reproductive characteristics [14]. Thus, studies that investigate population dynamics and reproductive biology of thalassinideans in several locations are of utmost importance, especially in priority conservation areas.

Despite the importance of thalassinidean species on Amazon coastal habitats, very little are known on their ecology, mostly regarding burrow morphology, physiology, population dynamics, behavior, and larval description.

Acknowledgements

The authors are grateful for the photos gently given by the colleagues: Dalila C. Silva (MSc) and Rory R. S. Oliveira (MSc). We also appreciate the valuable suggestions of the reviewers, which greatly improved the chapter. This study is part of the Ph.D. thesis of the first author (Danielly B. Oliveira) and was funded by the Brazilian National Research Council (CNPq) (Grant 553106/2005-8 to JMML), the Brazilian Higher Education Training Program (CAPES), and the Brazilian Carcinology Society (Grant SBC 01/2012). We also thank Daniela Tannus for translation of the original manuscript.

Author details

Danielly Brito de Oliveira^{1,3*}, Fernando Araújo Abrunhosa² and Jussara Moretto Martinelli-Lemos³

*Address all correspondence to: danybrito@gmail.com

1 Center for Research and Management of Fishing Resources of the North Coast, Chico Mendes Institute for the Biodiversity Conservation (CEPNOR/ICMBIO), Belém, Pará, Brazil

2 Carcinology Laboratory, Federal University of Pará, Bragança, Pará, Brazil

3 Laboratory for Fishery Biology and Management of Aquatic Resources, Ecology of Amazonian Crustaceans Research Group (GPECA), Federal University of Pará, Belém, Pará, Brazil

References

- Rodrigues SA, Shimizu RM. Autoecologia de *Callichirus major* (Say, 1818). In: Absalão RS, Esteves AM, editors. Ecologia de praias arenosas do litoral brasileiro. Rio de Janeiro, UFRJ. Vol. III. Oecologia Brasiliensis. 1997. pp. 155-170
- [2] Mclauhghlin PA, Camp DK, Angel MV, Bousfield EL, Brunei P, Brusca RC, Cadien D, Cohen AC, Conlan K, Eldredge LG, Felder DL, Goy JW, Haney T, Hann B, Heard RW,

Hendrycks A, Hobbs HH, Holsinger JR, Kensley B, Laubitz DR, Lecroy SE, Lemaitre R, Maddocks RF, Martin JW, Mikkelsen P, Nelson E, Newman WA, Overstreet RM, Poly WJ, Price WW, Reid JW, Robertson A, Rogers DC, Ross A, Schotte M, Schram FR, Shih C-T, Watling L, Wilson GDF. 2005. Common and scientific names of Aquatic Invertebrates from the United States and Canada: Crustaceans. Bethesda, Maryland: American Fisheries Society Special Publication 31. pp. 209-325

- [3] Kornienko ES. Burrowing shrimp of the infraorders gebiidea and axiidea (Crustacea: Decapoda). Russian Journal of Marine Biology. 2013;**39**(1):1-14
- [4] Dworschak PC. The biology of *Upogebia pusilla* (Petagna) (Decapoda, Thalassinidea). Growth and production. PSZNI Marine Ecology. 1988;9(1):51-77
- [5] Coelho PA, Rattacaso MCA. Revisão das espécies de *Upogebia* encontradas em Pernambuco, Brasil (Crustacea, Decapoda, Thalassinidea). Revista Brasileira de Zoologia. 1988;5(3): 381-392. (Cessou em 2008. Cont. ISSN 1984-4670 Zoologia)
- [6] Rodrigues SA, Pezzuto PR. Infraordem Thalassinidea (corruptos). In: Buckup L, Bond-Buckup G, editors. Os crustáceos do Rio Grande do Sul. Porto Alegre: Universidade/ UFRGS; 1999. pp. 328-335
- [7] Dworschak PC. Global diversity in the Thalassinidea (Decapoda). Journal of Crustacean Biology. 2000;**20**(2):238-245
- [8] Ayón-Parente M, Hendrickx ME, Ríos-Jara E, Salgado-Barragán J. Records of mud shrimps (Crustacea: Decapoda: Axiidea and Gebiidae) from Pacific Mexico. Journal of the Marine Biological Association of the United Kingdom. 2014;94(2):369-388
- [9] Silva DC, Martinelli-Lemos JM. Species composition and abundance of the benthic community of Axiidea and Gebiidea (Crustacea: Decapoda) in the Marapanim Bay, Amazon estuary, northern Brazil. Zoologia. 2012;**29**:144-158
- [10] Konishi K. Larval development of the mud shrimp Upogebia (Upogebia) major (De Haan) (Crustacea: Thalassinidea: Upogebiidae) under laboratory conditions, with comments on larval characters of thalassinid families. Bulletin of National Research Institute of Aquaculture. 1989;15:1-17
- [11] Hodgson AN, Allanson BR, Cretchley R. The exploitation of *Upogebia africana* (Crustacea: Thalassinidae) for bait in the Knysna Estuary. Transactions of the Royal Society of South Africa. 2000;55(2):197-204
- [12] Souza JRB, Borzone CA. A extração de corrupto, *Callichirus major* (Say) (Crustacea, Thalassinidea), para uso como isca em praias do litoral do Paraná: As populações exploradas. Revista Brasileira de Zoologia. 2003;20(4):625-630. (Cessou em 2008. Cont. ISSN 1984-4670 Zoologia)
- [13] Contessa L, Bird FL. The impact of bait-pumping on populations of the ghost shrimp *Trypaea australiensis* Dana (Decapoda: Callianassidae) and the sediment environment. Journal of Experimental Marine Biology and Ecology. 2004;304:75-97

- [14] Botter-Carvalho ML, Santos PJP, Carvalho PVVC. Population dynamics of *Callichirus major* (Say, 1818) (Crustacea, Thalassinidea) on a beach in northeastern Brazil. Estuarine, Coastal and Shelf Science. 2007;71:508-516
- [15] Conides AJ, Nicolaidou A, Apostolopoulou M, Thessalou-Legaki M. Growth, mortality and yield of the mudprawn *Upogebia pusilla* (Petagna, 1792) (Crustacea: Decapoda: Gebiidea) from western Greece. Acta Adriatica. 2012;53(1):87-103
- [16] Brown AC, McLachlan A. Sandy shore ecosystems and the threats facing them: some predictions for the year 2025. Environmental Conservation. 2002;**29**(1):62-77
- [17] Wynberg RP, Branch GM. Trampling associated with bait-collection for sandprawns *Callianassa kraussi* Stebbing: Effects on the biota of an intertidal sandflat. Environmental Conservation. 1997;24(2):139-148
- [18] Shy JYA, Chan TY. Complete larval development of the edible mud shrimp Upogebia edulis Ngoc-Ho and Chan, 1992 (Decapoda: Thalassinidea: Upogebiidae) reared in the laboratory. Crustaceana. 1996;69(2):175-186
- [19] Dumbauld BR, Armstrong DA, Feldman KL. Life-history characteristics of two sympatric thalassinidean shrimps, *Neotrypaea californiensis* and *Upogebia pugettensis*, with implications for oyster culture. Journal of Crustacean Biology. 1996;16(4):689-708
- [20] Felder DL. Diversity and ecological significance of deep-burrowing macrocrustaceans in coastal tropical waters of the Americas (Decapoda: Thalassinidea). Interciencia. 2001;26(10):440-449
- [21] Dumbauld BR, Wyllie-Echeverria S. The influence of burrowing thalassinid shrimps on the distribution of intertidal seagrasses in Willapa Bay, Washington, USA. Aquatic Botany. 2003;77:27-42
- [22] Thessalou-Legaki M. Advanced larval development of *Callianassa tyrrhena* (Decapoda: Thalassinidea) and the effect of environmental factors. Journal of Crustacean Biology. 1990;10(4):659-666
- [23] Candisani LC, Sumida PYG, Pires-Vanin AMS. Burrow morphology and mating behaviour of the thalassinidean shrimp *Upogebia noronhensis*. Journal of the Marine Biological Association of the United Kingdom. 2001;81(3795):1-5
- [24] Abrunhosa FA, Simith DJB, Palmeira CAM, Arruda DCB. Lecithotrophic behaviour in zoea and megalopa larvae of the ghost shrimp *Lepidophthalmus siriboia* Felder and Rodrigues, 1993 (Decapoda: Callianassidae). Anais da Academia Brasileira de Ciências. 2008;80(4):639-646
- [25] Coelho VR, Cooper RA, Rodrigues SA. Burrow morphology and behavior of the mud shrimp *Upogebia omissa* (Decapoda: Thalassinidea: Upogebiidae). Marine Ecology Progress Series. 2000;200:229-240
- [26] Borradaile LA. On the classification of the Thalassinidea. Annals and Magazine of Natural History. 1903;12(7):534-551

- [27] Hart JFL. Larval and adult stages of British Columbia Anomura. Canadian Journal of Research. 1937;15(10):179-220
- [28] Mclauhghlin PA. Comparative Morphology of Recent Crustacea. San Francisco: W.H. Freeman and Company; 1980. p. 177
- [29] Gurney R. Notes on some Decapod Crustacea from the Red Sea. VI–VIII. Proceedings of the Zoological Society of London. 1938;108B:73-84
- [30] Gurney R. Larvae of Decapod Crustacea. London: Ray Society; 1942. p. 306
- [31] Burkenroad MD. The higher taxonomy and evolution of Decapoda (Crustacea). Transactions of the San Diego Society of Natural History. 1981;**19**(17):251-268
- [32] Poore GCB. A phylogeny of the families of Thalassinidea (Crustacea: Decapoda) with keys to families and genera. Memoirs of the Museum of Victoria. 1994;**54**:79-120
- [33] Griffis RB, Suchanek TH. A model of burrow architecture and trophic modes in thalassinidean shrimp (Decapoda: Thalassinidea). Marine Ecology Progress Series. 1991;79:171-183
- [34] Martin JW, Davis GE. An Updated Classification of the Recent Crustacea. Natural History Museum of Los Angeles Country. Los Angeles, California: Science Series; 2001;**39**:47-48
- [35] Saint Laurent M. Sur la systématique et la phylogénie des Thalassinidea: définition des familles des Callianassidae et des Upogebiidae et diagnose de cinq genres nouveaux (Crustacea Decapoda). Comptes rendus hebdomadaires des séances de l'Académie des sciences, D. 1973;277:513-516
- [36] Robles R, Tudge CC, Dworschak PC, Poore GCB, Felder DL. Molecular phylogeny of the Thalassinidea based on nuclear and mitochondrial genes. In: Martin JW, Crandall KA, Felder DL, editors. Crustacean Issues 18: Decapod Crustacean Phylogenetics. Boca Raton, FL: Taylor and Francis/CRC Press; 2009. pp. 309-326
- [37] Lin FJ, Liu Y, Sha Z, Tsang LM, Chu KH, Chan TY, Liu R, Cui Z. Evolution and phylogeny of the mud shrimps (Crustacea: Decapoda) revealed from complete mitochondrial genomes. BMC Genomics. 2012;13:631-642
- [38] De Grave S, Pentcheff ND, Ahyong ST, Chan TY, Crandall KA, Dworschak PC, Felder DL, Feldmann RM, Fransen CHJM, Goulding LYD, Lemaitre R, Low MEY, Martin JW, Ng PKL, Schweitzer CE, Tan SH, Tshudy D, Wetzer R. A classification of living and fossil genera of decapod crustaceans. Raffles Bulletin of Zoology. 2009;21:1-109
- [39] Ahyong ST, Lowry JK, Alonso M, Bamber RN, Boxshall GA, Castro P, Gerken S, Karaman JS, Goy JW, Jones DS, Meland K, Rogers DC, Svavarsson J. subphylum crustacea brünnich, 1772. In: Zhang ZQ, editor. Animal Biodiversity: An Outline of Higherlevel Classification and Survey of Taxonomic Richness. Zootaxa. 2011;3148:165-191
- [40] Dworschak PC, Felder DL, Tudge CC. Infraorders Axiidea De Saint Laurent, 1979 and Gebiidea De Saint Laurent, 1979 (formerly known collectively as Thalassinidea). In:

Schran FR, Vaupel Klein JC, editors. The Crustacea – Treatise on Zoology: Anatomy, Taxonomy, Biology. Boston: Koninklijke Brill NV; 2012. Crustacea. 9B(69). pp. 109-219

- [41] Dworschak PC. Axiidea and Gebiidea (Crustacea: Decapoda) of Costa Rica. Annalen des Naturhistorischen Museums in Wien. 2013;115(B):37-55
- [42] Poore GCB, Ahyong ST, Bracken-Grissom HD, Chan TY, Chu KH, Crandall KA, Dworschak PC, Felder DL, Feldmann RM, Hyžný M, Karasawa H, Lemaitre R, Komai T, Li X, Mantelatto FL, Martin JW, Ngoc-Ho N, Robles R, Schweitzer CE, Tamaki A, Tsang LM, Tudge CC. On stabilising the names of the infraorders of thalassinidean shrimps, Axiidea de Saint Laurent, 1979 and Gebiidea de Saint Laurent, 1979 (Decapoda). Crustaceana. 2014;87:1258-1272
- [43] Sakai K, Turkay M. A review of the collections of the Infraorders Thalassinidea Latreille, 1831 and Callianassidea Dana, 1852 (Decapoda, Pleocyemata) lodged in three German museums, with revised keys to the genera and species. Crustaceana. 2014;87(2):129-211
- [44] Saint Laurent M. Vers une nouvelle classification des Crustacés Décapodes Reptantia. Bulletin de l'Office National des Pêches République Tunisienne. 1979;**3**:15-31
- [45] Dworschak PC. Methods collecting Axiidea and Gebiidea (Decapoda): A review. Annalen des Naturhistorischen Museums in Wien. 2015;**117**(B):5-21
- [46] Wear RG, Yaldwyn JC. Studies on Thalassinid Crustacea (Decapoda, Macrura Reptantia) with a Description of a New Jaxea from New Zelan and an Account of its Larval Development. Vol. 41. Zoology Publications from Victoria University of Wellington, New Zealand; 1966
- [47] Melo GAS. Manual de identificação dos Crustacea Decapoda do litoral brasileiro: Anomura, Thalassinidea, Palinuridea e Astacidea. São Paulo: Plêiade/FAPESP; 1999. p. 551
- [48] Oliveira DB, Martinelli-Lemos JM, Abrunhosa FA. The complete larval development of the mud shrimp *Upogebia vasquezi* (Gebiidea: Upogebiidae) reared in the laboratory. Zootaxa. 2014;**3826**(3):517-543
- [49] Dworschak PC. Global diversity in the Thalassinidea (Decapoda): An update (1998-2004). Nauplius. 2005;13(1):57-63
- [50] Sakai K. Upogebiidae of the world. Crustacean Monographs. 2006;6:1-185
- [51] Nucci PR, Melo GAS. First record of *Upogebia inomissa* Williams, 1993 (Decapoda, Thalassinidea, Upogebiidae) in Brazil. Nauplius. 2001;9(1):71-71
- [52] Barros MP, Pimentel FR. A fauna de Decapoda (Crustacea) do Estado do Pará, Brasil: lista preliminar das espécies. Boletim do Museu Paraense Emílio Goeldi, Zoologia. 2001;17(1):15-41
- [53] Oliveira DB, Silva DC, Martinelli JM. Density of larval and adult forms of the burrowing crustaceans *Lepidophthalmus siriboia* (Callianassidae) and *Upogebia vasquezi* (Upogebiidae) in an Amazon estuary, northern Brazil. Journal of the Marine Biological Association of the United Kingdom. 2012;92(2):295-303

- [54] Hernández-Ávila I, Lira C, Hernández G, Bolaños J. Upogebia vasquezi Ngoc-Ho, 1989 (Decapoda: Thalassinidea: Upogebiidae): First record for Venezuela. Boletin del Instituto Oceanografico de Venezuela. 2005;44(2):119-122
- [55] Souza Filho PWM. Costa de manguezais de macromaré da Amazônia: cenários morfológicos, mapeamento e quantificação de áreas usando dados de sensores remotos.
 Revista Brasileira de Geofísica. 2005;23(4):427-435
- [56] Burke L, Kura Y, Kassem K, Revenga C, Spalding M, Mcallister D. Coastal Ecosystems. Washington: World Resources Institute; 2001. disponível em: http://www.wri.org/ wr2000. (Acesso em fevereiro de 2015)
- [57] Vilhena MPSP, Costa ML, Berrêdo JF. Continental and marine contributions to formation of mangrove sediments in an Eastern Amazonian mudplain: The case of the Marapanim Estuary. Journal of South American Earth Sciences. 2010;29:427-438
- [58] Silva CA, Souza-Filho PW, Rodrigues SWP. Morphology and modern sedimentary deposits of the macrotidal Marapanim Estuary (Amazon, Brazil). Continental Shelf Research. 2008;**29**:619-631
- [59] Rosa-Filho JS, Girard TC, Frédou FL. Population dynamics of the burrowing shrimp *Lepidophthalmus siriboia* Felder and Rodrigues, 1993 (Reptantia: Axiidea: Callianassidae) on the Amazonian coast. Journal of Crustacean Biology. 2013;**33**(4):503-511
- [60] Laverock B, Smith CJ, Tait K, Osborn AM, Widdicombe S, Gilbert JA. Bioturbating shrimp alter the structure and diversity of bacterial communities in coastal marine sediments. The International Society for Microbial Ecology Journal. 2010;4:1531-1544
- [61] Laverock B, Kitidis V, Tait K, Gilbert JA, Osborn AM, Widdicombe S. Bioturbation determines the response of benthic ammonia-oxidizing microorganisms to ocean acidification. Philosophical Transactions of The Royal Society, Biological Sciences. 2013;368:20120441. Available from: http://dx.doi.org/10.1098/rstb.2012.0441
- [62] Laverock B, Tait K, Gilbert JA, Osborn AM, Widdicombe S. Impacts of bioturbation on temporal variation in bacterial and archaeal nitrogen-cycling gene abundance in coastal sediments. Environmental Microbiology Reports. 2014;6(1):113-121
- [63] Kornienko ES, Korn OM, Demchuk DD. The larval development of the mud shrimp Upogebia yokoyai Makarov, 1938 (Decapoda: Gebiidea: Upogebiidae) reared under laboratory conditions. Journal of Natural History. 2013;47(29-30):1933-1952
- [64] Sasaki A, Nakao H, Yoshitake S, Nakatsubo T. Effects of the burrowing mud shrimp, Upogebia yokoyai, on carbon flow and microbial activity on a tidal flat. Ecological Research. 2014;29:493-499
- [65] Webb AP, Eyre BD. Effect of natural populations of burrowing thalassinidean shrimp on sediment irrigation, benthic metabolism, nutrient fluxes and denitrification. Marine Ecology Progress Series. 2004;268:205-220
- [66] Kinoshita K. Burrow structure of the mud shrimp *Upogebia major* (Decapoda: Thalassinidea: Upogebiidae). Journal of Crustacean Biology. 2002;**22**(2):474-480

- [67] Berkenbusch K, Rowden AA. An examination of the spatial and temporal generality of the influence of ecosystem engineers on the composition of associated assemblages. Aquatic Ecology. 2007;41:129-147
- [68] Hernáez P, Wehrtmann IS. Population of the burrowing shrimp Callichirus seilacheri (Decapoda: Callianassidae) in northern Chile. Revista de Biología Tropical. 2007;55:141-152
- [69] Allanson BR, Skinner D, Imberger J. Flow in prawn burrows. Estuarine, Coastal and Shelf Science. 1992;**35**:253-266
- [70] Astall CM, Taylor AC, Atkinson RJA. Behavioural and physiological implications of a burrow-dwelling lifestyle for two species of Upogebiid mud-shrimp (Crustacea: Thalassinidea). Estuarine, Coastal and Shelf Science. 1997;44:155-168
- [71] Atkinson RJA, Froglia C, Arneri E, Antolini B. Observations on the burrows and burrowing behavior of *Brachynotus gemmellari* and on the burrows of several other species occurring on *Squilla* grounds off Ancona, Central Adriatic. Scientia Marina. 1998;62(1-2):91-100
- [72] Curran HA, Martin AJ. Complex decapod burrows and ecological relationships in modern and Pleistocene intertidal carbonate environments, San Salvador Island, Bahamas. Palaeogeography, Palaeoclimatology, Palaeoecology. 2003;192:229-245
- [73] Sepahvand V, Sari A, Tudge C, Bolouki M. A study of burrow morphology in representative axiidean and gebiidean mud shrimps, from the Persian Gulf and Gulf of Oman, Iran. Nauplius. 2014;22(2):137-144
- [74] Kinoshita K, Itani G, Uchino T. Burrow morphology and associated animals of the mud shrimp Upogebia yokoyai (Crustacea: Thalassinidea: Upogebiidae). Journal of the Marine Biological Association of the United Kingdom. 2010;90(5):947-952
- [75] Macginite GE. The natural history of *Callianassa californiensis* Dana. American Midland Naturalist. 1934;15(2):166-177
- [76] Santagata S. A waterborne behavioral cue for the actinotroch larva of *Phoronis pallida* (Phoronida) produced by *Upogebia pugettensis* (Decapoda: Thalassinidea). Biological Bulletin. 2004;**207**:103-115
- [77] Lima JF, Abrunhosa F, Coelho PA. The larval development of *Pinnixa gracilipes* Coelho (Decapoda, Pinnotheridae) reared in the laboratory. Revista Brasileira de Zoologia. 2006;23(2):480-489
- [78] Anker A, Jeng MS, Chan TY. Two unusual species of Alpheidae (Decapoda: Caridea) associated with upogebiid mud shrimps in the mudflats of Taiwan and Vietnam. Journal of Crustacean Biology. 2001;21(4):1048-1061
- [79] Peiró DF, Baeza JA, Mantelatto FL. Host-use pattern and sexual dimorphism reveals the mating system of the symbiotic pea crab *Austinixa aidae* (Crustacea: Brachyura: Pinnotheridae). Journal of the Marine Biological Association of the United Kingdom. 2013;93:715-723

- [80] Botter-Carvalho ML, Costa LB, Gomes LL, Clemente CCC, Carvalho PVVC. Reproductive biology and population structure of *Axianassa australis* (Crustacea, Axianassidae) on a sand-mud flat in north-eastern Brazil. Journal of the Marine Biological Association of the United Kingdom. 2015;95(4):735-745
- [81] Dumbauld BR, Chapman JW, Torchin ME, Kuris AM. Is the collapse of mud shrimp (*Upogebia pugettensis*) populations along the Pacific Coast of North America caused by outbreaks of a previously unknown bopyrid isopod parasit (*Orthione griffenis*)? Estuaries and Coasts. 2011;**34**:336-350
- [82] Chapman JW, Dumbauld BR, Itani G, Markham JC. An introduced Asian parasite threatens northeastern Pacific estuarine ecosystems. Biological Invasions. 2012;14:1221-1236
- [83] Ubaldo JP, Nanri T, Takada Y, Saigusa M. Prevalence and patterns of infection by the epicaridean parasite, *Gyge ovalis* and the emergence of intersex in the estuarine mud shrimp, *Upogebia major*. Journal of the Marine Biological Association of the United Kingdom. 2014;94(3):557-566
- [84] Selin NI. Some features of the biology of the mud shrimp Upogebia issaeffi (Balls, 1913) (Decapoda: Upogebiidae) from the Subtidal Zone of Vostok Bay, Sea of Japan. Russian Journal of Marine Biology. 2014;40(1):24-29
- [85] Henmi Y, Itani G. Burrow utilization in the goby *Eutaeniichthys gilli* associated with the Mud shrimp *Upogebia yokoyai*. Zoological Science. 2014;**31**(8):523-528
- [86] Pohle G, Santana W. Gebiidea and axiidea (= thalassinidea). In: Martin JW, Olesen J, Høeg JT, editors. Atlas of Crustacean Larvae. Baltimore, MD: Johns Hopkins University Press; 2014
- [87] Pohle G, Santana W, Jansen G, Greenlaw M. Plankton-caught zoeal stages and megalopa of the lobster shrimp *Axius serratus* (Decapoda: Axiidae) from the Bay of Fundy, Canada, with a summary of axiidean and gebiidean literature on larval descriptions. Journal of Crustacean Biology. 2011;**31**(1):82-99
- [88] Andryszak BL. *Upogebia affinis* (Say): Its postlarval stage described from Louisiana plankton, with a comparison to postlarvae of other species within the genus and notes on its distribution. Journal of Crustacean Biology. 1986;6(2):214-226
- [89] Melo SG, Brossi-Garcia AL. Postembryonic development of Upogebia paraffinis Williams, 1993 (Decapoda, Thalassinidea), reared under laboratory conditions. Nauplius. 2000;8(1):149-168
- [90] Abrunhosa FA, Pires MAB, Lima JF, Coelho-Filho PA. Larval development of *Lepidophthalmus siriboia* Felder and Rodrigues, 1993 (Thalassinidea) from the Amazon region, reared in the laboratory. Acta Amazonica. 2005;**35**(1):77-84
- [91] Gurney R. Notes on some Decapod Crustacea from the Red Sea. II. The larvae of *Upogebia* savignyi (Strahl). Proceedings of the Zoological Society of London. 1937;**107**B:85-101
- [92] Rabalais NN, Gore RH. Abbreviated development in decapods. In: Wenner AM, editor. Crustacean Growth: Larval Growth. (Gen. Ed. Frederik R. Schram, San Diego). Crustacean Issues. 1985;2:67-126. ISBN 9061912946

- [93] Forbes AT. An unusual abbreviated larval life in the estuarine burrowing prawn *Callianassa kraussi* (Crustacea: Decapoda: Thalassinidea). Marine Biology. Springer Berlin Heidelberg; 1973;22:361-365
- [94] Nates SF, Felder DL, Lemaitre R. Comparative larval development in two species of the burrowing ghost shrimp genus *Lepidophthalmus* (Decapoda: Callianassidae). Journal of Crustacean Biology. 1997;17(3):497-519
- [95] Strasser KM, Felder DL. Larval development in two populations of the ghost shrimp *Callichirus major* (Decapoda: Thalassinidea) under laboratory conditions. Journal of Crustacean Biology. 1999;19(4):844-878
- [96] Strasser KL, Felder DL. Larval development of the ghost shrimp *Callichirus islagrande* (Decapoda: Thalassinidea: Callianassidae) under laboratory conditions. Journal of Crustacean Biology. 2000;20(1):100-117
- [97] Kornienko ES, Korn OM, Golubinskaya DD. The complete larval development of the lobster shrimp *Boasaxius princeps* Boas, 1880 (Decapoda: Axiidea: Axiidae) obtained in the laboratory. Journal of Natural History. 2014;48(29-30):1737-1769
- [98] Kornienko ES, Korn OM, Golubinskaya DD. The number of zoeal stages in larval development of *Nihonotrypaea petalura* (Stimpson, 1860) (Decapoda: Axiidea: Callianassidae) from Russian waters of the Sea of Japan. Zootaxa. 2015;3919(2):343-361
- [99] Goy JW, Provenzano Jr. AJ. Larval development of the rare burrowing mud shrimp Naushonia crangonoides Kingsley (Decapoda: Thalassinidea; Laomediidae). Biological Bulletin. 1978;154(2):241-261
- [100] Strasser KM, Felder DL. Larval development of the mud shrimp Axianassa australis (Decapoda: Thalassinidea) under laboratory conditions. Journal of Natural History. 2005;39(25):2289-2306
- [101] Shenoy S. Studies on larval development in Anomura (Crustacea, Decapoda) II. Proceedings of the Symposium on Crustacea, II. Mandapam Camp: Marine Biological Association of India. 1967. p. 777-804
- [102] Ngoc-Ho N. The larval development of Upogebia darwini (Crustacea, Thalassinidea) reared in the laboratory, with a redescription of the adult. Proceeding of the Zoological Society of London. 1977;**181**:439-464
- [103] Santos A, Paula J. Redescription of the larval stages of *Upogebia pusilla* (Petagna, 1792) (Thalassinidea, Upogebiidae) from laboratory-reared material. Invertebrate Reproduction and Development. 2003;43(1):83-90
- [104] Kornienko ES, Korn OM, Demchuk DD. The larval development of the mud shrimp Upogebia issaeffi (Balss, 1913) (Decapoda: Gebiidea: Upogebiidae) reared under laboratory conditions. Zootaxa. 2012;3269:31-46
- [105] Abrunhosa FA, Melo M, Lima JF, Abrunhosa J. Developmental morphology of mouthparts and foregut of the larvae and postlarvae of *Lepidophthalmus siriboia* Felder and Rodrigues, 1993 (Decapoda: Callianassidae). Acta Amazonica. 2006;**36**(3):335-342

- [106] Abrunhosa FA, Arruda DCB, Simith DJB, Palmeira CAM. The importance of feeding in the larval development of the ghost shrimp *Callichirus major* (Decapoda: Callianassidae). Anais da Academia Brasileira de Ciências. 2008;80(3):445-453
- [107] Oliveira DB, Martinelli JM, Abrunhosa FA. Description of early larval stages of *Upogebia* vasquezi (Gebiidea: Upogebiidae) reared in laboratory. Journal of the Marine Biological Association of the United Kingdom. 2012;92(2):335-342
- [108] Oliveira DB, Martinelli-Lemos JM, Souza AS, Costa JR, Abrunhosa FA. Does retention or exportation occur in the larvae of the mud shrimp *Upogebia vasquezi* (Decapoda, Gebiidea)? Implications for the reproductive strategy of the species on the Amazon coast. Hydrobiologia. 2016;773:241-252
- [109] Costlow JD, Bookhout CG, Monroe R. The effect of salinity and temperature on larval development of *Sesarma cinereum* (Bosc) reared in the laboratory. Biological Bulletin. 1960;118(2):183-202
- [110] Anger K. The Biology of Decapod Crustacean Larvae. Crustacean Issues, Zoological Museum. University of Amsterdam, CRC Press; 2001. pp. 1-300
- [111] Hartnoll RG. Growth in Crustacea twenty years on. In: Paula JPM, Flores AAV, Fransen CHJM, editors. Advances in Decapod Crustacean Research. Hydrobiologia. Kluwer Academic Publishers, Netherlands; 2001;449:111-122
- [112] Paula J, Mendes RN, Paci S, Mclaughlin P, Gherardi F, Emmerson W. Combined effects of temperature and salinity on the larval development of the estuarine mud prawn Upogebia africana (Crustacea, Thalassinidea). Hydrobiologia. 2001;449:141-148
- [113] Anger K. Salinity as a key parameter in the larval biology of decapod crustaceans. Invertebrate Reproduction and Development. 2003;**43**(1):29-45
- [114] Queiroga H, Blanton J. Interactions between behaviour and physical forcing in the control of horizontal transport of decapod crustacean larvae. Advances in Marine Biology. 2005;47:107-214
- [115] Diele K, Simith DJB. Salinity tolerance of northern Brazilian mangrove crab larvae, Ucides cordatus (Ocypodidae): Necessity for larval export? Estuarine, Coastal and Shelf Science. 2006;68:600-608
- [116] Simith DJB, Diele K. O efeito da salinidade no desenvolvimento larval do caranguejo uçá, Ucides cordatus (Linnaeus, 1763) (Decapoda: Ocypodidae) no Norte do Brasil. Acta Amazonica. 2008;38(2):345-350
- [117] Simith DJB, Souza AS, Maciel CR, Abrunhosa FA, Diele K. Influence of salinity on the larval development of the fiddler crab *Uca vocator* (Ocypodidae) as an indicator of ontogenetic migration towards offshore waters. Helgolang Marine Research. 2012;66:77-85
- [118] Simith DJB, Pires MAB, Abrunhosa FA, Maciel CR, Diele K. Is larval dispersal a necessity for decapod crabs from the Amazon mangroves? Response of *Uca rapax* zoeae to different salinities and comparison with sympatric species. Journal of Experimental Marine Biology and Ecology. 2014;457:22-30

- [119] Tamaki A, Ingole B. Distribution of juvenile and adult ghost shrimps, *Callianassa japon-ica* Ortmann (Thalassinidea), on an intertidal sand flat: Intraspecific facilitation as a possible pattern-generating factor. Journal of Crustacean Biology. 1993;13(1):175-183
- [120] Kevrekidis T, Gouvis N, Koukouras A. Population dynamics, reproduction and growth of *Upogebia pusilla* (Decapoda, Thalassinidea) in the Evros Delta (North Aegean Sea).
 Crustaceana. 1997;70(7):799-812
- [121] Pezzuto PR. Population dynamics of Sergio mirim (Rodrigues 1971) (Decapoda: Thalassinidea: Callianassidae) in Cassino Beach, southern Brazil. Marine Ecology – Pubblicazioni del la Stazione Zoologica di Napoli I. 1998;19:89-109
- [122] Nates SF, Felder DL. Growth and maturation of the ghost shrimp *Lepidophthalmus sinu*ensis Lemaitre and Rodrigues, 1991 (Crustacea, Decapoda, Callianassidae), a burrowing pest in penaeid shrimp culture ponds. Fishery Bulletin. 1999;97:526-541
- [123] Berkenbusch K, Rowden AA. Latitudinal variation in the reproductive biology of the burrowing ghost shrimp *Callianassa filholi* (Decapoda: Thalassinidea). Marine Biology. 2000;**136**:497-504
- [124] Tamaki A, Miyabe S. Larval abundance patterns for three species of *Nihonotrypaea* (Decapoda: Thalassinidea: Callianassidae) along an estuary-to-open-sea gradient in Western Kyushu, Japan. Journal of Crustacean Biology. 2000;**20**(2):182-191
- [125] Kinoshita K, Nakayama S, Furota T. Life cycle characteristics of the deep-burrowing mud shrimp *Upogebia major* (Thalassinidea: Upogebiidae) on a tidal flat along the northern coast of Tokyo bay. Journal of Crustacean Biology. 2003;23(2):318-327
- [126] Rotherham D, West RJ. Patterns in reproductive dynamics of burrowing ghost shrimp *Trypaea australiensis* from small to intermediate scales. Marine Biology. 2009;156:1277-1287
- [127] Kaiser MJ, Attrill MJ, Jennings S, Thomas DN, Barnes DKA, Brierley AS, Hiddink JG, Kaartokallio H, Polunin NVC, Raffaelli DG. Marine Ecology: Processes, Systems, and Impacts. 2nd ed. Oxford: OUP; 2011. p. 528
- [128] Kramer MJ, Bellwood DR, Bellwood O. Benthic Crustacea on coral reefs: A quantitative survey. Marine Ecology Progress Series. 2014;511:105-116
- [129] Petracco M, Cardoso RS, Corbisier TN, Turra A. Brazilian sandy beach macrofauna production: A review. Brazilian Journal of Oceanography. 2012;60(4):473-484
- [130] Petracco M, Cardoso RS, Turra A. Patterns of sandy-beach macrofauna production. Journal of the Marine Biological Association of the United Kingdom. 2013;**93**(7):1717-1725
- [131] Souza JRB, Borzone CA, Brey T. Population dynamics and secondary production of *Callichirus major* (Crustacea: Thalassinidea) on a southern Brazilian sandy beach. Archives of Fisheries and Marine Research. 1998;46:151-164

- [132] Petracco M. Produção secundária da macrofauna bentônica da zona entremarés no segmento norte da praia do Uma, litoral sul do estado de São Paulo [Phd Thesis]. São Paulo: University of São Paulo; 2008. p. 254
- [133] Cardoso RS, Veloso VG. Population biology and secondary production of the sand-hopper *Pseudorchestoidea brasiliensis* (Amphipoda: Talitridae) at Prainha Beach, Brazil.
 Marine Ecology Progress Series. 1996;142:111-119
- [134] Petracco M, Veloso VG, Cardoso RS. Population dynamics and secondary production of *Emerita brasiliensis* (Crustacea: Hippidae) at Prainha Beach, Brazil. Marine Ecology. 2003;24(3):231-245
- [135] Veloso VG, Cardoso RS, Petracco M. Secondary production of the intertidal Macrofauna of Prainha Beach, Brazil. Journal of Coast Research. 2003;**35**:385-391
- [136] Souza JRB, Borzone CA. Population dynamics and secondary production of *Euzonus furciferus* Ehlers (Polychaeta, Opheliidae) in an exposed sandy beach of Southern Brazil. Revista Brasileira de Zoologia. 2007;24(4):1139-1144
- [137] Veloso VG, Sallorenzo IA. Differences in the secondary production of *Emerita brasiliensis* (Decapoda: Hippidae) on two sandy beaches in Rio de Janeiro State, Brazil. Nauplius. 2010;**18**(1):57-68
- [138] Petracco M, Cardoso RS, Corbisier TN, Turra A. Secondary production of sandy beach macrofauna: An evaluation of predictive models. Estuarine, Coastal and Shelf Science. 2012;115:359-365
- [139] Turra A, Petracco M, Amaral AC, Denadai MR. Population biology and secondary production of the harvested clam *Tivela mactroides* (Born, 1778) (Bivalvia, Veneridae) in Southeastern Brazil. Marine Ecology. 2015;36(2):221-234





IntechOpen