Fiddler crabs and their above-ground sedimentary structures: a review

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Abstract

Fiddler crabs are abundant, semi-terrestrial crustaceans inhabiting tropical, subtropical and warm temperate coasts worldwide. Some species build above-ground sedimentary structures at or near the opening to their burrows. The functions and shapes of these constructions vary interspecifically and according to the sex of the builder. Here, we compile the dispersed reports on these structures, suggest uniform naming for different shapes, review explanations for their functions and explore associations between the attributes of builders and their structures. We found that 47 fiddler crab species build at least one type (or subtype) of structure, including chimneys, hoods, pillars, semidomes, mudballs, and rims. Sedimentary structures show a strong association with sediment type as well as builder front type, genus and sex, but not with fiddler crab clade. Experimental studies have revealed distinct, sometimes multiple functions for some of these structures (e.g., female attraction, reduction of aggressive behavior and/or landmark orientation); however, most studies have been observational leaving the proposed functions of these structures for many species untested. Both field and laboratory experiments as well as observational studies can help us to better understand these sedimentary structures and their role in fiddler crab behavior and ecology.

Keywords External structures · Burrow · Chimney · Hood · Pillar · Semidome · Mudball · Rim

Introduction

Many animals build structures: ants and termites construct nests and underground galleries, beavers build dams, spiders weave webs, and so forth (Hansell 2007). Some structures have obvious benefits for their builders (e.g. shelter, display and ornamentation), yet the functions of many other structures remain unclear (Hansell 2007). Structures are regularly and appropriately viewed as physical manifestations of the behavioral phenotypes of builders (Dawkins 1982; Schaedelin and Taborsky 2009; Bailey 2012). When such constructions result from manipulation of the sediment in the form of tracks, burrows or pellets, they are called biogenic sedimentary structures, defined by Frey (1973) as “[a product of] the activity of an organism upon or within an unconsolidated particulate substrate”.

Fiddler crabs build biogenic above-ground sedimentary structures from mud or sand, sometimes more than one type per species, in or just above the intertidal zone (Crane 1975; Wolfrath 1992; Thurman et al. 2013; Mokhtari et al. 2015). There are 106 described species (102 living, 4 extinct) encompassing eleven genera and two subfamilies within Ocypodidae (Gelasiminae and Ocypodinae) (Rosenberg 2001; Shih et al. 2016). The group is distributed worldwide
The structures built by fiddler crabs are diverse in form and are built via different behaviors. Some are built only by males, others by females, and the social and ecological contexts in which they are built vary within and among species. Not surprisingly, there are several hypothesized functions for these structures. The most thoroughly and experimentally tested idea is that the relatively tall structures some courting males build by their burrows attract females because they elicit landmark orientation, i.e., the tendency of fiddler, ghost and related crabs, when away from their burrows, to move toward vertical objects to reduce their conspicuousness to predators and thereby reduce their predation risk (summarized by Christy and Wada 2015). Yet, recent experimental studies on other species showed that the attractive properties of a similar structure are sex- and context dependent, which challenges the claim that such structures elicit a common, shared and ancestral behavioral response (Kim et al. 2017). Far less well-studied are the structures erict a common, shared and ancestral behavioral response 

Literature survey and data analysis

Ethological studies have described and proposed functions for fiddler crab above-ground biogenic structures since the early twentieth century (see Yamaguchi et al.’s 2005 discussion for details). Among the most comprehensive studies, Crane (1975) described fiddler crab constructions as structures “erected beside the burrows, made of substrate” and classified them as three main shapes: chimneys, pillars and hoods. von Hagen (1968) highlighted that 12 fiddler crab species build structures, followed by Christy (1988a) who tallied a total of 14 species. Christy et al. (2001) then stated that at least 19 species construct sedimentary structures out of sand or mud including hoods, pillars, semidomes, and rims (or lips); subsequently, 20 chimney builders were identified by Shih et al. (2005) and 10 semidome builders were compiled by Carvalho et al. (2018).

Our review of the current literature found that at least 47 fiddler crab species (9 genera) build above-ground sedimentary structures (Table 1). We employed a narrative literature review strategy aimed to encompass all published literature on the topic: periodic searches from September 2014 to August 2019 using the term “fiddler crab” combined with “chimney”, “hood”, “shelter”, “oven”, “pillar”, “dome”, “semi-dome”, “semidome”, “mudball”, “rim”, “lip”, “above-ground structure” or “sedimentary structure” were performed using the ISI Web of Science and Google Scholar. Searches were followed by checking the reference list of papers for any possible missing studies. Seventy-six relevant papers on the topic were found (Table 1).

We employed a multiple correspondence analysis (MCA), used to identify patterns and associations of categorical variables (Greenacre and Blasius 2006), to explore relationships between descriptive categories of above-ground structure, sediment type, builder clade, genus, sex and front (i.e., width between the bases of eyestalks) (categories and data are specified in Table 1). Function of sedimentary structures was not included due to a lack of experimental data for some structures, including multifunctionality, which can lead to misinterpretation of MCA results. Sedimentary structures with vague and uncertain descriptions were also not included (all indicated with a ‘?’ in Table 1). Mixed sediments were categorized according to their predominant sediment type (sand for muddy sand; mud for sandy mud) to simplify data interpretation. Additionally, a Chi-Square test followed by Cramer’s V test, a measure of association for categorical variables, was performed to correlate types (subtypes) of sedimentary structures with the above-mentioned variables. The likelihood ratio (95% confidence interval) was assumed when the chi-square assumption was violated (McHugh 2013).

Types and subtypes of sedimentary structures: what is known?

Four types and two subtypes of above-ground sedimentary structures were recognized: chimney (Fig. 1a, 26 species); hood (or shelter/oven) (Fig. 1b, 11 species), further categorized into pillar (Fig. 1c, 6 species) and semidome (Fig. 1d, 11 species); mudball (Fig. 1e, 5 species); and rim (or lip) (Fig. 1f, 4 species). All fiddler crab genera, except for Petruca and Cranuca, represented by Petruca panamensis (Stimpson, 1859) and Cranuca inversa (Hoffmann, 1874), respectively, build at least one type of sedimentary structure (Fig. 2).

Chimney

Chimneys (also called funnels by Salmon 1987) are tall, tower-shaped structures with vertical walls that completely encircle the burrow entrance (Fig. 3a). The construction process is similar for all male and female builders (Table 1): crabs stack mud/muddy sand at the edge of the burrow entrance using their pereiopods and chelipeds (Shih et al. 2005; Gusmão-Junior et al. 2012). The source of chimney material is varied; some chimney builders use sediment dug

(Crane 1975; Rosenberg 2001) and is subdivided in two major clades, Atlantic-East Pacific (AEP) and Indo-West Pacific (IWP), estimated to have undergone differentiation around 17 million years ago (Sturmbauer et al. 1996; Shih et al. 2016).
Table 1 A summary of fiddler crabs that build above-ground sedimentary structures and their general characteristics (builder and sedimentary structure): clade (AEP: Atlantic-East Pacific, IWP: Indo-West Pacific), genus, species, front (B: broad, I: intermediate, N: narrow), sex (M: male, F: female, BS: both sexes), structure, function, type of study (E: experimental, O: observational, E/O: both), type of sediment used to build structures and associated references (bs: both sediments, *T. capricornis*, *T. elegans* and *T. flammula*).

| Clade | Genus | Species | Front type | Builder (sex) | Sedimentary structure | Sediment type | Function | Study type | References |
|-------|-------|---------|------------|---------------|----------------------|---------------|----------|------------|------------|
| IWP   | Austruca | *A. annulipes* (Milne-Edwards, 1837) | B           | BS*           | Mudball [both sexes, Burford et al., 2001b, Mokhlesi et al., 2011, Saher and Qureshi, 2017], pillar/hood/semidome [male, Mokhlesi et al., 2011] | Mud/sand      | –        | E/O        | Burford et al. (2001b), Mokhlesi et al. (2011), Saher and Qureshi (2017), Tina et al. (2017b) |
| IWP   | Austruca | *A. iranica* (Pretzmann, 1971) | B           | M             | Semidome             | Mud           | –        | E/O        | Saher and Qureshi (2017) |
| IWP   | Austruca | *A. lactea* (De Haan, 1835) | B           | M             | Hood/semidome        | Sand, mud     | Female attraction, landmark orientation, burrow defense | E/O | Crane (1975), Kim and Choe (2003), Kim et al. (2004), Yamaguchi et al. (2005), Muramatsu (2009), Muramatsu (2010a, 2010b), Kim et al. (2010, 2017), Zhu et al. (2012) |
| IWP   | Austruca | *A. mjoebergi* (Rathbun, 1924) | B           | M             | Semidome             | Mud           | –        | E/O        | Bourdiol et al. (2018) |
| IWP   | Austruca | *A. perplexa* (Milne-Edwards, 1852) | B           | M/F(?)        | Mudball (?), rim, pillar | Sand          | –        | E/O        | Christy (2007) |
| IWP   | Austruca | *A. occidentalis* (Naderloo, Schubart and Shih, 2016) | B           | M             | Semidome             | –            | –        | O          | P. R. Y. Backwell, pers. comm. in Bourdiol et al. (2018) |
| IWP   | Austruca | *A. sindensis* (Alcock, 1900) | B           | M             | Pillar/semidome      | Sandy mud     | –        | E/O        | Saher and Qureshi (2017) |
| IWP   | Austruca | *A. bengali* (Crane, 1975) | B           | BS            | Mudball              | Mud           | –        | O          | Chatterjee et al. (2014) |
| AEP   | Afruca  | *A. tangeri* (Eydoux, 1835) | B           | BS            | Mudball              | Mud           | Territorial marking, increase of male conspicuousness (?), female attraction | E/O | Günther (1963), Crane (1975), Oliveira et al. (1998), Latruffe et al. (1999), Burford et al. (2000, 2001a, 2001b, 2001c), Cummings et al. (2008) |
| IWP   | Gelasimus | *G. tetragonon* (Herbst, 1790) | N           | F             | Chimney, mudball     | Mud           | –        | O          | Crane (1975), Weis and Weis (2004) |
| IWP   | Gelasinus | *G. vocans* (Linnaeus, 1758) | N           | –             | Chimney              | Mud, silt, sand | –        | O          | De (2000) |
| Clade | Genus | Species | Front type | Builder (sex) | Sedimentary structure | Sediment type | Function | Study type | References |
|------|-------|---------|------------|---------------|-----------------------|---------------|----------|------------|------------|
| AEP  | Leptuca | L. beebei (Crane, 1941) | B | BS* | Pillar, rim (?) [male] // chimney [female] | Mud | Landmark orientation, female attraction [hood (pillar)], burrow defense | E/O | Peters (1955), Zucker (1974), Crane (1975), Christy (1987), Christy (1988a, 1988b), Christy and Schober (1994), Christy, pers. obs. [chimney], Backwell et al. (1995), Christy (1995), Koga et al. (1998), deRivera et al. (2003), Kim et al. (2009), Matsumasa et al. (2013) |
| AEP  | Leptuca | L. crenulata (Lockington, 1877) | B | M | Hood/rim | Silt, sand | – | O | deRivera (2003) |
| AEP  | Leptuca | L. cumulanta (Crane, 1943) | B | BS* | Hood [male] // chimney (?) [female] | Mud | – | O | Crane (1975) |
| AEP  | Leptuca | L. dorotheae (von Hagen, 1968) | B | M | Pillar | Sandy mud, mud | – | O | Crane (1975) |
| AEP  | Leptuca | L. latimanus (Rathbun, 1893) | B | M | Hood (?)/semidome | Mud, sand | Female attraction, reduction of aggressive behaviour | E/O | Zucker (1981), Pope and Haney (2008) |
| AEP  | Leptuca | L. leptodactyla (Rathbun, 1893) | B | M | Hood | Sand | Female attraction, burrow defense (?) | E/O | Matthews (1930), Crane (1975), Masunari (2012), de O. Rodrigues et al. (2016) |
| AEP  | Leptuca | L. musica (Rathbun, 1914) | B | M | Hood | Sand | – | O | Crane (1975), Pope and Haney, 2008 |
| AEP  | Leptuca | L. panacea (Novak and Salmon, 1974) | B | M | Rim | Muddy sand | – | O | Salmon et al. (1978), Christy (1988a) |
| AEP  | Leptuca | L. pugilator (Bosc, 1801) | B | M | Hood | Sand | Burrow defense (?) | E/O | Christy (1982) |
| AEP  | Leptuca | L. speciosa (Ives, 1891) | B | – | Chimney | Mud | – | O | Rodriguez-Tovar et al. (2014) |
| AEP  | Leptuca | L. stenodactylus (Milne-Edwards and Lucas, 1843) | B | F | Chimney, column/pillar | Muddy sand, sandy mud | – | O | Crane (1975), Müller (1989) |
| AEP  | Leptuca | L. subcylindrica (Stimpson, 1859) | B | – | Chimney (?) | Sand | Temperature and humidity regulation (?) | O | Thurman (1984) |
| Clade | Genus | Species          | Front type | Builder (sex) | Sedimentary structure | Sediment type | Function                                                                 | Study type | References |
|-------|-------|------------------|------------|---------------|-----------------------|---------------|---------------------------------------------------------------------------|------------|------------|
| AEP   | Leptuca | *L. terpsichores* (Crane, 1941) | B          | M             | Hood                  | Sand          | Female attraction, reduction of aggressive behaviour, landmark orientation | E/O        | Peters (1955), Zucker (1974, 1981), Christy et al. (2001, 2002, 2003a, b), Christy and Backwell (2006), Ribeiro et al. (2006), Kim et al. (2007), Kim and Christy (2015), Perez et al. (2016) |
| AEP   | Leptuca | *L. thayeri* (Rathbun, 1900) | I          | BS            | Chimney               | Mud           | Burrow defense (?), temperature and humidity regulation (?)              | E/O        | von Hagen (1970), Crane (1975), Salmon (1987), Gusmão-Júnior et al. (2012) |
| AEP   | Minuca | *M. galapagensis* (Rathbun, 1902) | B          | M             | Hood                  | Muddy sand    | Landmark orientation                                                     | E/O        | von Hagen (1968), Christy (1988a) |
| AEP   | Minuca | *M. herradurensis* (Bott, 1954) | B          | F             | Chimney               | Mud           |                                                                            | O          | Christy, pers. obs |
| AEP   | Minuca | *M. minax* (LeConte, 1855) | B          | M             | Chimney [Basan and Frey, 1976], chimney/dome [Montague, 1980], semidome (?) [Crane, 1975] | Mud          | Acoustic signal enhancement (?)                                           | E/O        | Crane (1975), Basan and Frey (1977), Montague (1980) |
| AEP   | Minuca | *M. pugnax* (Smith, 1870) | B          | BS*           | Chimney, semidome [male] | Mud         | Reduction of agonistic interaction (?), female attraction, acoustic signal enhancement (?), reduced erosion by tidal action (?) | E/O        | Crane (1975), Basan and Frey (1977), Montague (1980), Greenspan (1982) |
| AEP   | Minuca | *M. virens* (Salmon and Atsaiides, 1968) | B          | M             | Semidome              | Sand          |                                                                            | E/O        | Powers and Cole (1976) |
| AEP   | Minuca | *M. vocator* (Herbst, 1804) | B          | BS            | Chimney               | Mud           |                                                                            | O          | Crane (1943, 1975) |
| AEP   | Minuca | *M. rapax* (Smith, 1870) | B          | M             | Semidome              | Sand, muddy sand |                                                                            | O          | Carvalho et al. (2018) |
| IWP   | Paraleptuca | *P. chloropthalmus* (Milne-Edwards, 1837) | B          | F             | Chimney               | Mud           |                                                                            | O          | Crane (1975) |
| IWP   | Paraleptuca | *P. crassipes* (Adams and White, 1848) | B          | F             | Chimney               | Mud           |                                                                            | O          | Crane (1975) |
Table 1 (continued)

| Clade | Genus | Species | Front type | Builder (sex) | Sedimentary structure | Sediment type | Function | Study type | References |
|-------|-------|---------|------------|---------------|-----------------------|---------------|----------|------------|------------|
| IWP   | Tubuca| T. arcuata (De Haan, 1835) | N            | BS            | Chimney               | Mud           | Burrow defense | E/O        | Crane (1975), Wada and Murata (2000) |
| IWP   | Tubuca| T. capricornis (Crane, 1975) | N            | BS            | Chimney               | Mud           | Burrow defense | E/O        | Slatyer et al. (2008) |
| IWP   | Tubuca| T. coarctata (Milne-Edwards, 1852) | N            | BS            | Chimney               | Mud           | –         | O          | Crane (1975) |
| IWP   | Tubuca| T. dussumieri (Milne-Edwards, 1852) | N            | F             | Chimney               | Mud           | –         | O          | Crane (1975) |
| IWP   | Tubuca| T. elegans (George and Jones, 1982) | N            | M             | Chimney               | Mud, sandy mud | –         | O          | George and Jones (1982), Booksmythe et al. (2008) |
| IWP   | Tubuca| T. flammula (Crane, 1975) | N            | BS            | Chimney               | Mud           | –         | O          | Crane (1975) |
| IWP   | Tubuca| T. forcipata (Adams and White, 1848) | N            | F             | Chimney               | Mud           | –         | O          | Crane (1975) |
| IWP   | Tubuca| T. paradussumieri (Bott, 1973) | N            | BS            | Chimney               | Mud           | –         | O          | Ng and Sivasothi (1999) |
| IWP   | Tubuca| T. urvillei (Milne-Edwards, 1852) | N            | F             | Chimney               | Mud           | –         | O          | Crane (1975) |
| IWP   | Tubuca| T. rosea (Tweedie, 1937) | N            | BS            | Chimney               | Mud           | Burrow defense, reduction of agonistic interaction (?) | E/O        | Tina et al. (2017a, 2018a, b) |
| AEP   | Uca   | U. heteropleura (Smith, 1870) | N            | F             | Chimney               | Mud           | –         | O          | Christy, pers. obs |
| IWP   | Xeruca| X. formosensis (Rathbun, 1921) | N            | BS            | Chimney               | Mud           | Burrow defense (?) | E/O        | Shih et al. (2005) |

“(?)” indicates either possibly misidentified structure or an untested hypothesis for a function. Additional information is added in brackets when differences among papers are present. Asterisk in “builder (sex)” column indicates that males and females build different types of structures.
from within their burrows, while others collect with their walking legs parcels of sediment from the surface around the burrow entrance (Shih et al. 2005).

Several experimental studies have explored the functions of chimneys. Wada and Murata (2000) observed that Tubuca arcuata (De Haan, 1835) of both sexes build chimneys with higher abundance in the reproductive season of spring to summer. The authors suggested that chimneys may avoid burrow loss to an opponent based on an experiment where burrowless crabs were released near burrows with and without chimneys. Shih et al. (2005) described similar behavioral patterns in Xeruca formosensis (Rathbun, 1921), testing different hypotheses for chimney function (e.g., sexual attraction, sunshade effect). Both studies indicated that chimneys hide males from predators, while protecting females from other males.

Slatyer et al. (2008) suggested that chimney owners are “shy” crabs; crabs with chimneys generally spend more time in their burrows and less time feeding, minimizing their chances of being chased by intruders. It is still unclear whether chimneys directly or indirectly affect such behavior. The authors stated that Tubuca capricornis (Crane, 1975) chimneys hide the entrance to the burrow, making it more difficult for occasional intruders to find it, as previously mentioned for other species.

Chimney building in Leptuca thayeri (Rathbun, 1900) may also be related to burrow defense, mainly to females at late gonadal stages, thus enhancing reproductive success (Salmon

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**Fig. 1** The types (and subtypes) of above-ground biogenic sedimentary structures built by fiddler crabs: chimney (a), hood (b), pillar (subtype) (c), semi-dome (subtype) (d), mudball (e), rim (f) (drawings by FR de Grande). Drawings are not in the same scale.
Gusmão-Junior et al. (2012) stated that social context does not influence chimney construction, attributing this behavior to endogenous factors. Their main hypothesis is that chimneys may act in maintaining internal burrow conditions (Gusmão-Junior et al. 2012). As observed in Fallicambarus fodiens (Cottle, 1863) lobsters (Cambaridae) (Williams et al. 1974) and Cleistostoma dilatatum (De Haan, 1833) manicure crabs (Camptandriidae) (Kim et al. 2011), sedimentary structures (chimney and tower, respectively) may also help in the regulation of burrow evaporation and preservation of internal humidity. In this context, L. subcylindrica (Stimpson, 1859) also appear to use their above-ground sedimentary structures to maintain abiotic conditions (humidity and/or temperature), especially during dry periods (Thurman et al. 1984).

A series of studies on T. rosea (Tweedie, 1937) tested several functions for chimneys. The chimneys this species builds seem to reduce agonistic interactions and improve burrow defense against intruders (Tina et al. 2017a). Consistent with results by Slatyer et al. (2008), chimney owners spend less time feeding compared to crabs with no chimneys (Tina et al. 2018a). The energetic cost of building a chimney may require builders to feed faster, also as suggested by Slatyer et al. (2008). Tubuca rosea females tend to build longer and deeper burrows with higher chimneys than males’s chimneys, supporting the hypotheses that chimneys may help to maintain abiotic conditions necessary for successful incubation in the burrow (Gusmão-Junior et al. 2012; Tina et al. 2018b). In sum, these experimental studies indicate that chimneys help their builders defend their burrows. Chimneys may also affect the temperature or humidity of burrows but such effects and their possible benefits to builders have yet to be demonstrated.

Hood

Hoods are morphologically varied structures typically built by males, the only fiddler crab constructions to encompass three largely different forms, including semidomes, pillars and “typical” hoods. Generally, a hood (also known as shelter or oven) consists of a cupped, roughly semi-circular mound of sediment which extends above the burrow entrance (Fig. 3b, c). While some researchers initially referred to

![Fig. 2 Phylogenetic tree of the Ocypodidae family (Shih et al. 2016) with corresponding above-ground sedimentary structures described for each genus, clade (Atlantic-East Pacific, AEP and Indo-West Pacific, IWP) and subfamily, including the pyramid structures built by the ghost crabs Ocypode ceratophthalma (Pallas, 1772), O. kuhlii (De Haan, 1835) and O. saratan (Forskal, 1775) (Linsenmair 1967; Jones 1972)]
hoods as shelters (Yamaguchi 1971; Zucker 1974), shelters and hoods were later recognized as being the same structure (Crane 1975; Yamaguchi 2001). Zucker (1978) explained that “shelter” implies a protective function, whereas “hood” is a functionally neutral term for these structures. Semidomes, also known as half-domes, are more massive and robust structures (Zucker 1981) (Fig. 3d). On the other hand, a pillar is a narrow, tower-like structure—sometimes conical—which is also positioned at the edge of the burrow, often with a skirt encircling the burrow opening (Fig. 3e). The terms “semidome” and “pillar” will be used throughout this text as hood subtypes, even though these constructions are also considered to be in their own category in other studies.

Typical hood builders use their chelipeds, pereiopods and the dorsal region of the carapace to manipulate sediment collected from the surface or inside their burrows into the characteristic arching shape (Christy 1982; Yamaguchi et al. 2005). As seen by Christy et al. (2001), L. terpsichores (Crane, 1941) build their hoods in one to three activity bouts, during which they do not feed. Individual males build one or two hoods per courtship cycle and from 15% to about 60% of males will build a hood on days of peak courtship. Semidome and hood construction appears to be similar (Matthews 1930; Christy et al. 2001; Chatterjee et al. 2014). Interestingly, all courting male L. pugilator (Bosc, 1801) (Christy 1982) and L. latimanus (Rathbun, 1893) (Zucker 1981) build hood-shaped structures and Kim et al. (2004) found that over 80% of Austruca lactea (De Haan, 1835) males completed their semidome structures during low tide. As for the more delicate pillars, crabs scrape surface sandy mud, carry it to the burrow with their pereiopods and pile it at the edge of the burrow’s entrance with their legs and minor chelipeds (Crane 1975; Christy 1988a).

Using an experimental approach, Zucker (1974, 1981) provided evidence that L. beebei (Crane, 1941), L. latimanus and L. terpsichores with hoods are active primarily in the semi-circular area in front of these structures and that L. latimanus and L. terpsichores with hoods interact aggressively less often with their neighbors. Hoods built by L. terpsichores are also vital for juveniles avoiding conflict with adults (Zucker 1981). In contrast, crabs that do not build hoods defend a territory encompassing the entire area around the burrow (Zucker 1981). Christy (1988a) analyzed lengthy records of the locations and behavior of individual courting male L. beebei before and after they built pillars and was unable to corroborate the effects reported by Zucker (1981): pillars did not affect the distribution of male activity around the burrows nor reduce rates of aggressiveness toward males located “behind” these structures. Christy (1988a) discussed possible reasons why these two studies yielded different results.

Christy (1982) reported that burrow resident L. pugilator and M. pugnax (Smith, 1870) tend to brace themselves during fights for burrows by pressing their body against their hoods. Male L. pugilator usually build hoods from wet sand dug from the bottom of their burrows; as this sand dries out, salt crystals and desiccated silt act like cement increasing the rigidity of hoods and their resistance to fracture, giving residents a clear advantage over invading challengers (Christy 1982). Courting L. pugilator males always build hoods which are immediately repaired when damaged, e.g., during fights at the burrow entrance, highlighting the importance of this structure to resident builders independent of the presence of mate-searching females. In contrast, courting L. terpsichores and L. beebei build hoods and pillars, respectively, but far less often, and they usually do not rebuild these structures even when damaged 1.5–2 h after low tide, when most sexually receptive females would have found mates but burrow defense still remains important (Christy 1988b; Christy et al. 2001). Greenspan (1982) suggested that hoods in M. pugnax may also protect burrows from tidal erosion. Yet, at least for the hooded “breeding burrows” of L. pugilator found in the supra-tidal zone, tidal erosion is not a problem (Christy 1982).

Hoods also attract females and hood building is sexually selected by female choice (Christy et al. 2002). Courting male L. terpsichores build hoods at the time of the monthly and daily tidal cycles when most females seek mates (Christy et al. 2001). Small males tend to build relatively high structures, increasing their conspicuousness (Christy et al. 2001). Christy et al. (2002) experimentally showed that L. terpsichores females are differentially attracted to courting males that build hoods (approach rates: with hood = 0.84; without hood = 0.66; n = 354 approaches) even when replicas of hoods are added to the burrows of males that do not build hoods, suggesting that hoods contribute directly to male attractiveness. Most convincing were experiments in which hood replicas are positioned about 3 cm away from the entrances of males’s burrows; females orient to these structures rather than to the courting male about 40% of the time when the male does not directly lead the female to the entrance to his burrow. In contrast, Perez et al. (2016) did not find a female preference for L. terpsichores males with hoods, possibly due to the much smaller sample size in that study (n = 28 approaches). Similarly, de O. Rodrigues et al. (2016) also showed that hoods built by male L. leptodactyla attract females.

Male crabs also orient to their own hoods. In field experiments using displaced hoods, some males that move far from their burrows to intercept passing females lead the females not to their burrow entrance but to the nearby hood replica (Christy et al. 2002). This simple observation prompted a new line of study indicating that hood building is under sexual selection by indirect male–male competition to maintain burrow ownership as well as natural selection for the ability to visually relocate a burrow in a quick manner (Ribeiro
et al. 2006). Hoods not only assist males in orienting back to their burrow, but also enable males to move more freely when courting, suggesting they are visual beacons marking a male’s home burrow (Kim and Christy 2015). Heatwole et al. (2018) found, however, that males with hoods are no more likely to reach passing females and do not move greater distances from their burrows compared to males without hoods. Since males of this species typically move no more than 17 cm away from their burrows to attract females, hoods may not be beneficial as home beacons during most courtship interactions (Heatwole et al. 2018).

Hood building by male L. terpsichores appears to be selected by female choice, male–male competition for resource (burrow) holding, and predation on both sexes (Ribeiro et al. 2006). Sexual signals that are favored by both modes of sexual selection, typically operating in sequence, are not unusual (Hunt et al. 2009). It has been argued that females should prefer male traits that have been tested in combat because they will reliably advertise male condition, which might benefit females directly or via indirect genetic benefits to the offspring (Berglund et al. 1996). Although structure building (Backwell et al. 1995; Matsumasa et al. 2013) and investment in other courtship behavior (Kim et al. 2010) are condition dependent, hoods may attract both sexes and facilitate burrow location because they elicit landmark orientation. This “sensory trap” hypothesis (Christy 1995) as it pertains to female preference has received considerable experimental support (summarized by Christy 2007), including demonstration that the preference for pillars built by L. beebei (Kim et al. 2006) and the preference for hoods built by L. terpsichores (Kim et al. 2007) increase with the perceived risk of predation in females. Although females that choose males with structures may enjoy reproductive benefits by mating with males in good condition, such benefits cannot explain why males, females not seeking mates and females of species that do not build structures are attracted to them (Christy 2007).

The effects and functions of the semidomes built by A. lactea have also been well-described. Studies of this species in Japan use the word “hood” (e.g., Yamaguchi et al. 2005; Muramatsu 2010a, b); while Korean studies prefer “semidome” (e.g., Kim et al. 2004; Zhu et al. 2012). Field experiments showed that semidomes are sexual signals, reduce aggressive interactions and are landmarks for male orientation (see references in Table 1). Kim and Choe (2003) also showed that semidome building in A. lactea is related to the semilunar cycle of tidal inundation, which in turn affects food abundance: males that are given supplemental food build more semidomes. The frequency of semidome building, however, seems to be independent of food abundance, being instead dependent on environmental factors such as temperature (Takeshita et al. 2018).

Zhu et al. (2012) explored the effects of semidome size on female orientation. Females prefer courting males with normal-sized semidomes over those with half-sized or no semidome when seeking mates. When startled by a mock predator, females orient preferentially to empty burrows with full-sized semidomes over those with half-size or no semidome both during and outside the breeding period. When no predator is present, females differentially orient to burrows with structures (equally to full- and half-sized structures) over those without structures during the breeding period but not during the nonbreeding period. Hence, both level of risk and the reproductive state of the female may potentiate her orientation response to structures. Under reduced predation risk, female preference for larger structures appears to be contingent upon other stimuli from the courting male. Landmark orientation in service of mate choice (the response to the mimic in a sensory trap) and in service of risk reduction (the response selected by the model) may be flexibly linked to the benefit to females in both contexts. A recent study by Kim et al. (2017) further supports such flexibility; female A. lactea preferentially approach sedimentary structures (in test arenas without males) only during reproductive periods and males do not differentially orient to structures. Kim et al. (2017) suggested a divergent evolutionary path in the group for this behavior, where orientation to hoods by females during courtship and to vertical objects for risk reduction evolved independently. Interestingly, the closely related species A. mjoebergi rarely builds semidomes, due to loss of attractive function that is possibly mediated by predation and environmental changes (Bourdil et al. 2018).

Pillar building is a condition-dependent behavior, carried out by six species (see Table 1); male L. beebei, for example, build more pillars when given supplemental food (Backwell et al. 1995) and pillar builders allocate more energy and time to claw waving and less to feeding compared to non-builders. Females of this species approach structures not only for mating but also for protection (Christy 1988b; Christy et al. 2003b), which is consistent with the sensory trap hypothesis (Christy 1995; Kim et al. 2009). Although males with pillars attract more females to their burrows, once attracted to a male and burrow, females are equally likely to stay and mate whether or not the male has a pillar (Christy 1987). Pillars, thus, signal both male condition and the location of male...
burrows to breeding females (Christy 1988a, b; Backwell et al. 1995; Matsumasa et al. 2013). Under high predation risk, male L. beebei tend to build fewer pillars, either because males spend more time sheltering in their burrows or because fewer females wander to find mates, reducing the benefits of having a pillar (Koga et al. 1998; deRivera et al. 2003) despite the low energetic cost to build these structures (Matsumasa et al. 2013). Finally, pillars in other species are often regarded as “poorly-formed hoods” (Crane 1975).

Hoods in general, thus, appear to attract females, orient male movement and reduce antagonistic behavior and reduce predation risk for both sexes. Semidomes, as well as the possible functions mentioned above, may further aid in burrow defense. Pillars, on the other hand, seem to be exclusively related to orientation and mate attraction. Although the sensory trap hypothesis has helped explain why both sexes orient to sedimentary structures, recent studies suggest that orientation for risk reduction and in service of mate choice can be uncoupled and evolve independently. The sensory trap hypothesis should not be uncritically used to explain how structure building benefits males in all species.

**Mudball**

Mudballs are large, roughly spherical pellets of substrate that crabs remove from the burrow during excavation and deposit on the surface around the burrow entrance (Fig. 3f, g). All fiddler crabs that dig burrows make similar balls of sand or mud during burrow construction. Mudballing, when thought to have a social function, is the term used for this behavior in both male and female *Afruca annulipes*, *A. bengali* (Crane, 1975), *A. tangeri* (Eydoux, 1835), and *Gelasimus tetragonon* (Herbst, 1790). Mudballs and their deposition differ between sexes. Oliveira et al. (1998) stated that mudballs produced by male *A. tangeri* are larger than females’s, although this could be related to the larger volume of male burrows (Latruffe et al. 1999). Chatterjee et al. (2014) showed similar results in *A. bengali* mudballs and Tina et al. (2017b) observed a positive correlation between *A. annulipes* carapace width and number of mudballs. Also, females of *A. tangeri* and *A. annulipes* tend to deposit theirs in a mound near the burrow opening, while males usually position their mudballs far from the burrow entrance (Oliveira et al. 1998; Burford et al. 2001b; Tina et al. 2017b). Interestingly, Burford et al. (2001c) observed that some females exhibit a male-like mudballing behavior, with similar structural arrangement of mudballs; the authors suggested that their masculinization may either be a consequence of parasite infestation or constitute a behavior to avoid harassment. Mudball function remains unclear in *A. bengali* and *G. tetragonon* (Weis and Weis 2004; Chatterjee et al. 2014).

Mudballing behavior in *A. tangeri* was first described in the early 1960s (Günther 1963). In previous studies, mudballs were described as a consequent product of excavation (Crane 1975). Behavioral studies with this species, however, showed that both sexes spend more than two hours on mudball maintenance (Burford et al. 2001c), and also suggested that their function is to define territorial boundaries and prevent invasion and/or conflict with other individuals (Oliveira et al. 1998; Burford et al. 2000, 2001a).

Sharing a similar purpose, structures built by the genus *Hyoploax* (Stimpson, 1858), which are made from surface sediments, have also been extensively studied. Barricades and/or fences near burrows are built to deter the approach of potential competitors and maintain territory (Wada 1984, 1994; Wada et al. 1998; Christy and Wada 2015).

Oliveira et al. (1998) proposed that *A. tangeri* mudballs attract females, with males producing larger and more numerous mudballs than females. The number of mudballs may be also related to the degree to which the burrow collapsed during the previous high tide. Latruffe et al. (1999) claimed that females may relate male structures and mudball distance to the depth of the male’s burrow and, consequently, to burrow quality for breeding (Latruffe et al. 1999), but subsequent studies found little support for this function (Burford et al. 2001a). Males invest more in the claw waving display to attract females during the courtship phase than in mudball maintenance (Burford et al. 2001a).

Mudballs may also increase male conspicuousness when exposed to different backgrounds, although Cummings et al. (2008) stated that these hypotheses required more studies to determine that specific role. At present, the likely function of mudballs lies in territorial behavior.

**Rim**

Rims or lips are circular, elevated rings of sediment built around the edge of the burrow opening (Fig. 3h). Male crabs usually mold rims from sand with their major claw and ambulatory legs (Salmon et al. 1978; deRivera 2003; Christy 2007). No experiments to determine the functions of rims have been done. Salmon et al. (1978) noticed that intruder male *L. panacea* often destroy the owner’s burrow entrance. Additionally, Christy (2007) suggested that *A. perplexa* rims may make burrow openings more visible from a distance and help both males and approaching females locate the burrow.

**Results and discussion**

**Are fiddler crab above-ground sedimentary structures associated with general attributes?**

The first two factorial axes of the MCA explained 28.89% of the variance (F1: 18.30%, F2: 11.59%, Fig. 4) and the complementary Cramer’s V test showed that types (subtypes) of
sedimentary structures are strongly associated with front, genus, sex and sediment, but not clade (Tables 2 and 3).

Table 2: Chi-Square and Cramer’s V results comparing sedimentary structures with clade, front, genus, sex and sediment data

| Variables     | Chi-square | df | p   | Cramer’s V | Value |
|---------------|------------|----|-----|------------|-------|
| Sedimentary   | Clade      | 4.831 | 5   | 0.437      | 0.289 |
|               | Front      | 26.352 | 10  | **0.003**  | 0.443 |
|               | Genus      | 59.321 | 40  | **0.025**  | 0.446 |
|               | Sex        | 37.112 | 10  | **0.001**  | 0.533 |
|               | Sediment   | 33.134 | 10  | **0.001**  | 0.515 |

Statistically significant p-values are in bold.

A variety of types (subtypes) of structures are associated in Cluster 1, being explained by the AEP clade and the

Fig. 4: Multiple correspondence analysis. Plots of the following variables on the first two dimensions (F1 and F2) with the eigenvalue: clade (AEP: Atlantic-East Pacific, IWP: Indo-West Pacific), genus (Austruca, Afruca, Gelasimus, Leptuca, Minuca, Paraleptuca, Tubuca, Uca, Xeruca), front type (broad, narrow, intermediate), sex (male, female, both sexes), structure (chimney, hood, pillar, semidome, mudball, rim), type of sediment used to build structures [mud (including sandy mud), sand (including muddy sand), both types].
genera *Leptuca* and *Minuca*. Otherwise, with few exceptions, builders of the IWP clade form two groups: hood-builders of the genus *Austruca* (Cluster 2) and chimney-builders of *Gelasimus, Tubuca* and *Xeruca* (Cluster 3). As mentioned, clade was the only attribute that did not correlate with structure type, where all shapes are present at least once in both clades (Fig. 2). Clades (AEP and IWP) are expressively divergent and paraphyletic (Shih et al. 2016) indicating a non-phylogenetic relationship in the evolutionary history of fiddler crab structure types. However, fiddler crab genera correlated with structure type, which suggests that structures readily diversified due perhaps to social and environmental factors. From these novel inferences, further studies could explore intra-generic relationships using phylogeny-based comparative approaches.

There is an apparent relationship between above-ground sedimentary structures and sediment type. Upon close inspection of the summarized data, chimneys seem to be more common in muddy/muddy sand habitats, pillars are more common in sandy mud, hoods/semidomes and rims are usually built in sand, and mudballs are a by-product of burrow construction in muddy sediments (see references in Table 1). These associations indicate that the cohesiveness of sandy mud may facilitate construction of delicate structures, such as pillars, which probably cannot be built of more friable sand (Christy 1988b). Field experiments in which replicas of hoods were placed on the burrows of courting male *L. beebei* and replicas of pillars were placed on the burrows of courting male *L. terpsichores* showed that hoods, the larger of the two structures (structure height was equal), are most attractive to females of both species (Christy et al. 2003a). However, *L. beebei* males with hoods do not mate at a greater rate than those with pillars because hoods may visually shield some passing females, affecting male–female encounter rates. Hence, there was no evidence that signal competition currently selects for hood building by pillar building *L. beebei*. Experimental studies manipulating sediment type and properties are needed to test these ideas. Sediment saturation also influences structure shape: Kim et al. (2004) hypothesized that semidome building is usually limited by water content of the soil, showing that *A. lactea* males are stimulated to build semidomes when the surrounding area is not dry and hard. This may explain why hood building typically occurs before low tide in *L. terpsichores* (Christy et al. 2001) and why *L. pugilator* use moist sand from the bottom of their burrows to build hoods (Christy 1982). Additionally, Mokhlesi et al. (2011) observed that the muddy areas inhabited by *A. annulipes*, a multi-structure builder, allow building behavior; whereas the sandy and drier sediments in which *A. sindensis* (Alcock 1900) burrows limit structure building. These effects of soil moisture on construction behavior have been observed in other groups; for example, Grow (1981) found that chimney building by the crayfish *Cambarus diogenes diogenes* (Girard, 1852) depends on the percentage of soil moisture. Experimental studies on the effects of sediment type and moisture content on fiddler crab structure building have yet to be conducted.

The majority of chimney builders gathered in Cluster 3 seems strongly associated with the narrow-front characteristic and the genera *Gelasimus, Tubuca* and *Xeruca*. Both sexes build chimneys, with these structures serving as barriers against potential invaders (Slatyer et al. 2008; Wada and Murata 2000; Gusmão-Junior et al. 2012; Tina et al. 2017a). Narrow-front species tend to mate on the surface close to burrows built by females, and these females are typically more aggressive toward intruders than are females of broad-front species (Christy and Salmon 1991). As females tend to build more robust and taller structures than males, chimneys are likely to play an important role in female incubation behavior in these species.

**Toward a common nomenclature**

Authors have used different terms for the same structures built by fiddler crabs. Peters (1955) and Montague (1980) attributed chimney building (“towers”) to *M. pugnax*; while, both Crane (1975) and Greenspan (1982) showed that this species builds hoods. Crane (1975, page 284) observed chimney building behavior in one population of *L. stenodactylus* in the Panama Canal Zone, but Müller (1989) called

| Variable          | F1     | F2     |
|-------------------|--------|--------|
| Clade—AEP        | −0.626 | −0.741 |
| Clade—IWP        | 0.626  | 0.741  |
| Genus—*Afruca*   | 0.101  | −1.182 |
| Genus—*Austruca* | −0.302 | 1.499  |
| Genus—*Gelasimus*| 1.650  | −0.349 |
| Genus—*Leptuca*  | −0.841 | −0.615 |
| Genus—*Minuca*   | −0.464 | −0.817 |
| Genus—*Paraleptuca* | 0.669  | −0.107 |
| Genus—*Tubuca*   | 1.607  | 0.131  |
| Genus—*Uca*      | 1.057  | −1.876 |
| Genus—*Xeruca*   | 1.676  | 0.103  |
| Front—Broad       | −0.513 | 0.071  |
| Front—Intermediate| −0.055 | −1.889 |
| Front—Narrow      | 1.559  | −0.165 |
| Sex—Both sexes    | 0.874  | −0.201 |
| Sex—Female        | 0.821  | −0.640 |
| Sex—Male          | −0.695 | 0.248  |
| Sediment—Mud      | 0.477  | −0.276 |
| Sediment—Sand     | −0.932 | 0.413  |
| Explained variance (%) | 18.30  | 11.59  |
such structures “pillars or columns”. *Leptuca stenodactylus* populations, however, have been extensively observed in Panama with no indications of such structure building behavior (JH Christy, pers. obs.). This species often overlaps with the hood builder *L. terpsichores*, which may have caused misinterpretations. Some species were nevertheless observed building more than one type of structure, highlighting the importance of detailed and concise ethological descriptions.

The lack of simplified definitions for fiddler crab structures may explain some inconsistencies in the current literature. Pope and Haney (2008) described in detail both *L. musica* (Rathbun, 1914) and *L. latimanus* hood structures showing their differences and context-dependent variations in shape. Upon close inspection of the authors’ figures and description of *L. latimanus* structures, we believe that *L. latimanus* hoods could be instead semidomes; the same applying to Zucker (1981) when using hoods instead of semidomes. Similarly, Mokhlesi et al. (2011) described four distinct types and subtypes of structures built by *A. annulipes* (Milne-Edwards, 1837), differentiating between hoods, pillars, mudballs and semidomes; the semidome structures shown in their figures appear to be yet another type of above-ground sedimentary structure. This species, thus, builds four structures in addition to this novel structure. Moreover, in their review of African fiddler crabs, Peer et al. (2015) ascribed dome building to *G. hesperiae* (Crane, 1975) [previously *U. (T.) vocans hesperiae*] but we have been unable to find this passage in Crane’s work.

Above-ground sedimentary structures seem to have a crucial role for juvenile and smaller crabs. Benefits such as burrow protection may lead them to build higher or even a larger number of structures than adults in some species (Günther 1963; Zucker 1974, 1981; Wada and Murata 2000). There is, however, an additional behavior observed by Günther (1963) where smaller-sized *A. tangeri* rarely build “turret buildings”. This unusual and undescribed type of structure was mentioned by Crane (1975) and more recently attributed to *C. inversa* (Atkinson and Eastman 2015). Günther (1963) distinguished that structure from the “walled holes” (chimneys) constructed by *M. vocator* (Crane, 1943). Until now, no other studies have provided detailed descriptions of the “turret” structures built by fiddler crabs. Future ethological studies could reveal if turrets are just a by-product of excavated pellets or an actual functional structure.

## Conclusions and future directions

Fiddler crabs and their above-ground sedimentary structures have been used as a model in ecological and ethological studies. Researchers have proposed several hypotheses concerning structure function for a few species, yet most remain untested experimentally. Explanations for structure function that have been experimentally tested include courtship signaling and biasing of activity space, reduction of territorial overlap, visual orientation, refuge from predation, intruder exclusion and reduction of aggression between neighbors. Future studies should evaluate structure function using equivalent experimental designs on species that build similar constructions. This, along with the use of simplified terms for these structures, would facilitate comparisons between groups, especially when referring to hood subtypes, i.e., pillars and semidomes (Table 4).

We envisage an integration between field observations and experiments in which structures are altered or removed to determine their effects on social interactions and burrow microclimate, with varying sediment characteristics. Apart from their functions in relation to fiddler crabs, we also lack an understanding of how these structures may affect sediment biogeochemistry and habitat for associated groups, especially meiofauna and microbial communities (Bell et al. 1978; Kristensen 2008; González-Ortiz et al. 2014; Nobbs and Blamires 2015; Citadin et al. 2016). A combined approach should give us a better understanding of the relationship between architecture, function and ecological effects of these biogenic constructions.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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